Eukaryotic chaperonin CCT stabilizes actin and tubulin folding intermediates in open quasi-native conformations

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Three-dimensional reconstruction from cryoelectron micrographs of the eukaryotic cytosolic chaperonin CCT complexed to tubulin shows that CCT interacts with tubulin (both the α and β isoforms) using five specific CCT subunits. The CCT–tubulin interaction has a different geometry to the CCT–actin interaction, and a mixture of shared and unique CCT subunits is used in binding the two substrates. Docking of the atomic structures of both actin and tubulin to their CCT-bound conformation suggests a common mode of chaperonin–substrate interaction. CCT stabilizes quasi-native structures in both proteins that are open through their domain-connecting hinge regions, suggesting a novel mechanism and function of CCT in assisted protein folding.

Keywords: actin/chaperonin/electron microscopy/protein folding/tubulin

Introduction

Folding of many proteins in vivo requires interaction with macromolecular complexes known as chaperonins. These proteins are ubiquitous oligomeric assemblies that have been classified into two distinct families that share limited but significant sequence homology: type I, present in eubacteria and endosymbiotic organelles, and of which the bacterial GroEL is the best known representative; and type II, present in archaebacteria and the eukaryotic cytosol, which are represented by the thermosome and CCT (chaperonin containing TCP-1), respectively (Bukau and Horwich, 1998; Gutsche et al., 1999; Willison et al., 1999). Most of the chaperonins share a common architecture, a cylinder made up of two back-to-back stacked rings, each one enclosing a cavity where folding takes place. The atomic structures of GroEL (Braig et al., 1994) and the type II thermosome (Ditzel et al., 1998) have revealed a common subunit architecture consisting of three domains: apical, intermediate and equatorial. The equatorial domain provides most of the intra- and inter-ring interactions and contains the binding site for ATP, the hydrolysis of which is necessary for the working cycle of the chaperonin, while the apical domain is involved in substrate binding and undergoes large conformational changes during the folding cycle. There are, however, numerous differences between type I and type II chaperonins, one of which is the absence of co-chaperonins for type II family members, whose role in the closure of the cavity during the chaperonin working cycle is fulfilled instead by a helical protrusion in the apical domain (Klumpp et al., 1997; Ditzel et al., 1998; Llorca et al., 1999a). Another important difference is related to the degree of complexity of the chaperonin ring, ranging from the seven identical subunits of type I chaperonin GroEL to eight different polypeptide subunits in the case of the type II chaperonin CCT. The most important difference between these two chaperonins is, however, related to their substrate specificity: whereas GroEL interacts with a broad range of substrates (Houry et al., 1999) using a non-specific recognition mechanism based on hydrophobic interactions (Bukau and Horwich, 1998; Chen and Sigler, 1999; Shiltelman et al., 1999), the main in vivo substrates of CCT are actins and tubulins (although other proteins that bind to CCT are continuously being found, suggesting a possible broader role of CCT in protein folding; Leroux and Hartl, 2000). CCT has already been shown to bind actin through a mechanism that is both geometry dependent and subunit specific (CCTα, CCTβ and CCTε subunits are involved in actin binding; Llorca et al., 1999b). To gain further insight into the folding mechanism of CCT and to search for a common pattern of interaction with tubulin, the other major in vivo substrate of CCT, we have carried out electron microscopy and biochemical analysis of CCT–tubulin complexes. From docking analyses performed on actin and tubulin folding intermediates bound to CCT, we propose a mechanism of interaction of CCT with folding substrates that is different from the mechanism proposed for GroEL.

