Three-dimensional structures of the flagellar dynein–microtubule complex by cryoelectron microscopy

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Introduction

Cilia and flagella are elaborate organelles that play important roles in generating extracellular fluid currents in eukaryotes (Haimo and Rosenbaum, 1981; Goodenough and Heuser, 1985; Satir and Christensen, 2007). For example, left-right asymmetry in mammals is established by the nodal flow generated by the rotary movement of cilia (Hirokawa et al., 2006). The beating motion of cilia/flagella is driven by ciliary dyneins, which produce sliding forces between outer doublet microtubules (MTs) using ATP as an energy source (Gibbons, 1996). Elucidation of the mechanisms underlying the chemomechanical energy conversion by the dyneins is of great importance.

Since its discovery four decades ago (Gibbons and Rowe, 1965), the outer dynein arm (ODA) has been extensively studied by biochemical and structural methods. The *Chlamydomonas reinhardtii* ODA molecule is composed of three heavy chains and several smaller proteins (Mitchell, 2000; DiBella and King, 2001; Kamiya, 2002), and its total mass is ~2 MD. Isolated ODAs have three heads connected to a common base through thin stems (Goodenough and Heuser, 1982, 1984; Tsukita et al., 1983; Witman et al., 1983). The head domain contains six tandemly linked AAA+ (ATPases associated with diverse cellular activities) modules, which form a ringlike structure (Neuwald et al., 1999; Samso and Koonce, 2004).

Quick-freeze deep-etch (QFDE) EM studies of axonemes revealed that ODA has an elliptical head, which binds to the A tubule through two spherical feet (P foot and D foot), and a slender stalk that binds to the B tubule in an ATP-dependent manner (Goodenough and Heuser, 1982, 1984; Tsukita et al., 1983; Burgess, 1995; Lupetti et al., 2005). It has been hypothesized that the N-terminal stem and/or the stalk domains serve as lever arms that amplify ATP hydrolysis–dependent conformational changes within the head domain (Vallee and Gee, 1998; King, 2000; Asai and Koonce, 2001; Burgess et al., 2003; Kon et al., 2005; Reck-Peterson et al., 2006), but these hypotheses are based on studies of isolated dyneins in the absence of MTs. To elucidate the mechanism by which dynein generates force under physiological conditions, it is necessary to investigate the structural changes of MT-bound dyneins.

In this study, we established a method for reconstructing the 3D structure of the in vitro reconstituted ODA–microtubule complex and visualize nucleotide-dependent conformational changes using cryoelectron microscopy and image analysis. As the complex went from the rigor state to the relaxed state, the head domain of the β heavy chain shifted by 3.7 nm toward the B tubule and inclined 44° inwards. These observations suggest that there is a mechanism that converts head movement into the axonemal sliding motion.
Two MTs of the ODA–CB-MT complex is 24 nm, which is a uniform 24-nm periodicity (Fig. 2 F). The distance between computed diffraction pattern of the cryo-EM image again shows many views of the complex, allowing 3D reconstruction. The effective staining and cryo-EM, we found that two MTs twist over each other (Fig. 2, D and E; arrowheads). This twisting provides many views of the complex, allowing 3D reconstruction. The computed diffraction pattern of the cryo-EM image again shows a uniform 24-nm periodicity (Fig. 2 F). The distance between the two MTs of the ODA–CB-MT complex is 24 nm, which is the same as that observed by cryo-electron tomography of the axoneme (Nicastro et al., 2006). The moiré pattern of the MTs shows that both of the MTs in the complex have the same polarity (Sosa and Chretien, 1998).

3D reconstruction of the ODA-CB-MT complex
To reconstruct the 3D structure of the ODA–CB-MT complex, we first determined the helical arrangement of the complex from its cryo-EM images. In the layer line–filtered image (Fig. 3 A), the two rows of ODA molecules are staggered. This arrangement was also quantitatively confirmed by comparing the near- and far-side layer line at 24 nm. In principle, the phase difference between the two sides allows the determination of whether the left- and right-side ODAs is in phase (0°) or out of phase (180°; Moore et al., 1970). Judging from the phase difference at the amplitude peak (Fig. 3 B, arrowhead), ODAs are staggered as shown in Fig. 3 A. Therefore, we concluded that the complex has a twofold screw symmetry along its longitudinal axis.

