

Near-atomic cryo-EM structure of PRC1 bound to the microtubule

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Proteins that associate with microtubules (MTs) are crucial to generate MT arrays and establish different cellular architectures. One example is PRC1 (protein regulator of cytokinesis 1), which cross-links antiparallel MTs and is essential for the completion of mitosis and cytokinesis. Here we describe a 4-Å-resolution cryo-EM structure of monomeric PRC1 bound to MTs. Residues in the spectrin domain of PRC1 contacting the MT are highly conserved and interact with the same pocket recognized by kinesin. We additionally found that PRC1 promotes MT assembly even in the presence of the MT stabilizer taxol. Interestingly, the angle of the spectrin domain on the MT surface corresponds to the previously observed cross-bridge angle between MTs cross-linked by full-length, dimeric PRC1. This finding, together with molecular dynamic simulations describing the intrinsic flexibility of PRC1, suggests that the MT-spectrin domain interface determines the geometry of the MT arrays cross-linked by PRC1.

PRC1 | microtubules | cryo-EM | MAPs | cytoskeleton

Cells rely on the microtubule (MT) cytoskeleton to help organize organelles (1), control cellular morphology (2), provide mechanical stability (3, 4), and form the spindle apparatus used to segregate chromosomes during cell division (5–8). The diverse functions of MTs are made possible through the action of motors (i.e., kinesin and dynein) and nonmotor MT-associated proteins (MAPs) that tightly regulate the MT network. Some of these proteins act by binding directly to the ends of MTs, either to the highly dynamic plus end, such as the conserved end-binding proteins (9, 10), or to the minus end (11, 12). Other MT regulators, such as MAP1 and MAP2/Tau, bind along the MT lattice, stabilizing it and helping build parallel arrays, most notably in axons (13, 14). Members of the MAP65 family, which includes human protein regulator of cytokinesis 1 (PRC1) and its budding yeast ortholog Ase1, form antiparallel MT arrays important for setting the spindle midzone and determining the location of the cytokinetic ring (8, 15–19). In addition to binding selectively to antiparallel MTs, PRC1 recruits other spindle-organizing factors and therefore is an essential component of the mitotic spindle (15, 20). Proper functioning of PRC1 requires cell-cycle-dependent localization and regulation. PRC1 contains two nuclear localization signals (NLSs) in its C-terminal domain (Fig. 1A) and is found almost exclusively in the nucleus during interphase (21–23). As the cell enters mitosis, PRC1 localizes to the spindle and becomes concentrated at the midzone by late anaphase (22), similar to observations for Ase1 (24). PRC1 is subject to phosphorylation by several cyclin-cyclin-dependent kinase (CDK) complexes (21) and together with kinesin-4 can regulate MT antiparallel overlap at the spindle midzone (20, 23, 25, 26).

Previous studies have identified that the dimerization domain of PRC1 is within the N-terminal domain (Fig. 1A) and that the primary MT binding region relies on a set of conserved residues within the central spectrin domain (PRC1-S) (15, 22, 26). However, the C-terminal domain (PRC1-C) also was found to contribute to the

MT-binding activity (15), and the specific contacts made between the MT and PRC1 were unclear. We used cryo-EM to visualize a monomeric PRC1 construct containing the spectrin domain and the full C-terminal domain (PRC1-SC) bound to the MT. The near-atomic resolution of our cryo-EM reconstruction allowed us to build an atomic model into the density and to explain, at the single amino acid level, the key interactions between PRC1 and the MT surface. We also found that PRC1-SC promotes MT assembly for both dynamic and taxol-stabilized MTs, suggesting that PRC1-SC stabilizes the MT lattice in a manner distinct from that of taxol. We propose that the unstructured C-terminal domain of PRC1-SC likely forms electrostatic contacts with the neighboring protofilament, thereby accounting for the increase in binding affinity when this C-terminal domain is included. Additionally, we found that the binding mode of the spectrin domain of PRC1-SC dictates the cross-bridge angle between antiparallel, cross-linked MTs, and we further investigated the validity of such a model using MD simulations.

Results

Features of the PRC1–MT Cryo-EM Reconstruction. Full-length PRC1 consists of four domains: a dimerization domain, a rod domain, a spectrin domain, and a C-terminal domain (Fig. 1A). The dimerization and rod domains are required to orient the spectrin and C-terminal domains at opposite ends of an extended homodimer

Significance

PRC1 (protein regulator of cytokinesis 1) is critical to cellular architecture through its interaction with microtubules to form antiparallel microtubule arrays, like those in the spindle midzone. Here, cryo-EM studies describe, in close to atomic detail, the interaction of PRC1 with the microtubule surface. Together with previous studies, our structure leads to a model of how PRC1 promotes the establishment of stable, higher-order microtubule arrays.

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Data deposition: Atomic coordinates have been deposited in the Protein Data Bank (PDB ID code [5KMG](#)). The cryo-EM reconstruction has been deposited in the Electron Microscopy Data Bank (accession code [EMD-8266](#)).

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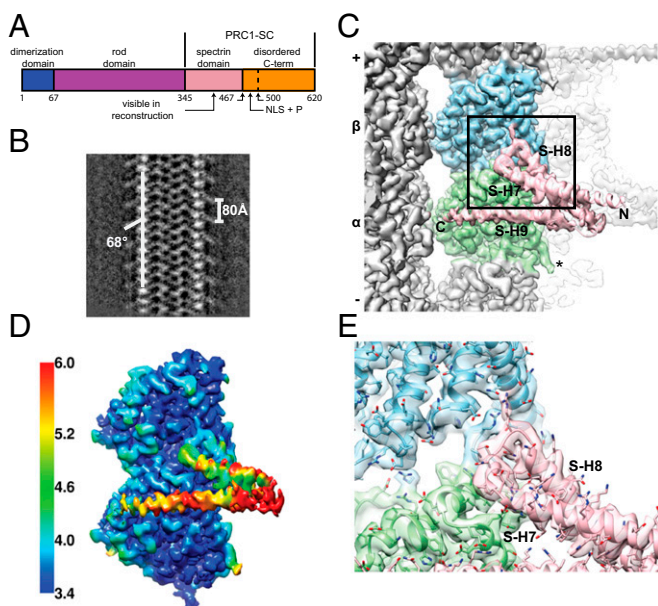


Fig. 1. The 4-Å resolution cryo-EM structure of the spectrin domain of PRC1 bound to the microtubule. (A) Domain structure of full-length PRC1 and of the PRC1-SC construct used in this study. Residues 470–500 contain the NLSs and phosphorylation sites (P). (B) 2D class average for an MT segment bound to PRC1-SC showing the $\sim 70^\circ$ angle between the spectrin domain of PRC1 and the MT wall. (C) Side view of the cryo-EM reconstruction showing the bound PRC1 spectrin domain in pink and the tubulin dimer to which it is bound in green (α -tubulin) and blue (β -tubulin). The tail of α -tubulin is indicated by *. Although the construct used contains both the spectrin and C-terminal domain, only the spectrin domain (residues 345–467) is visible. (D) Segmented cryo-EM density map of the asymmetric unit, PRC1-SC bound to a tubulin dimer, displaying local resolution estimation. (E) Close-up of the box in C showing the density map and the atomic models for the spectrin domain of PRC1 (pink) and α - and β -tubulin (green and blue, respectively).