Results and discussion

CCT–tubulin interaction

When either recombinant β-tubulin or a mixture of α- and β-tubulin purified from microtubules (tubulin unless stated otherwise) is chemically denatured and incubated with CCT in a diluting buffer, a binary complex is formed that can be visualized by negative staining or cryoelectron microscopy (NS and CR in Figure 1A, respectively). These top views show the characteristic circular shape of CCT with some material crossing the cavity. The two-dimensional average of the stained particles shows that tubulin (β-tubulin in Figure 1B; tubulin not shown but identical to Figure 1B) is an asymmetrical mass that crosses the cavity and contacts two regions of CCT in an apparent 1,5 interaction, clearly distinct from the 1,4 interaction found for actin (Llorca et al., 1999b). A more detailed view of this interaction is observed in a
three-dimensional reconstruction of negatively stained CCT-β-tubulin complexes carried out by the random conical tilt method (Frank, 1996) (Figure 1C). Due to the fact that the staining agent cannot contrast the whole height of the CCT particle, only the apical domains of the ring in contact with the grid are reconstructed. The three-dimensional reconstruction reveals the basic features already observed in the two-dimensional average; tubulin is divided into two asymmetrical domains connected by a small linker, which interact with two diametrically opposite regions of CCT. This three-dimensional reconstruction shows that the tubulin interaction with apparently only two CCT subunits observed in the two-dimensional average (Figure 1B) is a simplification of a more complex interaction involving several subunits (see below).

**Three-dimensional structure of the CCT-tubulin complex**

CCT–tubulin complexes were observed by cryoelectron microscopy, a technique that allows reconstruction of the whole particle. Chemically denatured tubulin was reacted with CCT for 15 min in a diluting buffer and aliquots of this solution were vitrified and observed at low temperature (−170°C) under an electron microscope. 5937 tilted top views were processed. Using neural network methods that searched for variability inside the chaperonin cavity, 2007 substrate-free and 3930 substrate-bound particles were classified. Each group was used for independent three-dimensional reconstructions. The substrate-free CCT three-dimensional structure (Figure 2A and B) reveals a similar architecture, albeit at higher resolution, to that of a previous three-dimensional reconstruction of CCT (Llorca et al., 1999a). The differences observed are due to the low number of particles analysed in the former reconstruction (Llorca et al., 1999a) and the better quality of the electron microscopy performed in this study.

The three-dimensional reconstruction of the CCT–tubulin complex (Figure 2C–H) reveals a barrel-shaped structure with a tubulin molecule bound to the apical domains of one of the rings. As in the case of the CCT–α-actin complex (Llorca et al., 1999b), only one molecule of tubulin is bound to CCT, despite the 10-fold excess of unfolded tubulin incubated with CCT. This strengthens the notion that inter-ring communication occurs upon substrate binding that prevents interaction of a second polypeptide with the opposite ring. The shape of the bound tubulin was found to be reproducible after several independent reconstructions and is almost identical to that obtained using random conical tilt of negatively stained CCT–β-tubulin complexes (compare the views shown in Figures 1C and 2H). Tubulin binds to CCT following a 1.5 geometrical interaction and shows two
domains connected by a small linker. One of these domains (right side in Figure 2G and H) has the shape of a truncated cone and seems to interact with two CCT subunits. The other domain, which looks smaller in the two-dimensional projection average (left tubulin domain in Figure 1B) or the two three-dimensional reconstructions (left tubulin domains in Figures 1C and 2H), in fact contains a similar mass to the apparently larger domain (compare the two tubulin domains in Figure 2D). The left tubulin domain seems to interact with three CCT subunits. The conformation of all eight apical domains in the tubulin-bound ring of CCT (Figure 2G and H) is more closed than that obtained for the substrate-free CCT (Figure 2A and B) and for the CCT-α-actin complex (Llorca et al., 1999b), and may be a consequence of a stronger interaction between tubulin and the five CCT subunits than the one generated between actin and just two CCT subunits (Llorca et al., 1999b). The differences found are again strongly indicative of CCT displaying a considerable degree of flexibility, as observed for other type II chaperonins (Ditzel et al., 1998; Nitsch et al., 1998; Llorca et al., 1999a,b; Grantham et al., 2000). The structure of the substrate-free ring in the CCT–tubulin complex (Figure 2E and F) clearly displays a clockwise twist in the apical domains and begins to reveal the helical protrusions pointing towards the adjacent subunit, undetected in our previous lower resolution reconstruction (Llorca et al., 1999a).