Using this helical arrangement, we reconstructed the 3D structure of the ODA–CB-MT complex (Fig. 4 A and Video 1, available at http://www.jcb.org/cgi/content/full/jcb.200609038/DC1). Because the long twisting of the two MTs does not always follow the ideal helical symmetry, we combined single-particle analysis and helical image analysis to correct the distortions. We divided the images into 48-nm-long segments and calculated the rotation on the longitudinal axis from the distance between the two MTs. Translational alignment of the segments along the longitudinal axis was based on the 24-nm periodicity. Therefore, no assumption was made regarding the ODA structure. After 10–15 cycles of refinements, the 3D structures converged, and the effective resolution was determined to be 2.7 nm for the wild-type complex in the rigor state from the Fourier shell correlation (Fig. 3 C; van Heel, 1987).

Domain organization of ODA
Understanding the domain organization of ODA is essential for interpreting its structural changes. The end-on view of the ODA–CB-MT complex shows a pair of MTs cross-bridged by two rows of ODA molecules (Fig. 5 A). The 3D structure of the ODA–CB-MT complex in the rigor state is roughly divided into four domains: the α, β, and γ head domains and a base complex (Fig. 5, A–C). The stalk domain, which binds to MTs in a nucleotide-dependent manner (Johnson, 1985), should be located between the head domains and the β tubule. Although the stalk is visible in the quick-freeze replica (Fig. 2 C), it did not appear in the 3D reconstruction, possibly because of its flexibility and its thin structure. An ODA molecule binds to the α tubule through a thicker end, which we refer to as a base complex (Fig. 5, A–C; faint green densities). The base complex may contain two intermediate chains, which have WD repeat domains (Fig. 5, A–C; green bodies; Ogawa et al., 1995; Wilkerson et al., 1995; Madrona and Wilson, 2004).

To determine which side of the ODA–CB-MT complex harbors the ATP-sensitive MT-binding sites of ODA (Johnson, 1985), the complex was treated with 20 μM ATP and observed by cryo-EM and image analysis. By comparing the relaxed and rigor states of the ODA–MT complex, we observed the conformational changes that occur in the head domains. These results suggest that our system has a high potential to reveal the mechanism of dynein power generation.

Results
Characterization of the ODA–MT complex
To obtain dynein–MT complexes suitable for 3D reconstruction, we reconstituted the ODA–MT complex in vitro. This complex has a structure similar to the axoneme, is periodically decorated with ODAs, provides many views necessary for 3D reconstruction, and has MT translocation activity.

The dynein–MT complex is formed by copolymerizing tubulin in the presence of axonemal dyneins from Chlamydomonas reinhardtii (Haimo et al., 1979). SDS-PAGE analysis of the complex suggests that it contains the heavy, intermediate, and light chains as well as the docking complex (Fig. 1; Sakato and King, 2004). Because two MTs are cross-bridged by ODA, we refer to the complex as the ODA–cross-bridged MT (ODA–CB-MT) complex. QFDE-EM observation of the ODA–CB-MT complex (Com) were separated by SDS-PAGE and stained with Coomassie brilliant blue. HCs, heavy chains; DC1–3, docking complex protein 1–3; IC1/2, intermediate chain 1/2; LCs, light chains; MW, molecular weight marker.

Figure 1. SDS-PAGE analysis of the ODA–CB-MT complex. The high-salt extract from Chlamydomonas axonemes (Ex) and the purified ODA–CB-MT complex [Com] were separated by SDS-PAGE and stained with Coomassie brilliant blue. HCs, heavy chains; DC1–3, docking complex protein 1–3; IC1/2, intermediate chain 1/2; LCs, light chains; MW, molecular weight marker.

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by cryo-EM (Fig. 6 A). Because ATP treatment disrupts the ATP-sensitive binding of the ODAs, in the presence of ATP, the cross-bridged MTs separate, whereas the ATP-insensitive binding remains intact (Haimo and Rosenbaum, 1981). Haimo and Rosenbaum (1981) also reported that ATP depolymerizes the MTs. To prevent this depolymerization, the MTs were stabilized by the addition of taxol. The cryo-EM image of this hemicomplex showed that the 24-nm periodicity of the ODAs was unaffected by ATP treatment. We compared the averaged cryo-EM image of the hemicomplex with different projections from the 3D reconstruction of the ODA–CB-MT complex. We found that the characteristic triangular shape of the ODA density is also seen in the projection of the ODA bound to the MT by its base complex side (Fig. 6 B). The similarity in the 2D projection was also supported by the high cross-correlation value between the cryo-EM image of the hemicomplex and the projection of ODA. These results demonstrate that the base complex side of ODA–MT binding is ATP insensitive, which is consistent with the findings by Johnson (1985).