and for PRC1’s cross-linking and bundling of MTs (15). MT bundling would result in MT images that cannot be processed because of overlap; therefore, to obtain a reconstruction with the highest possible resolution, we chose to focus on the PRC1-SC construct consisting of only the spectrin and C-terminal domains (Fig. 1A), which are the MT-binding regions. Raw images showed MT binding by PRC1-SC (Fig. S1A), and 2D class averages display clear density corresponding to PRC1-SC decorating the MT lattice with an ~ 80 -Å repeat (i.e., binding once per $\alpha\beta$ -tubulin dimer) (Fig. 1B and Fig. S1B). Using $\sim 230,000$ tubulin heterodimers, corresponding to 17,069 overlapping MT segments (Materials and Methods), we generated a 3D reconstruction of an MT with bound PRC1-SC with an overall resolution better than 4 Å (Fig. 1 and Fig. S1C). The final density map (Fig. 1C) shows high-resolution features corresponding to α -helical register, β -strand separation, and side-chain densities for larger amino acids (Fig. 1 and Fig. S2), as expected for a nearly atomic-resolution cryo-EM map. However, local resolution estimation reports a range of resolutions throughout the density map that reflects conformational or compositional heterogeneity (Fig. 1D). The resolution for tubulin is generally between 3.4 and 4 Å, but the resolution for the PRC1-SC molecule is lower, in the range of 3.8–6 Å, and is progressively poorer as the radius becomes larger, i.e., as the molecule becomes more distant from its contact with the MT surface. This lower resolution could be caused by two different factors: PRC1-SC occupancy on the MT and/or its flexibility. We therefore attempted to quantify the percentage of tubulin-binding sites occupied by PRC1-SC on the MT lattice. Because PRC1-SC recognizes a single binding site per tubulin heterodimer, and tubulin dimers are regularly spaced at ~ 80 Å along the MT lattice, the repeating pattern of PRC1-SC binding will contribute to the intensity

of a layer line in Fourier space corresponding to that distance (i.e., $\sim 1/80$ Å $^{-1}$). Note that this layer line is very weak for naked MTs without a “decorating” binding factor because of the similarity between α - and β -tubulin. By comparing the intensity of this layer line (and higher orders) in our final reconstruction with the layer lines obtained from a simulated map containing full PRC1-SC decoration (27, 28) (Materials and Methods), we estimate that PRC1-SC occupied $\sim 49\%$ of all possible MT-binding sites in the MT images used for the reconstruction (Fig. 1 and Fig. S1D). Thus, this “compositional” heterogeneity (i.e., partial occupancy) must be a factor contributing to the comparatively lower resolution of PRC1-SC as compared with tubulin. However, we suspect that conformational heterogeneity also may contribute to this disparity. Although the local resolution for PRC1-SC is highest (3.8 Å) at the point of interaction with tubulin, it is noticeably lower (5–6 Å) near the N and C termini. In fact, the density for the C-terminal domain beyond residue 467 (the end of the spectrin domain) is not visible in our high-resolution map. We reasoned that although the C-terminal domain may not adopt a single, well-defined structure, it could adopt a limited range of compact conformations that would be observable at 10- to 20-Å resolution. However, even low-pass filtering the reconstruction to this resolution did not reveal any additional density attributable to this domain, confirming predictions based on results from Phyre2 (29) and META-Disorder (30) that PRC1’s C terminus is disordered.

In the reconstruction we can clearly identify the fold of the PRC1 spectrin domain (PRC1-S), consisting of a three-helix bundle (Fig. 1C) of helices [S-H7, S-H8, and S-H9, following previous nomenclature (26)]. Furthermore, the previously reported crystal structure (15, 26) fits well within the observed density. In a previous cryo-EM reconstruction of PRC1-SC limited to ~ 20 -Å resolution, docking of the crystal structure suggested that the PRC1 spectrin domain was at an $\sim 90^\circ$ angle with respect to the MT surface (15). In contrast, both our 2D class averages (Fig. 1B and Fig. S1B) and our high-resolution 3D reconstruction (Fig. 1C) clearly show that PRC1-SC projects from the MT surface at an angle of $\sim 70^\circ$. Interestingly, previous tomographic studies reported a 70° cross-bridge angle between antiparallel MTs cross-linked by full-length PRC1 (i.e., PRC1 including the N-terminal dimerization domain) (15). Our present work now indicates that the cross-bridge angle does in fact originate from the orientation of the spectrin domain with respect to the MT lattice.

Molecular Basis of PRC1-SC Binding Specificity. To understand the atomic details of the MT–PRC1-SC interaction, we generated an atomic model of PRC1-SC bound to tubulin using previous atomic models of PRC1 (15) and the MT (27) as starting points. From these initial models, we retraced and refined the atomic coordinates to fit the final density map better, using methods optimally suited for near-atomic (4 Å or better) cryo-EM reconstructions (Materials and Methods). In particular, the loop between S-H7 and S-H8 (residues 374–384, from now on referred to as S-H7H8 loop) was either absent or involved in crystal contacts in the previously determined crystal structures (15, 26), necessitating de novo loop-tracing. The presence of clear density for this region in our reconstruction supports the idea that the S-H7H8 loop is likely disordered in solution but becomes ordered upon binding to the MT lattice. Our resulting atomic structure shows how the spectrin domain binds at the intradimer interface along a protofilament. Interactions with tubulin involve primarily the S-H7H8 loop (Figs. 1E and 2A and B) [notice that side-chain densities for negatively charged residues are missing from cryo-EM density maps, likely because of radiation damage during early electron exposure (31)], with additional contacts made through S-H9 (Fig. 2C). The S-H7H8 loop makes a critical contact with helix H12 in β -tubulin. R381 in PRC1-S interacts with N424 in β -tubulin and potentially with D427 (Fig. 2A). Additionally, N380 in PRC1-S is well positioned (3 Å between donor and acceptor atoms) to hydrogen bond with

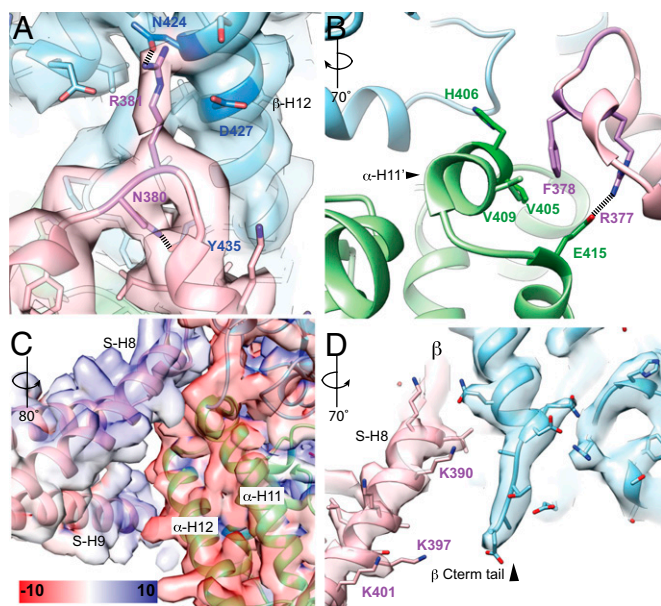


Fig. 2. Model of PRC1–tubulin interactions. (A) R381 and N380 in the loop between helices S-H7 and S-H8 of PRC1-SC (pink) are positioned to contact D427, N424, and Y435 in the H12 helix of β -tubulin (blue). (B) F378 of PRC1-SC nestles into a hydrophobic pocket formed by V405 and V409 in α -tubulin (green), and R377 of PRC1-SC likely contacts E415 of α -tubulin. (C) Helix H9 of PRC1-SC forms a complementary electrostatic surface with tubulin (the colored surface shown corresponds to the atomic model). Helix H9 is rich in basic residues (blue), whereas tubulin surface in its close proximity is negatively charged. Electrostatic units are $k_B T/e^-$. (D) Part of the β -tubulin tail is structured in our MT–PRC1-SC reconstruction, likely as a result of its stabilization by interactions with the multiple lysines (K390, 397, 401) in S-H8 of PRC1. All rotations for the views displayed in B–D are relative to the orientation shown in A.