**Comparison between the three-dimensional reconstructions of apo-CCT, CCT–α-actin and CCT–tubulin**

A visual inspection of the structures of apo-CCT and CCT–α-actin (Llorca et al., 1999b) reveals a very similar conformation of the chaperonin. Both structures have their apical domains in an open conformation characteristic of an open state with their helical extensions pointing upwards. This state is similar to one observed for the thermosome (see Figures 1 and 2 of Gutsche et al., 1999). However, the conformation of the CCT–tubulin complex is somewhat different, with the apical domains being in a more closed conformation. In order to analyse the structural changes driven by tubulin binding, a difference map was calculated following the alignment of the apo-CCT and CCT–tubulin structures. The most relevant features of the difference map between the apo-CCT minus the CCT–tubulin structure are shown in red shading in Figure 3A. The differences are mainly localized to the tips of the apical domains, which point upwards in the substrate-free CCT structure. The differences between the CCT–tubulin minus the apo-CCT structure (Figure 3B) are due mainly to the presence of tubulin in the cavity and an extra mass in the region of the apical domains surrounding the substrate. Although some changes can also be found in the equatorial domains, it seems clear that upon tubulin binding CCT modifies the conformation of the tips of its apical domains to generate a semi-closed conformation.

The statistical significance of the tubulin mass observed within the cavity was tested by a Student’s t-test carried out between the cavities of the tubulin-bound and substrate-free three-dimensional reconstructions. Figure 3C shows the volume where no significant differences are found between the CCT–tubulin and substrate-free CCT cavities (blue shading). On the other hand, the red shading in Figure 3D indicates the volume where significant differences (P <0.0005) in the cavity are found between the two three-dimensional reconstructions. This mass corresponds to the location of tubulin. From these results it seems that tubulin (but not actin) binding to CCT induces a downward movement of the tips of the apical domains that is in fact maintained in the following steps of the CCT cycle (O.Llorca, J.Martín-Benito, G.Hynes, K.R.Willison, J.L.Carrascosa and J.M.Valpuesta, manuscript in preparation).

Another important difference between the three-dimensional reconstructions of CCT–α-actin and CCT–tubulin concerns the location of the substrate binding sites on CCT. In the structure of the CCT–α-actin complex, actin seems to be binding below the helical protrusions, but in the CCT–tubulin structure tubulin seems to interact with CCT over a much wider area encompassing the base of the apical domains and parts of the helical protrusions. This may reflect differences in the structures of the two proteins when bound to CCT. Additionally, there may be conformational differences due to the sequence heterogeneity among CCT subunits, which mostly reside in the apical domains. Although a direct comparison between CCT and GroEL points to the region at the base of the helical protrusions as being the one involved in substrate binding (Willison, 1999), the fact that it contains a high proportion of charged residues has led some authors
(Klumpp et al., 1997) to propose that the hydrophobic region of the helical protrusions is a novel substrate binding site for type II chaperonins. From our results with actin and tubulin it seems that one or both regions of the CCT apical domains could play a role in substrate interaction. A recent study (Bosch et al., 2000) describes structural plasticity within the helical protrusions of thermosome apical domains. These authors hypothesize that differences in amino acid sequence of the CCT helical protrusions could give rise to different conformations in this region of the chaperonin and they predict that, due to the presence of a proline residue in the helical protrusions of the CCTβ and CCTε subunits, they should not be able to adopt a helical conformation. Curiously, these two subunits are involved in actin binding (Llorca et al., 1999b).

The structural studies presented here and in Llorca et al. (1999b) show that tubulin binds to more CCT subunits than actin to generate a more complex set of interactions. It therefore seems reasonable to suggest that the different conformational changes being induced in the CCT structure (mainly at the tips of the apical domains) upon binding of either actin or tubulin are to do not only with differences in the regions of CCT interacting with the two substrates but also with the inherently different conformations of the CCT subunits interacting with these two cytoskeletal proteins.

**Docking analyses of actin and tubulin**

Even at the low resolution of this study the structures of α-actin and tubulin obtained in the three-dimensional reconstructions of these two proteins complexed with CCT (Llorca et al., 1999b and this work, respectively) are very suggestive of a mostly folded but open conformation in which two domains of these two proteins interact with opposite sides of CCT. The native structure of actin (Kabsch et al., 1990) can be divided into two equally sized domains, the so-called small domain (red in Figure 4A) and the large domain (yellow and white in Figure 4A). The nucleotide sits in the cleft between the two domains (blue in Figure 4A). The two domain structure of tubulin (Nogales et al., 1998a) is less obvious, but a bilobar structure is visible upon inspection of the native structure. In all tubulins (Nogales et al., 1998b) there is an N-terminal domain that contains the GTP binding site (red in Figure 4A) and a C-terminal domain of similar size (green, white and yellow in Figure 4A). To test the hypothesis that open conformations of both proteins
**CCT subunits involved in tubulin binding**