To examine the motor activity of the ODAs in the ODA–CB-MT complex, we performed a motility assay using the ATP-treated ODA–MT complex. We attached the AlexaFluor543-labeled ODA–MT complex onto a glass coverslip (Fig. 6 D, red) and applied AlexaFluor488-labeled plain MTs (Fig. 6 E, green). MT translocation was initiated by the addition of 100 μM ATP. All of the sliding MTs translocated along the ODA–MT complex (Fig. 6 E, arrowheads; and Video 3, available at http://www.jcb.org/cgi/content/full/jcb.200609038/DC1). The sliding velocity by the ODA–MT complex was 4.0 ± 2.6 μm/s (n = 20), which is similar to that by the αβ particle of ODA (Sakakibara and Nakayama, 1998). This result demonstrates that the row of ODAs in the ODA–CB-MT complex has motor activity.

Figure 2. EM images of the ODA–CB-MT complex by QFDE, negative-staining, and cryo-EM. A–C, QFDE; D, negative staining; E, cryo-EM. (A) A pair of cross-bridged MTs, as indicated by the arrowheads. The white lines show the row of ODAs. (B) The 24-nm periodicity is indicated by brackets. (C) The globular heads and the thin stalks (arrows) are shown. (D and E) The crossover points are indicated with arrowheads. The black lines indicate the ODAs. (F) The averaged diffraction pattern of the cryo-EM images of five ODA–CB-MT complexes showing the prominent 24-nm layer line. The layer line periodicities are indicated. (G) The interdynein connections observed by cryo-EM. The row of ODAs extending past the end of the MT is indicated with arrowheads. Bars, 100 nm.
and by Sakakibara et al. (1991), this allowed assignment of the outermost density in the top view of the ODA–CB-MT complex (Fig. 7 B, arrowhead). Together with the thin-section EM study of our 3D structure (Fig. 9; Goodenough and Heuser, 1982). The elliptical head corresponds to the α head domain and functions (King et al., 1985; Marchese-Ragona et al., 1989; Sakakibara et al., 1991, 1993; Sakakibara and Nakayama, 1998). To assign α, β, and γ heavy chains, we compared the 3D structures between the wild type and the oda-11 mutant (Fig. 7). The lack of an α heavy chain in oda-11 appeared as a loss of the outermost density in the top view of the ODA–CB-MT complex (Fig. 7 B, arrowhead). Together with the thin-section EM study by Sakakibara et al. (1991), this allowed assignment of the α, β, and γ heavy chains from the outermost to innermost positions (Fig. 5 A). If viewed from the side, the planes of the head domain rings appear parallel to the longitudinal axis of the MT (Figs. 5 C and 8 B), which is consistent with recent observations by Nicastro et al. (2006). The top view (Fig. 5 A) and cross section (Fig. 8 A) of our reconstruction also revealed that the planes of the head domain rings are not exactly parallel but rather tangential to the circumference of the B tubule (Figs. 5 A and 8 A). In addition, the three head domains are offset from each other longitudinally (Figs. 5 C and 8 B). The disk-shaped objects in the densities of the head domains (Fig. 5) summarize the aforementioned interpretations. The positions and orientations of the disks were determined from the center of gravity and the long axis of the densities, respectively (Fig. 8, A, B, D, and E).

We correlated the surface structures observed by QFDE-EM with our 3D structure (Fig. 9; Goodenough and Heuser, 1982). The elliptical head corresponds to the α head domain and a part of the β head domain. The boundary between the α and β head domains appears as a medial cleft within the head by QFDE-EM (see Figs. 7 and 8 in Goodenough and Heuser, 1982). The D foot corresponds to the γ head domain and part of the β head domain, and the P foot corresponds to the base complex.