β -tubulin's Y435 (Fig. 2A). F378 in PRC1-S nestles into a hydrophobic pocket in α -tubulin formed by V405 and V409 (within the segment connecting α -H11 and α -H12, denoted " α -H11'") (Fig. 2B). Additionally, R377 in PRC1-S is positioned to form a salt bridge with E415 in α -tubulin (within H11) (Fig. 2B).

Although S-H9 lies over the α -tubulin subunit, only its C-terminal portion is within interacting distance of α -H12 (Fig. 2C). The position of S-H9 and the abundance of basic residues across homologs in this region (residues 439–456) (Fig. S3) suggest that S-H9 interacts via electrostatic interactions with negative charges on the nearby MT surface.

The acidic C-terminal tails of tubulin are generally unstructured and, with a few notable exceptions, are not visible in most MT cryo-EM reconstructions (32, 33). It is thus noteworthy that we can clearly trace up to five extra amino acids from the last residue seen in previous cryo-EM studies (27, 34) for both α - and β -tubulin (Figs. 1C and 2D). The length and conformation of the tubulin tails we see here are similar to previous reports of structured tubulin tails (32, 33) indicating that the tubulin tails become partially structured upon PRC1-SC binding. The β -tubulin tail is near lysines K390, K397, and K401 on the surface of S-H8. The α -tubulin tail, in contrast, has no structured PRC1-SC density in close contact (Fig. 1C). However, the C terminus of PRC1-S is ~ 30 Å from the C-terminal tail of α -tubulin, placing it within an appropriate range to interact with the disordered C-terminal domain of PRC1-SC (which comprises >100 amino acids). Together, these features suggested that PRC1-SC binding could be mediated in part by the tubulin tails.

Because of the importance of PRC1 in mitosis, it is reasonable to assume that residues critical for MT binding should be well conserved across species. To gauge the importance of the ascribed PRC1-S–tubulin interactions, we aligned several PRC1 sequences

across a diverse set of eukaryotes and mapped the conservation onto our atomic model (Fig. S3). We find that the described critical interactions (displayed as spheres) correspond well to highly conserved residues. PRC1-S residues R377, F378, N380, and R381, observed to make direct contacts with tubulin, are 100% conserved in all examined homologs ranging from yeast to human. Residue K390, positioned to interact with the C-terminal tubulin tail, is also highly conserved and is followed by a conserved stretch of positively charged residues, supporting our proposal that basic residues in this region are required to interact with tubulin's acidic C-terminal tail. These observations are further supported by alanine mutagenesis showing that residues R377, R381, and K390 contribute significantly to PRC1-S's MT-binding activity (15). Some residues that are highly conserved but do not participate in PRC1-S–tubulin interactions appear to play a supportive structural role (e.g., PRC1-S E388 hydrogen bonds with R377, which in turn interacts with α -tubulin E415). The residues in helix S-H9 (439–456) are less conserved than the S-H7/H8 loop but maintain general basic character, supporting our model in which S-H9 makes electrostatic interactions with the acidic MT surface.

Molecular Basis of PRC1-SC's Specificity for the Intradimer Interface.

Each PRC1-SC binds the MT specifically at an intradimer interface (within an $\alpha\beta$ -tubulin heterodimer), using a small footprint on the tubulin surface that is significantly conserved between α - and β -tubulin ($\sim 70\%$ overall). The calponin homology (CH) domain of Ndc80, a kinetochore component, also binds tubulin using a very small region of the MT surface. In that case, the conservation of the tubulin-binding site between α - and β -tubulin is so significant that human Ndc80 also binds at the interdimer interface, where the α - and β -tubulin segments of the footprint are swapped with respect to those in the intradimer contact (35, 36). Thus, we decided to investigate the basis of PRC1-SC's specificity for the intradimer over the equivalent interdimer interface. Although the residues involved in contact with PRC1-S are generally conserved in α - and β -tubulin, a number of notable exceptions may explain the observed specificity (Fig. 3A): (i) β -tubulin D427, which is close to PRC1-S R381, is substituted at the interdimer interface by α -tubulin A427, thus disrupting a likely salt bridge; (ii) β -tubulin Y435, which hydrogen bonds with PRC1-S N380, corresponds to V435 in α -tubulin at the interdimer interface; (iii) the hydrophobic tubulin pocket filled by PRC1-S F378 at the intradimer interface is disrupted in the interdimer interface by the presence of a polar residue, β -tubulin T409 (in place of α -tubulin V409 at the intradimer interface) (Fig. 3B). Finally, α -tubulin G416, whose small size allows PRC1-S R450 to interact with neighboring charged amino acids (α -tubulin E414 and α -tubulin E420), is substituted by β -tubulin M416 and likely sterically occludes PRC1-S R450 from interacting with these residues at the interdimer interface (Fig. 3C). We propose that these critical sequence changes between α - and β -tubulin both sterically and chemically confer PRC1's selectivity for the tubulin intradimer interface.

PRC1's Disordered C-Terminal Domain Forms Electrostatic Interactions with the Neighboring Protofilament

In our reconstruction, the C-terminal domain beyond residue 467 is not visible, verifying that it has no regular structure (see *Features of the PRC1-MT Cryo-EM Reconstruction*). However, the reconstruction shows that the end of S-H9, which precedes the C-terminal domain, is oriented so that the C-terminal domain of PRC1-SC could interact with a neighboring α -subunit close to α -H3. Electrostatic maps, computed using the Adaptive Poisson–Boltzmann Solver (37), reveal that the site in the neighboring protofilament, potentially adjacent to the C terminus of PRC1 (i.e., α -H11 and the loop connecting α -H9 and α -S8), is negatively charged (Fig. 4), further suggesting that it forms a complementary electrostatic surface for the predominantly positively charged C-terminal domain of PRC1. The lack of density for the C-terminal domain means it does not make

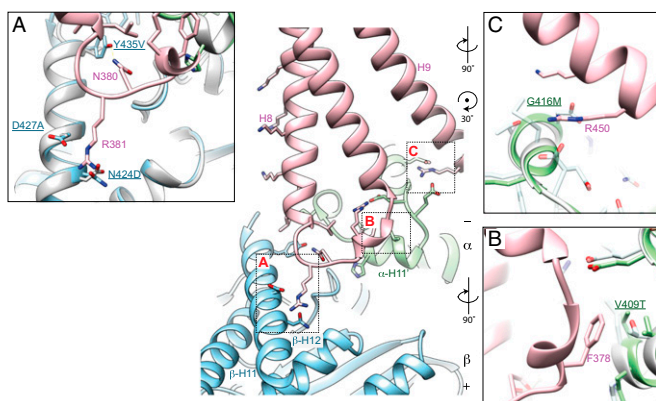


Fig. 3. Molecular basis for the specificity of PRC1-SC for the intradimer interface. A number of critical PRC1-S–tubulin contacts are present only at the intradimer interface. PRC1-S–tubulin in the center panel is rotated 180° in-plane from the view shown in Fig. 2A, and the MT is tilted out of plane to be viewed from the plus end of the microtubule. (A) β -Tubulin D427 at the intradimer site corresponds to α -tubulin A427 at the interdimer site, eliminating electrostatic interactions with R381 in PRC1. β -Tubulin Y435 is α -tubulin V435 at the interdimer site, which eliminates hydrogen bonding to N380 in PRC1-S. (B) α -Tubulin V409 is β -tubulin T409 at the interdimer site, reducing the hydrophobicity of the tubulin pocket that cradles PRC1-S F378. (C) α -Tubulin G416 is β -tubulin M416 at the interdimer site, potentially occluding PRC1-S R450 sterically from interacting with tubulin.

specific contacts with the MT, indicating instead that it can make multiple interactions to increase the affinity of PRC1-SC for MTs. Because the interface between PRC1-S and tubulin is relatively small ($\sim 500 \text{ \AA}^2$), the C terminus would provide favorable interactions to target PRC1-SC to the MT lattice and could increase PRC1-SC's binding affinity for MTs.