In order to validate the model of tubulin binding to CCT and also to determine which CCT subunits are involved in the interaction, several immunomicroscopy experiments were carried out. In the first set of assays, immunocomplexes of CCT–tubulin and each of four monoclonal antibodies against specific subunits of CCT ($\alpha$, $\delta$, $\eta$, and $\theta$) were negatively stained, recorded, processed and averaged. In each antibody, labelling two different conformations that follow the same I,5 geometry can be observed (Figure 5A and B; only the average images corresponding to immunocomplexes generated with the anti-CCT$\theta$ antibody are shown; however, the same results were obtained with antibodies against CCT$\alpha$, CCT$\eta$ and CCT$\theta$). From these results and the published model of CCT subunit arrangement (Liou and Willison, 1997) we can hypothesize two modes of interaction (see model in Figure 5I). One involves CCT$\beta$ interaction with one tubulin domain and CCT$\eta$ interaction with the other (Figure 5A). The second possible interaction involves CCT$\epsilon$ with one tubulin domain and CCT$\theta$ with the other (Figure 5B). These two modes of interaction could reflect two ways for tubulin to interact with CCT, as observed for CCT–actin (Llorca et al., 1999b), or they could be related to the presence of two types of tubulin ($\alpha$ and $\beta$) that interact in different ways. To test this hypothesis, similar immunolabelling experiments were carried out with complexes of CCT and recombinant $\beta$-tubulin, and the same two types of interactions were observed (results not shown), indicating clearly that both $\alpha$- and $\beta$-tubulin share the same interaction pattern with the CCT subunits.

**Tubulin domains involved in CCT binding**

To test the validity of the tubulin interaction suggested by the docking results, the $\beta$-tubulin C-terminal domain was divided into three fragments that, according to the model (Figure 4C and E), should interact with different CCT subunits. Three chimeric proteins of these fragments fused with Ha-Ras (but not Ha-Ras itself) bind to CCT upon *in vitro* translation in reticulocyte lysate (see Figure 6A and B). The first chimeric protein, Ha-Ras-$\beta$-tubulin fragment 5, contains residues P263–I384 of $\beta$-tubulin (white in Figure 4C and E) and binds strongly to CCT (Figure 6B). Immunolabelling experiments were performed with CCT–Ha-Ras–$\beta$-tubulin fragment 5 complexes and antibodies against the CCT$\eta$ or CCT$\theta$ subunits to determine which CCT subunits are involved in the interaction (Figure 5C and D; only the average images of immunocomplexes generated with antibody against CCT$\theta$ are shown; however, the same results were obtained with antibodies against CCT$\eta$). The results not only confirmed the interaction between Ha-Ras–$\beta$-tubulin fragment 5 and CCT but also showed that it binds to CCT$\beta$ (Figure 5C) and CCT$\epsilon$ (Figure 5D). A second chimeric protein contains a fragment of $\beta$-tubulin that encompasses residues D205–L265 (Ha-Ras–$\beta$-tubulin fragment 3; green in Figure 4C). Immunolabelling experiments performed with antibodies against CCT$\beta$ (Figure 5E and F) showed that Ha-Ras–$\beta$-tubulin fragment 3, which according to the docking model (Figure 4C) binds to the first subunit clockwise to CCT$\beta$ or CCT$\epsilon$, does indeed bind to CCT$\gamma$ (Figure 5E) and CCT$\eta$ (Figure 5F), respectively (see model in Figure 5I). Finally, immunolabelling
experiments performed with antibodies against CCTδ (Figure 5G and H) and the third chimeric protein (Ha-Ras–β-tubulin fragment 4), containing residues I384–A455 of β-tubulin (yellow in Figure 4C), showed that Ha-Ras–β-tubulin fragment 4 binds, as suggested by the docking model (Figure 4C), to the second subunit clockwise to CCTβ (CCTδ; Figure 5G) and CCTe (CCTβ; Figure 5H). The last two hybrid proteins interact more weakly with CCT than fragment 5 in reticulocyte lysate (Figure 6A) in equilibrium between an unbound population and a CCT-bound species, which is very similar to the behaviour of the Ha-Ras–β-actin, subdomain4 fusion protein when interacting with CCT (Llorca et al., 1999b). The immunolabeling results are therefore consistent with the docking model for tubulin (Figure 4C and E) and this enables us to propose that five specific CCT subunits (using two modes of interaction) are involved in the binding of tubulin to the open conformation of CCT (Figure 5I). This represents the first step in the functional cycle of CCT.