PRC1-SC Promotes MT Assembly. The results of our electrostatic calculations lead to the prediction that PRC1-SC should facilitate MT assembly by bridging across protofilaments. We tested whether PRC1-SC enhances MT assembly by performing cosedimentation assays. PRC1-SC and MTs were mixed together, and the reaction mixtures were centrifuged to separate large complexes (i.e., MTs and bound proteins) from unbound, soluble proteins (i.e., unpolymerized tubulin and unbound PRC1-SC) to determine how the proteins partitioned between the pellet and supernatant fractions (*Materials and Methods*).

Under our experimental conditions, PRC1-SC self-pellets to some extent (Fig. 5A and B, lanes PRC1), but most of the protein remains in the supernatant fraction. When using dynamic MTs and substoichiometric amounts of PRC1-SC, we observed significantly greater MT assembly in the presence of PRC1-SC, as shown by the increase in the amount of tubulin in the pellet fractions and a corresponding decrease in the supernatant fractions (Fig. 5A and B, lanes MTs and MTs+PRC1-SC). To replicate better the sample conditions that we imaged, we performed the experiment with taxol-stabilized MTs at much lower concentrations of MTs (as allowed by this stabilizing drug) and with a molar excess of PRC1-SC. We again observed an increase in MT assembly in the presence of PRC1-SC (Fig. 5C and D, lanes TMTs and TMTs+PRC1).

Based on the presence of density for the C-terminal tails of $\alpha\beta$ -tubulin in our reconstruction (Fig. 2D), we also tested whether these tails contributed to PRC1-SC binding and if they were required for PRC1-SC to promote MT assembly. MTs were assembled and then treated with subtilisin to cleave the C-terminal tails of tubulin (38). A similar amount of PRC1-SC bound to both native and subtilisin-cleaved MTs (Fig. S4A and B), clearly indicating that, at least at the concentrations used for this study, the C-terminal tails of tubulin do not contribute significantly to the apparent binding

affinity of PRC1-SC. Concerning PRC1 promotion of tubulin assembly, an increase in tubulin polymerization was observed in the presence of PRC1-SC for both the dynamic subtilisin-treated MTs (Fig. S4A and B) and the taxol- and subtilisin-treated MTs (Fig. S4C and D) similar to that observed for full-length tubulin. [Notice that subtilisin-treated MTs polymerize to a greater extent than untreated MTs, as reported previously (39).] This result indicates that the C-terminal tails of tubulin are not needed for PRC1-SC promotion of MT assembly, at least under the conditions we tested.

Kinesin, Dynein, and PRC1 Partially Share a Tubulin-Binding Site.

PRC1 is not the only protein that has been found to bind to MTs by recognizing the tubulin intradimer interface (see ref. 40 for a recent review). In particular, the MT-interacting domains of the motors dynein and kinesin are both known also to bind across the tubulin intradimer interface. Specifically, the H11' connection between the C-terminal α -H11 and α -H12 has been previously identified as a major part of the binding interface for both the kinesin motor head (41, 42) and the dynein MT-binding domain (MTBD) (43). This tubulin region is near the site where we observe the S-H7H8 loop in PRC1-S interacting with tubulin. To investigate whether there are specific residues in tubulin that are shared in the interaction with kinesin, PRC1, and dynein, we performed sequence and structural alignments for these proteins, using tubulin as the common element to superimpose the MT-MAP models (*Materials and Methods*). Although as yet there is no atomic-resolution information on the binding of the dynein MTBD to MTs, hybrid models of this interaction have been generated by integrating diverse sources of data, including crystal structures (41–47), medium-resolution EM maps (43, 47), and biochemical data (48, 49). Superposition of the MT–kinesin atomic model with our MT–PRC1-SC atomic model identified kinesin R278 and PRC1 R381 as functionally similar in their interaction with tubulin (potentially with β -tubulin D420 and β -tubulin E427) (Fig. 6A). Superposition of the MT–dynein MTBD model with our MT–PRC1-SC model revealed that dynein has two lysine residues (dynein K3298 and dynein K3299) in a position approximately equivalent to PRC1 R381, one of which may interact with tubulin in a similar manner (Fig. 6B). Previous mutational analyses identified these dynein residues as critical to the dynein–tubulin interaction (48). To identify additional residues that may be important for the interaction of tubulin with these motors, we chose a 4.0- \AA cutoff for the distance between tubulin and the binding partner to inspect potential contacts (dark green residues in Fig. 6). α -Tubulin H406 and α -tubulin V409, within the same hydrophobic pocket, are both within range potentially

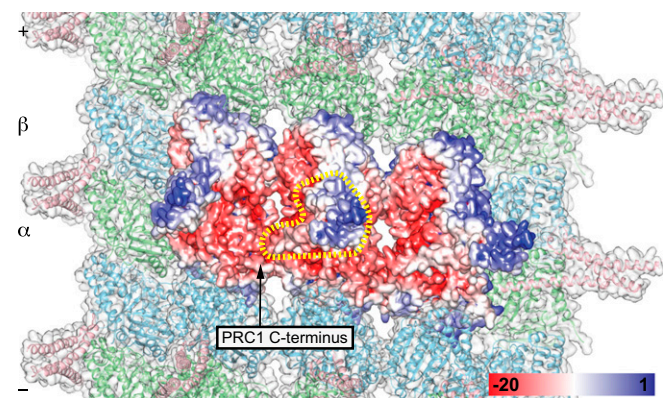


Fig. 4. Electrostatic map of three PRC1-SCs bound to three tubulin dimers in adjacent protofilaments. The positively charged, unstructured C terminus of PRC1-SC (composed primarily of arginine and lysine) is positioned to form complementary electrostatic interactions with the negatively charged (red) tubulin surface. The dashed yellow line outlines PRC1-S. Electrostatic units are $k_B T/e^-$.