**Biochemical correlates of the CCT–tubulin interaction**

We have shown recently that certain interactions between β-actin folding intermediates and specific CCT subunits can be maintained during immunoprecipitation experiments using mixed micelle detergent buffers that cause complete disruption of CCT into its constituent monomers (Hynes and Willison, 2000). In this assay the two strongest interactions occurred between β-actin and CCTβ or CCTe. The fusion protein between Ha-Ras and subdomain 4 of actin also interacted most strongly with CCTβ and CCTe. This biochemical analysis is consistent with the two three-dimensional reconstructions of the CCT–α-actin complexes (Llorca et al., 1999b) and the docking analysis carried out here. Figure 6 shows the results of a similar analysis carried out with the tubulins. Complexes between CCT and [35S]α-tubulin or [35S]β-tubulin were prepared under the approximate in vivo conditions of in vitro translation in reticulocyte lysate. These complexes were immunoprecipitated with each of the eight anti-CCT subunit antibodies and the relative recoveries of 35S-labelled tubulins with individual CCT subunits were measured by quantitation of SDS–PAGE signals (Figure 6C–E). Both tubulin isoforms were recovered most effectively with the anti-CCTβ and anti-CCTe antibodies (Figure 6D and E). This data is remarkably consistent with the structural data in which CCTβ and CCTe both interact with fragment 5 of tubulin (white in Figure 4A, C and E). Of all the tubulin subdomains tested,
fragment 5 targets Ha-Ras most strongly to CCT upon in vitro translation (Figure 6A and B). Based on these results and those provided by immunomicroscopy (Figure 5A–H), a model seems to emerge in which CCT shares strong interactions with actin (Llorca et al., 1999b) and tubulin (this work) through either its CCTβ or CCTε subunits. This allows us to suggest that these two subunits are the ones driving the high affinity interaction between CCT and the two cytoskeletal proteins and provides a simple explanation for the ability of actin and tubulin to compete for binding to CCT, as observed by Melki et al. (1993).

The role of CCT in actin and tubulin folding
A common pattern for the interaction between CCT and the two cytoskeletal proteins is revealed: both proteins have two topological domains separated by a hinge with the nucleotide placed in its vicinity (Figure 4A). During the folding process, the two proteins reach (by themselves or with the help of other cofactors) quasi-native conformations that cannot proceed to their final native structures without the assistance of CCT (Willson, 1999). The eukaryotic cytosolic chaperonin may have evolved to overcome a specific kinetic barrier in the folding of actin and tubulin by stabilizing an open conformation through the binding of their two topological domains by opposite sides of the chaperonin ring. In the case of actin the critical step in the folding process may have to do with correct folding of the nucleotide binding site. Unfolding experiments with actin have shown that its native conformation depends on the degree to which the nucleotide contributes to the connectivity between the two domains of the protein (Schüler et al., 2000), and the fact that most of the inter-domain bonds occur through the ATP and cation binding sites strengthens our idea that CCT is involved in actin nucleotide binding or loading. In the case of tubulin a large percentage of the bonds that connect the two domains also occur through the GTP binding site, and some results from biochemical studies carried out with CCT and tubulin clearly indicate that at least one function of GTP binding is to stabilize the quasi-native tubulin molecule during CCT-facilitated folding (Tian et al., 1995). Nevertheless, it cannot be ruled out at this stage that the requirement for a CCT-bound open conformation may be due to other folding limiting processes.

Co-evolution of CCT with actin and tubulin
Phylogenetic analysis of the CCT gene family (Kubota et al., 1994, 1995) suggested that CCT evolved to completion, having eight subunit species, during the early evolution of eukaryotes and we have argued that the modern actin and tubulin proteins and CCT may have co-evolved (Willison and Horwich, 1996; Willison, 1999). These new structures of CCT–tubulin complexes support this view, since tubulin binds to five CCT subunits in two arrangements, thus utilizing all eight CCT subunits. Apparently, actin interacts more simply with CCT, using only three CCT subunits for its two modes of initial interaction with CCT (Llorca et al., 1999b). There also appear to be differences in the location of the binding sites of these two substrates on CCT subunit apical domains, as explained above. Actin binds low down on the apical domains of the CCT subunits, well below the helical protrusions (Klupp et al., 1997), whereas tubulin seems to interact directly with the helical protrusions. The helical protrusions have been suggested to be the location of type II chaperonin substrate binding sites (Klupp et al., 1997; Bosch et al., 2000). Tubulin binding also seems to create a significantly different conformation in the CCT apical domains to that observed for actin.

Willison and Grantham (2000) have proposed that, since many archaean species appear to lack the actin/hexokinase/dnaK fold, the primordial actin fold may have entered the evolving eukaryote by horizontal gene transfer from a Gram-negative bacterium. Thus it seems possible that actin-like molecules first interacted with CCT after it had already evolved to fold tubulins. If an evolving actin protein essentially adopted an already evolving CCT–tubulin folding machinery it is perhaps understandable that the mechanism of CCT binding and folding of actin could be different to that of tubulins. Some support for this idea emerges from recent phylogenetic analysis of a large CCT gene set that includes CCT genes from amitochondrial protist lineages thought to have diverged early from other eukaryotes (Archibald et al., 2000). The data suggest that the CCTβ/ε clade is the latest evolving pair of CCT subunit genes in the stepwise evolution of the CCT gene family (Archibald et al., 2000). Since both of these two ‘newer’ subunits interact with actin, CCTβ also being the common subunit used for both geometrical binding arrangements between actin and CCT (Llorca et al., 1999b), it is possible that the CCTβ/CCTε gene duplication occurred during establishment of the actin folding mechanism. Because modern actin is smaller than tubulin, 375 versus 450 amino acid residues, it may have been a requirement to establish specific actin binding sites in the shorter 1.4 configuration compared with the 1.5 configuration, across the diameter of the CCT cavity, required by tubulins. Indeed, the absolute dimensions of the actin and tubulin systems are presumably a direct function of the geometric constraints introduced by the dimensions of the CCT folding chamber.

During its earliest stages of evolution a primitive CCT machinery that had not yet evolved to complete heterooligomerism might have provided beneficial effects firstly for the evolving tubulins. Then CCT evolved further to cope with actin folding to finally reach the modern arrangement of eight CCT subunit species per ring, each having different functions in substrate binding and processing. The entire system then became fixed and the actins and tubulins hardly changed in sequence from then on (Willison, 1999). We must now establish the biological contributions of CCT to the biophysical properties of the tubulin and actin polymer systems. Does CCT merely increase folding efficiency and prevent aggregation of monomers or, more likely, has CCT enabled a qualitatively new folding process that produces actins and tubulins that have unique biophysical properties compared with their precursors? These are exciting questions now to pursue.

Materials and methods
Antibodies
The anti-CCTβ 8g and CCTε 91A monoclonal antibodies were prepared as described by Llorca et al. (1999b). The anti-CCTβ 81a and anti-CCT9 PK-5h antibodies were prepared by immunizing rats with recombinant Escherichia coli produced apical domain fragments of subunits CCTβ.
Preparation of complexes
Recombinant β-tubulin was prepared by insertion of human β-tubulin cDNA into pET21a vector (Novagen) and expression in E. coli BL21 (DE3) competent cells (Stratagene). Either β-tubulin or bovine brain tubulin (from Cytoskeleton) were denatured in 7 M guanidine chloride and incubated in diluting buffer (100-fold) containing 0.4 M murine CCT purified as described by Liou and Willison (1997). The Ha-Ras-β-tubulin fragments 1–5 fusion proteins were constructed by linking residues M1–L168 of human Ha-Ras to residues M1–D205, D205–I384, D205–L265, I384–A455 and P263–I384 of human β-tubulin, respectively. Residue numbers correspond to the aligned sequences of α- and β-tubulin as described by Nogales et al. (1998a). The fusion proteins were inserted into a Bluescript SKII+ vector (Stratagene) for *in vitro* translation analysis and into a pET11b vector (Stratagene) for expression in E. coli BL21 (DE3) competent cells (Stratagene). Complexes of CCT–tubulin and CCT–chimeric proteins were prepared by incubation for 15 min at room temperature at a 1:10 molar ratio. Immunoblotting was performed as described previously (Llorca et al., 1999b).