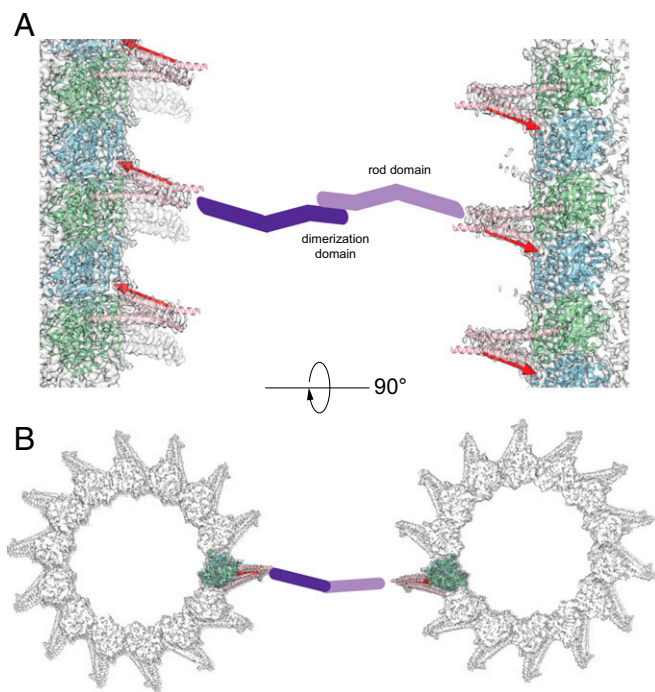


Fig. 7. The spectrin domain is oriented to interact preferentially with antiparallel microtubules. Atomic models of $\alpha\beta$ -tubulin and PRC1-SC (green, blue, and pink, respectively) for one protofilament are docked into the full-density maps (gray). The dimerization and rod domains are shown as cartoons in two purple hues. The side view (A) and end-on view (B) of two antiparallel microtubules (spaced by 30 nm to reflect the spacing and cross-bridge angle experimentally observed in previous cryo-tomography studies of PRC1-cross-linked MTs) (15) show how the orientation of the spectrin domains promotes antiparallel microtubule cross-linking.

angle (Ω) defined by two planes, each containing a spectrin domain (Fig. 8B), starts at $\Omega \sim 50^\circ$ in the crystallographic structure and can range from $30\text{--}210^\circ$ (a torsional angle of 180° is required for the antiparallel arrangement of cross-linked MTs as displayed in Fig. 7), with values around $70\text{--}100^\circ$ at equilibrium. Interestingly, conformations with spectrin domains pointing in the same direction ($\Omega = 0^\circ$) are not observed in our long-scale simulations, indicating that their relative orientation is partially constrained against parallel configurations. The conformational variability of the molecule, including the relative orientations of the spectrin domains, can be visualized directly in [Movie S1](#). We argue that the natural tendency of PRC1 toward an elongated conformation, together with an antiparallel predisposition of the spectrin domains as observed in the simulations, agrees with previous tomographic data on PRC1-bundled MTs and supports the selectivity of PRC1 in cross-linking antiparallel MTs.

Discussion

Using cryo-EM and atomic modeling, we have visualized the near-atomic details of the PRC1–MT interaction and thus have defined how PRC1 specifically recognizes the tubulin–intradimer interface. Critical specific interactions involve residues in the S-H7H8 loop of PRC1 that are nearly 100% conserved from yeast to human. In contrast to these specific contacts, helix S-H9, which is visualized in our study at lower resolution, likely because of intrinsic flexibility, uses a conserved basic character (i.e., lysine/histidine/arginine) for electrostatic interactions with the negative charge on the interacting surface of tubulin, which likely extends to the acidic C-terminal tails.

The specific contacts between PRC1 and tubulin identified by our atomic model are still likely to be insufficient to account for PRC1's binding affinity for MTs. Previous cosedimentation assays that

measured the binding affinities of various PRC1 constructs for assembled MTs demonstrated that the spectrin domain alone (PRC1-S) has weak affinity for the MT lattice (15, 22). A fivefold increase in binding affinity for MTs is achieved by including the arginine/lysine-rich C-terminal disordered domain ($K_d \sim 3.3 \mu\text{M}$ for PRC1-S vs. $0.6 \mu\text{M}$ for PRC1-SC) (15). Despite the presence of nearly 100 residues beyond helix S-H9 in our construct, we do not see density for any of these residues, as is consistent with the previous proposal that the C-terminal domain of PRC1 is unstructured. Because of the orientation of PRC1's H9, which precedes the C-terminal domain, and as supported by electrostatic calculations, we propose that PRC1's disordered basic C terminus interacts with a negative patch on the neighboring protofilament to increase PRC1's binding affinity for the MT surface. Logical extension of this model dictates that, by binding across protofilaments, PRC1 should facilitate MT assembly. Indeed, our cosedimentation assays show that more tubulin polymerizes in the presence of PRC1-SC. Interestingly, PRC1 promotes the polymerization even in the presence of the potent MT-stabilizing drug taxol, detectably reducing the critical concentration of assembly in the presence of the drug. This result suggests that the action of PRC1-SC on MT assembly may involve a molecular mechanism different from that of taxol, making the effects additive.

The PRC1 C-terminal domain contains phosphorylation sites at T470 and T481 that are known substrates for cyclin–CDK and whose phosphorylation state is tightly coupled to the cell cycle (21, 22). When PRC1 is phosphorylated, its MT bundling activity is reduced (22), as is consistent with a model in which the negative charge of phosphate groups would disrupt the electrostatic interactions between tubulin and PRC1. The presence of a disordered region under phosphoregulation is commonly observed for other MT-binding proteins, such as Ndc80 (33), CENP-E (52), and Bim1 (53).

The surface of tubulin recognized by PRC1's S-H7H8 loop is partially shared by kinesin and the dynein MTBD. Strikingly, both PRC1 and kinesin position an absolutely conserved arginine to interact with conserved residues in β -tubulin H12. Although PRC1 and the motors kinesin and dynein all have a single binding site per tubulin dimer, at the intradimer interface, the CH domain of human Ndc80 within the Ndc80 kinetochore complex binds with double that stoichiometry at both the intra- and the interdimer interfaces (33, 35, 54). The Ndc80 CH lacks the arginine residue that may be important to determine the specificity for β -tubulin N424 (with which PRC1 R381 and kinesin R278 interact). Instead, residues such as the conserved glutamic acid E415, present in both α - and β -tubulin, and lysine/arginine residues on the C-terminal end

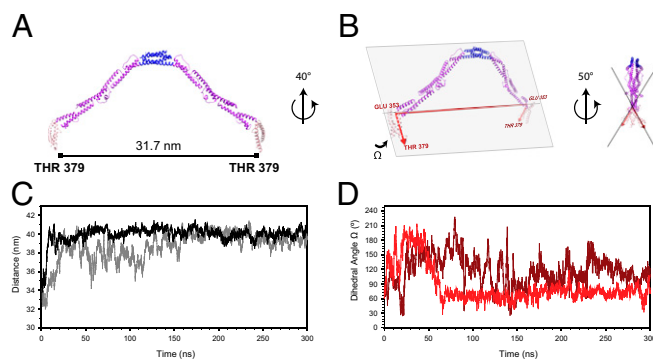


Fig. 8. Relaxation of the PRC1 dimer structure along the MD trajectories into an extended conformation. (A) Crystallographic structure of PRC1NS Δ C with the C α –C α distance between two residues (T379) located in the L7 loop at both ends of the homodimer. (B) The dihedral angle (Ω) between the two spectrin domains monitored from the initial crystallographic conformation. (C and D) Time evolution of the distance (C) and angle Ω (D) during two independent MD simulations showing the distance increasing to a more extended conformation and the constrained angle fluctuating in the range of $70\text{--}100^\circ$ at equilibrium.

of H11 for both α - and β -tubulin mediate critical interactions with Ndc80 at the intradimer and interdimer interfaces. Because several of the well-studied MT-binding proteins appear to recognize tubulin in overlapping regions of the MT surface (40), and given the sheer number of MT-binding proteins still to be characterized structurally and in their binding to tubulin, it is tempting to speculate that there are classes of MAPs that use defined sets of binding motifs and therefore compete with each other for MT real estate (40). This will be an important area for future cryo-EM-based structural studies.

We find that the spectrin domain of PRC1 binds to the MT surface at an $\sim 70^\circ$ angle, pointing to the minus end. This angle is consistent with the angle of the cross-bridges between cross-linked MTs previously observed by cryo-electron tomography of MT-PRC1 bundles (15). Although initially this angle was attributed to the dimerization and rod domains, we find here that the geometry is determined by the orientation of the spectrin domain with respect to the MT lattice. The conformation of the full-length homodimer adopted in the reported crystal structure (26) is incompatible with the antiparallel arrangement that would be inferred from our model, and we hypothesize that this conformation likely results from the effect of crystal contacts on the dimer structure. Our MD simulations show that when the structure is allowed to relax unconstrained, it quickly adopts an even more extended configuration that exhibits a large degree of torsional flexibility. The equilibrium separation between the spectrin domains was consistent in both our simulations and is compatible with the 35 ± 2 -nm spacing observed by tomography (15). Our simulations also revealed that the relative orientation of the spectrin domains is partially restricted to a range of 30 – 210° . The modular, restrained flexibility of the PRC1 molecule limits the range of preferred orientations of the spectrin domain, likely increasing the efficiency of MT capture and the antiparallel cross-linking of MTs while reducing the requirement for preexisting antiparallel MTs to establish a cross-bridge. On the other hand, the antiparallel binding orientation of the PRC1 spectrin domain (Fig. 7) and the conformations observed in the MD simulations of the PRC1 dimer help explain how PRC1 would selectively cross-link antiparallel MTs. In the spindle midzone, where multiple PRC1 molecules are present on each MT, the antiparallel bias of PRC1 dimers would accumulate to help arrange and stabilize regions of overlapping MTs. Furthermore, the ability of PRC1 to promote tubulin assembly likely contributes to the increased stability of the spindle midzone.

The findings of this work are summarized in Fig. 9. When bound to an MT, the spectrin domain of PRC1 adopts a fixed 70° angle with respect to the MT, but the rest of the dimer exhibits modular flexibility, increasing the tolerance for the possible orientations of the second MT when an initial cross-bridge is formed (Fig. 9A). When a second MT is bound, the preference for PRC1 to adopt antiparallel spectrin configurations would bias the antiparallel MT cross-linking (Fig. 9B). Binding of multiple PRC1 molecules would further steer the MTs into an antiparallel arrangement (Fig. 9D). Nevertheless, PRC1 remains flexible enough to bind with different protofilament registers on each MT. The spindle midzone is subject to many forces during mitosis (55) and PRC1's ability to promote polymerization (Fig. 9C) likely increases the stability of MTs that are critical for positioning the mitotic spindle as well as providing some degree of compliance.

Materials and Methods

Protein Purification. All materials purchased from Sigma-Aldrich unless indicated otherwise. A sequence encoding a tobacco etch virus (TEV) protease site and a hexahistidine tag was fused to the 5' end of PRC1-SC (amino acids 303–620). The overexpression of His₆-tagged PRC1-SC in BL21 (DE3) Rosetta cells was induced by 0.5 mM isopropyl β -D-1-thiogalactopyranoside at 18 °C for approximately 12–16 h. The following steps were done on ice or at 4 °C. We lysed cells in lysis buffer [50 mM phosphate buffer, 20 mM imidazole, 300 mM NaCl, 0.5% Tween-20, 1 mM TCEP, 4 mM Benzamidinium-HCl, 1 mg/mL Lysozyme

supplemented with 1 \times cComplete EDTA-free protease inhibitor (Roche), pH 8.0] by sonication (40% amplitude, six 1-min cycles) and subsequently clarified the lysate by centrifugation at 40,000 rpm in a Ti45 rotor (Beckman Coulter) for 50 min. The supernatant was mixed with Ni-NTA (catalog no. 30250; QIAGEN) at 4 °C for 50 min. The Ni-NTA resin was washed with nickel washing buffer [50 mM phosphate buffer, 20 mM imidazole, 300 mM NaCl, 0.1% Tween-20, 0.5 mM Tris(2-carboxyethyl)phosphine (TCEP), pH 8.0], and then PRC1-SC was eluted with elution buffer (50 mM phosphate buffer, 500 mM imidazole, 100 mM NaCl, 1 mM TCEP, pH 8.0). The fractions containing proteins were pooled, mixed with TEV protease, and dialyzed against 500 mL storage buffer [50 mM phosphate buffer, 300 mM KCl, 20% (wt/vol) sucrose, 10 mM β -mercaptoethanol, pH 8.0] for 12 h. The TEV-digested protein solution was then gel-filtered through Superdex 200 16/60 (catalog no. 28989335; GE Healthcare) that was equilibrated in gel filtration buffer (50 mM Tris, 300 mM KCl, 10 mM β -mercaptoethanol, pH 8.0). Fractions containing PRC1-SC were pooled, mixed with a fivefold volume of low-salt buffer (50 mM Tris, 10 mM β -mercaptoethanol, pH 8.0), and then loaded onto a 1-mL HiTrap SP Sepharose FF column (catalog no. 17-5054-01; GE Life Sciences). Proteins were eluted with a gradient containing 10–100% high-salt buffer (50 mM Tris, 500 mM KCl, 10 mM β -mercaptoethanol, pH 8.0). Fractions containing PRC1-SC were pooled and dialyzed against 1 L storage buffer [50 mM phosphate buffer, 300 mM KCl, 20% (wt/vol) sucrose, 10 mM β -mercaptoethanol, pH 8.0] for 12 h. The purified protein was snap-frozen with liquid nitrogen at a final concentration of 340 μ M and stored at -80° C.

Porcine tubulin (catalog no. T240; Cytoskeleton) was reconstituted to 10 mg/mL in BRB80 buffer [80 mM piperazine-*N,N'*-bis (Pipes) (pH 6.9), 1 mM EGTA, 1 mM MgCl₂, 1 mM DTT] with 10% (vol/vol) glycerol and 1 mM GTP and was flash-frozen in 5- μ L aliquots until needed. Subtilisin (catalog no. P5380; Sigma-Aldrich) was reconstituted to 2 mg/mL in BRB80 buffer with 10% glycerol and was flash-frozen in 5- μ L aliquots until needed.

Cosedimentation Assays. Equilibrium binding between PRC1-SC and MTs was measured using cosedimentation assays. A 15- μ L aliquot of PRC1-SC at 340 μ M was diluted to 40 μ L in cold BRB80 buffer with 10% glycerol and desalted using a 0.5-mL Zeba column (catalog no. P189882; WVR). Aggregates were subsequently removed by cold filtration using a spin filter (catalog no. UFC3 0V V 25; Fisher Scientific) and kept at 4 °C. To make dynamic MTs, two 5- μ L aliquots of tubulin were thawed, combined, and warmed to 37 °C for 15 min. The MTs were