Immunoprecipitation of α-tubulin–CCT and β-tubulin–CCT complexes
*In vitro* transcription/translation reactions of full-length human α- and β-tubulin cDNAs (Bluescript SKII+ vector; Stratagene) were performed at 30°C for 30 min in TNT rabbit reticulocyte lysate (Promega) in the presence of 40 μCi (40 pmol) of [35S]-methionine (Amersham) per 50 μl volume. Following sucrose gradient fractionation of the rabbit reticulocyte lysate, the [35S]-labelled α- and β-tubulin produced by *in vitro* translation were recovered by immunoprecipitation with anti-CCT subunit antibodies as described by Hynes and Willison (2000). [35S]-labelled protein bands were visualized by autoradiography and quantitated by PhosphorImager analysis on a Storm 860 (Molecular Dynamics).

Electron microscopy
Aliquots of the CCT–tubulin complexes or the immunocomplexes were applied to carbon grids, negatively stained with 2% uranyl acetate and recorded at 60°C. For the three-dimensional reconstruction of negatively stained specimens using the random conical tilt method (see below), pairs of untilted and tilted (45°) images were taken. For cryoelectron microscopy, samples were applied to the grids for 1 min, blotted for 5 s and frozen quickly in liquid ethane at −180°C. Images were recorded at 20° tilt in a Joell 1200EX-II electron microscope equipped with a Gatan cold stage operated at 120 kV and recorded on Kodak SO-163 film at a 60 000× nominal magnification and ±1.5 μm underfocus.

Image processing and three-dimensional reconstruction
In both the negative stain and frozen hydrated studies, top views were selected and two-dimensional processing was carried out as described by Llorca et al. (1999a). A self-organizing map algorithm (Marabini and Carazo, 1994) was used to separate the particles into those having no substrate in the cavity of the particle and those having density in the cavity to carry out two independent three-dimensional reconstructions using angular refinement algorithms provided by SPIDER (Frank et al., 1996). The volumes were generated using ART with blobs (Algebraic Reconstruction Techniques; Marabini et al., 1998). Eight-fold symmetry was applied to the CCT volume except in the inner cylinder comprising the CCT cavity where the substrate is located, as described previously (Llorca et al., 1999b). The final resolution was calculated by Fourier ring correlation of two independent reconstructions and the value obtained (27 Å for apo-CCT and 25 Å for the CCT–tubulin complex) was used to low pass filter the volume. The three-dimensional reconstruction generated with negatively stained particles of CCT–β-tubulin complexes (417 particles) was carried out using the random conical tilt method (Frank, 1996), and the subsequent and refined volumes using the angular refinement method described above.

Docking of the cryoelectron microscopy structures of α-actin and tubulin bound to CCT with their atomic counterparts
Docking was performed using SITUS (Wrighers et al., 1999). The volume corresponding to actin or tubulin was extracted from their cryoelectron microscopy three-dimensional reconstructions and quantified using a self-organizing algorithm that reduces the volume to a small number of codebook vectors. The atomic structures of α-actin and tubulin were cut into two domains across their hinge regions as described in the text and independently quantified using in total the same number of codebook vectors as for the cryoelectron microscopy volume. The two domains for both proteins were docked independently into the quantitated low resolution data. Among the several docking solutions generated by the docking program, the criteria to select the best fit were based on the root mean square deviation between the codebook vectors from the high resolution X-ray data and low resolution electron microscopy data and the proximity of the cut ends of the two domains. The best docking solutions were merged in a unique pdb file that was subjected to a conjugate gradient energy minimization using CNS (Brünger et al., 1998) in order to relax the molecule. Visualization of the dockings was carried out using VMD (Humphrey et al., 1996).

Difference maps between the apo-CCT and CCT–tubulin structures
Difference maps between the two three-dimensional reconstructions were carried out using the programs implemented in SPIDER (Frank et al., 1996) after previous alignment of the two structures. To test the significance of the differences revealed by the difference maps between the apo-CCT and CCT–tubulin three-dimensional reconstructions, the data sets were each divided into six groups and compared using Student’s t-test, as previously described by Roseman et al. (1996). Differences were considered significant when *P* <0.0005. As a control, internal comparisons within the same data set gave no significant differences.

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