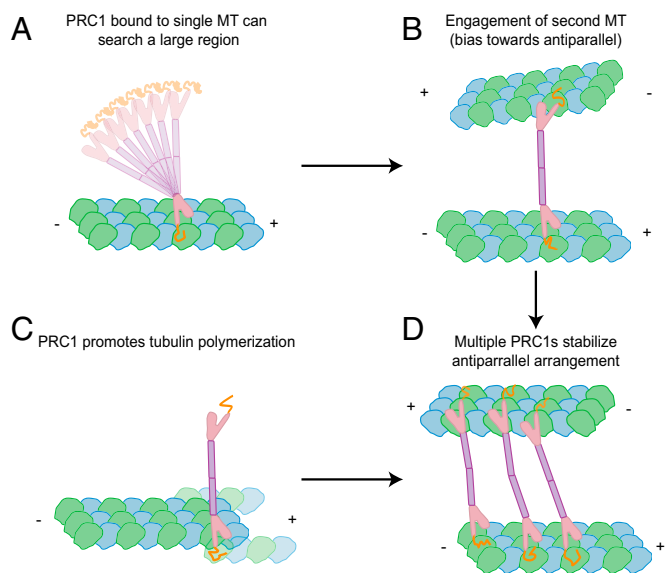


Fig. 9. PRC1 flexibility, selectivity for antiparallel MTs, and ability to promote MT polymerization contribute to the formation of stable arrays. (A) An isolated PRC1 molecule on an MT can flex significantly to search a large region of space to find an antiparallel MT. (B) MTs that are not perfectly antiparallel are biased toward an antiparallel arrangement by PRC1. (C) Tubulin polymerization further stabilizes MT regions decorated with PRC1. (D) Binding of multiple PRC1 molecules requires MTs to be antiparallel, but PRC1 retains enough flexibility to bind across multiple protofilaments. PRC1 is depicted with the spectrin domain in pink, rod and dimerization domains in purple, and the C-terminal domain in orange. The MT is depicted as three protofilaments with β -tubulin in blue and α -tubulin in green.

pelleted by centrifugation at $17,000 \times g$ for 20 min and were resuspended in warm BRB80 buffer with 1 mM GTP and 10% glycerol to a concentration of $\sim 50 \mu\text{M}$. MTs were kept at 37°C at all times to prevent cold-induced depolymerization.

Taxol/paclitaxel (catalog no. TXD01; Cytoskeleton) was reconstituted to 2 mM in anhydrous DMSO and was frozen in $5\text{-}\mu\text{L}$ aliquots until needed. To make taxol-stabilized MTs, one $5\text{-}\mu\text{L}$ tubulin aliquot was thawed and warmed to 37°C , and $0.5 \mu\text{L}$ taxol was added at 10 and 20 min after the tubulin had reached 37°C . MTs were pelleted as above and resuspended in room-temperature BRB80 buffer with $50 \mu\text{M}$ taxol.

MTs treated with subtilisin to remove the C-terminal tails of tubulin were prepared by making dynamic or taxol-stabilized MTs, as above, and adding subtilisin to a final concentration of 0.05 mg/mL . Proteolysis was carried out for 20 min at 37°C . Subtilisin activity was stopped by the addition of PMSF (freshly dissolved in DMSO to 20 mM) to a final concentration of 2 mM in the tubulin proteolysis mixture. Treated MTs were pelleted as above and were resuspended in warm BRB80 buffer with 10% (vol/vol) glycerol and either 1 mM GTP (for dynamic MTs) or $50 \mu\text{M}$ taxol (for taxol-stabilized MTs). The extent of cleavage was assessed to be $62 \pm 2\%$ from the relative intensity of the two tubulin bands clearly separated in the SDS gel following the pelleting assay of the subtilisin-treated taxol-stabilized MTs (Fig. S4). MTs not treated with subtilisin were mock-treated by adding the same volume of subtilisin-free buffer and PMSF at the same time points and also were pelleted a second time.

Protein concentrations for tubulin (after depolymerization by dilution into BRB80 buffer containing 50 mM CaCl_2 for taxol-stabilized MTs or cooling on ice for dynamic MTs) and PRC1-SC were measured using absorbance at 595 nm in the presence of Bradford reagent (catalog no. PI-23236; Fisher Scientific). Absorbance values were converted to concentrations using a calibration curve generated using BSA.

For cosedimentation assays with dynamic MTs, tubulin and PRC1-SC were mixed together in BRB80 buffer with 10% glycerol and 1 mM GTP to final concentrations of $10 \mu\text{M}$ and $5 \mu\text{M}$, respectively. Binding reactions were kept at 37°C at all times. For assays with taxol-stabilized MTs, tubulin and PRC1-SC were mixed together in BRB80 buffer with 10% glycerol and $20 \mu\text{M}$ taxol at $2 \mu\text{M}$ and $10 \mu\text{M}$, respectively, and were kept at 4°C to replicate the temperature used when preparing the EM grids. All binding reactions were carried out in $40\text{-}\mu\text{L}$ volumes and were left for 20 min to reach binding equilibrium. The $40\text{-}\mu\text{L}$ binding reaction was layered on top of a 50% glycerol cushion of the same binding buffer and was spun at $310,000 \times g$ in a TLA-100 rotor (Beckman Coulter) for 10 min. After centrifugation, the top $45 \mu\text{L}$ was taken as the supernatant. The middle $70 \mu\text{L}$ was discarded, and $20 \mu\text{L}$ of the binding buffer was added to the remaining $25 \mu\text{L}$ to resuspend the pellet in a volume equal to the supernatant. Then $15 \mu\text{L}$ of 4 \times LDS sample buffer (catalog no. NP0007; Life Technologies) supplemented with 80 mM DTT was added to each sample and boiled for 5 min. All the supernatant and the pellet were loaded onto Bolt Bis-Tris 10% polyacrylamide gels (catalog no. NW00100BOX; Life Technologies) and run at 150 V. Gels were stained with SYPRO ruby (catalog no. 170-3125; Bio-Rad) or Flamingo (catalog no. 161-0491; Bio-Rad) overnight and were visualized using a Gel Doc EZ imager (Bio-Rad). Exposure times for gel imaging were adjusted to the longest possible time that did not saturate camera pixels for intense bands to ensure accurate quantitation while maximizing sensitivity for weak bands. Degradation products for PRC1-SC were observed, as reported previously (15). Quantification of protein bands was performed using the built-in Analyze/Gels function of ImageJ (56). A rectangle was drawn covering the protein bands for each lane, and the pixel intensities were integrated within the rectangle for each band to measure the total amount of protein. The raw integrated intensities were used to calculate the fraction of protein in supernatant and pellet fractions for each gel.

Cryo-EM Sample Preparation and Imaging. A $10\text{-}\mu\text{L}$ aliquot of PRC1-SC protein was diluted to $50 \mu\text{L}$, desalted, and filtered as above, except that glycerol was omitted. A $5\text{-}\mu\text{L}$ aliquot of porcine tubulin was warmed to 37°C and allowed to polymerize for 15 min. MTs were pelleted by centrifugation at $17,000 \times g$ for 20 min and were resuspended in warm EM buffer with $66 \mu\text{M}$ peloruside A (supplied by John H. Miller and Peter Northcote, Centre for Biodiscovery, Victoria University of Wellington, Wellington, New Zealand).

MTs were diluted to 0.4 mg/mL , and $2 \mu\text{L}$ of the solution was applied to a glow-discharged C-flat grid with $1.2\text{-}\mu\text{m}$ holes (Protochips). The Mark IV Vitrobot (FEI) used for sample vitrification was set to 4°C and 100% relative humidity. Maintaining PRC1-SC at low temperature during all experimental procedures minimized protein aggregation and increased occupancy on the MT lattice. MTs were allowed to adhere to the grid for 30 s and then were washed twice with $4 \mu\text{L}$ of the cold PRC1-SC solution. The grid then was blotted for 4 s with a blot force of 20 Vitrobot units and was plunged into ethane slush.

Images were collected on a Titan electron microscope operated at 300 kV and equipped with a Gatan K2 direct detector (Gatan) using the Legion automated data-collection pipeline (57). The micrographs have a nominal magnification of $27,500\times$, resulting in a final pixel size of 1.32 \AA per pixel. Twenty frames of 300 ms each were collected at a dose rate of 8 e^- per pixel per second, with a total dose of $28 \text{ e}^-/\text{\AA}^2$.

Image Analysis and Data Processing. Images were processed using the Appion pipeline (58), including individual frame alignment using MotionCorr (59) and contrast transfer function estimation using CTFIND4 (60). Following previous MT reconstruction methods (27), regions of the raw micrographs containing decorated MTs were extracted using overlapping $675\text{-}\text{\AA}^2$ boxes spaced 80 \AA apart. Each of these boxes was treated as an independent, single particle using iterative helical real space reconstruction (IHRSR) (61). The boxed MT segments were sorted by protofilament number, and initial 3D alignment parameters were generated using EMAN2 multimodel refinement (62) with models of 12-, 13-, 14-, and 15-prot filament MTs (63) low-pass filtered to $20\text{-}\text{\AA}$ resolution, followed by IHRSR using a resolution range of $40\text{-}10 \text{ \AA}$ to obtain initial 3D reconstructions. FREALIGN (64) was then used to process the 13- and 14-prot filament MTs further to obtain better alignment parameters. Finally, we applied in-house scripts to verify and enforce the seam location (i.e., to align α - and β -tubulin and thus PRC1-SC) for particles from the same MT (28). The overall resolution of the resulting map was not significantly better after this last step, but the protein density for PRC1-SC was notably improved. Local resolution estimates were performed using the Bsoft software package (65) using the whole MT. Half-maps were generated using MT segments separated by MT, rather than even/odd images, to ensure that each half-map did not contain segments from the same MT. The reconstruction was B-factor sharpened (-125 \AA^{-1}) and low-pass filtered to the estimated overall resolution (4 \AA). 2D class averages were obtained by applying the alignment parameters from the 3D reconstruction to the overlapping boxes and averaging them to generate "superparticles" that represent the average of eight boxes and have improved signal. The superparticles were binned by 2 and were subjected to iterative (multivariate statistical analysis/multi-reference alignment) using IMAGIC (66) to generate the final class averages shown in Fig. 1D and Fig. S1B.

Previous crystallographic structures of PRC1-S [Protein Data Bank (PDB) ID code 3NRX] (15) and of the dimerization, rod, and spectrin domains of PRC1 (PRC1NS Δ C, PDB ID code 4L6Y) (26) were used as starting points for our model. Neither structure included the full loop between PRC1 S-H7 and S-H8 (all secondary structures follow the nomenclature established in ref. 26). Initial atomic models of PRC1-SC were generated using the iTASSER server (67), focusing on building this missing loop. Coarse refinement of the predicted structures was performed using iMODFIT (68) to fit the three best starting models from the iTASSER server into the 14-prot filament electron density map. The best model (as judged by agreement with the density) was then refined further using Rosetta (69) to optimize the fit and stereochemistry of the atomic model. The Fourier shell correlation for a simulated map based on this model and the final reconstruction (Fig. 1 and Fig. S1C) matches the average local resolution estimates for the PRC1 density and tubulin.

PRC1-SC occupancy was estimated following previously described procedures (27, 28), by comparing the final reconstruction with a simulated map that had 100% PRC1-SC occupancy. The simulated map was built by docking heterodimers of tubulin with the bound PRC1-S into the final reconstruction using University of California, San Francisco Chimera (70) for a single protofilament. The molmap command was used to generate electron density for a single protofilament. This density was symmetrized using the same scripts used to generate a whole microtubule during the density refinement. The Fourier transforms for the simulated map and for the final reconstruction were computed using EMAN2 libraries, and the ratio of the amplitudes of the line at the $80\text{-}\text{\AA}$ layer (and its four higher-order repeats up to $10\text{-}\text{\AA}$ resolution) on the simulated map and the final reconstruction was used to determine the PRC1-SC occupancy.

MD Simulations.

System preparation. The starting structures were prepared from the X-ray crystal structure of the PRC1 protein (PDB ID code 4L6Y) (26) using the Protein Preparation Wizard implemented in Maestro, version 9.8 (Schrodinger). This procedure included the modeling of missing side chains and loops using Prime (71). The system was prepared for MD simulations using the GROMACS simulation package version 4.6.3 (72). The Amber99sb-ILDN force field (73) was used for the protein, and the prepared structure was placed in triclinic boxes of explicit TIP3P water molecules (74) with a minimum distance of 20 \AA between the protein surface and the border of the box. Some water molecules were replaced with Mg^{2+} ions to neutralize the systems. Additional Mg^{2+} and Cl^- ions were added to reproduce approximately the experimental ion concentration of 1 mM.

Preparatory simulations. The starting structures were energy minimized, and subsequently the solvent was equilibrated in three phases. For the first phase of equilibration, a canonical/NVT (number of particles, volume, and temperature are kept constant) ensemble was applied for 5 ns. Position restraints were applied to all protein atoms using a harmonic potential with a force constant of $1,000 \text{ kJ}\cdot\text{mol}^{-1}\cdot\text{nm}^{-2}$. Following NVT, two phases of isothermal-isobaric/NPT (number of particles, pressure, and temperature are kept constant) equilibration were applied for 5 ns each. In the first phase all protein atoms were restrained; in the second phase the position restraint was limited to backbone atoms only. The pressure of the simulation box was kept at an average of the barometric pressure at sea level (1 bar) using the isotropic Berendsen barostat (75) with a time constant of 1 ps and a compressibility of 4.5×10^{-5} /bar. During the equilibration a 2-fs integrations time-step was used, and the neighbor lists were updated every 10th time-step. Short-range nonbonded van der Waals (Lennard–Jones) and Coulombic interactions were calculated within a cutoff radius of 1.15 nm. The particle mesh Ewald method (76) was used for long-range electrostatic calculations. The long-range Lennard–Jones interactions were analytically corrected for the pressure and the energy calculations. The solvent and the protein were coupled separately to an external heat bath at 300 K with the velocity-rescaling thermostat (77) using a time constant of 0.5 ps. Water molecules were constrained using the SETTLE algorithm (78), and the covalent bonds in the proteins were constrained using the LINCS algorithm (79). Boundaries were treated periodically.

MD simulations. Production MD simulations were carried out for 300 ns using the NPT ensemble in the absence of any restraints. Two replicate simulations were conducted, initiated using different random starting velocities. The isotropic Parrinello–Rahman barostat (80, 81) was used to keep the average pressure at 1 bar with a time constant of 1 ps. All other simulation parameters were the same as the equilibrium simulations. The trajectories were sampled every 80 ps for

analysis. In all cases, the simulations reached a steady state, and the integrity of the structure was well maintained over the long-simulation times. Superposition of PRC1 structures at different time points was performed using THESEUS (82).

Comparisons with Other MT-Binding Domains. Previous atomic models of motor domains bound to MTs derived from cryo-EM reconstructions (43, 45–47) and crystallography (41) were superimposed on our final atomic model using the H11' helix on α -tubulin to align the binding pockets locally. The following structures were inspected to identify tubulin residues that potentially share binding activity for PRC1, kinesin, or dynein: 2P4N (42), 4HNA (41), 4CK5 (44), 4UXO (45), 3J8Y (46), 3J6P (47), 3J1U, and 3J1T (43). A cutoff distance of 4 Å was used to identify potentially interacting residues.

Accession Codes. The atomic coordinates have been deposited in the PDB under the accession code 5KMG. The cryo-EM reconstruction has been deposited in the Electron Microscopy Data Bank under the accession code EMD-8266.

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