### Interdomain connections

The interdomain connections (Fig. 5, A–C; arrowheads) indicate that the three heavy chains are functionally distinct. The α head domain is connected to the base complex (Fig. 5 C, red arrowhead). The β head domain has two densities extending to the base complex: one density extends toward the base complex within the same ODA molecule (Fig. 5, B and C; orange arrowhead), and the other extends toward the adjacent base complex (Fig. 5, B and C; and Fig. 8, C and F; arrowheads). ODAs are connected even in the absence of MTs (Fig. 2 G), suggesting that the inter-ODA connection is stable. The γ head domain is connected to the base complex (Fig. 5 C, yellow arrowhead) and also directly binds to the MT (Fig. 5 A, arrowhead). These observations suggest functional differentiation among the three head domains (see Discussion).

### Nucleotide-induced conformational changes

To observe the ODA–CB-MT complex in the relaxed state, we analyzed it in the presence of ATP and vanadate (Fig. 4 B and Video 2, available at http://www.jcb.org/cgi/content/full/jcb.200609038/DC1). It is known that the addition of ATP to vanadate-treated axonemes causes them to straighten and reduces their elastic bending resistance, which is referred to as the relaxation of axonemes (Sale and Gibbons, 1979; Okuno, 1980). Treatment with ATP and vanadate also causes large structural changes of ODA (Goodenough and Heuser, 1982; Tsukita et al., 1983; Burgess, 1995). To prepare the ODA–CB-MT complex in the relaxed state, we treated the complex in the rigor state with vanadate and ATP. The complex retained the cross-bridging, and the radius of the complex in the relaxed state did not change. Therefore, we applied the same image analysis methods as for the rigor state.

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**Figure 3.** Arrangement of ODAs within the ODA–CB-MT complex and resolution estimation of the reconstructions. (A) A 24-nm filtered cryo-EM image of the ODA–CB-MT complex showing the staggered arrangement of the ODAs. Lines indicate the positions of ODAs. (B) An amplitude and phase plot of the 24-nm layer line from the ODA–CB-MT complex. Phase diff: the phase difference between the near and far sides. The theoretical phase difference is 180° if the two rows of ODAs are staggered. Amplitude: the amplitude of the layer line. The phase value at the amplitude peak (arrowhead) indicates the staggered arrangement of the ODAs. (C) The Fourier shell correlation curves for each reconstruction. The intersection between each curve and the horizontal line at 0.3 was taken as the effective resolution. The effective resolutions are 2.7 nm, 2.7 nm, and 3.5 nm for the rigor state, the relaxed state, and the oda-11 mutant, respectively.

**Figure 4.** 3D reconstructions of the ODA–CB-MT complex from the wild type in two nucleotide states. A, rigor state; B, relaxed state. Stereo pairs are presented. The color reflects the distance from the center of the MT. The surfaces represent the isosurface of 120% of the volume estimated from the molecular weight of ODA.
The conformational changes are different among the three head domains (Figs. 5 and 8, D–F). The density of the α head domain is unclear (Fig. 5 F), probably because of high heterogeneity. The center of gravity of the β head domain shifted 3.7 nm toward the B tubule (Fig. 8, A and D), and its head plane inclined 44° inward (Fig. 8, B and E) compared with its conformation in the rigor state. In contrast, the γ head domain remained in almost the same position as in the rigor state (Fig. 8, A and D). As a result of the conformational change, the distance between the β head domain and the B tubule became shorter (Fig. 5, A and D; and Fig. 10, B–D), from 14 nm in the rigor state to 10 nm in the relaxed state. Considering the reported length of the stalk domain (H1160 15 nm; Goodenough and Heuser, 1982, 1984; Gee et al., 1997), the change in the distance may affect the orientation of the stalk domain.

Our reconstruction and the QFDE-EM image in the relaxed state are similar except for the D foot (Fig. 9, C and D). The D foot in the QFDE-EM image is apart from the P foot of the adjacent ODA. In our reconstruction, the D foot, which corresponds to the γ head domain and part of the β head domain, is close to the next P foot, which corresponds to the base complex. The difference in ATP concentration (1 mM for the QFDE-EM image and 2 μM for our reconstruction) may be the reason for the different appearance.

Discussion

This study establishes that the ODA–CB-MT complex is suitable for reconstructing the 3D structure of dynein from cryo-EM images. Our reconstruction revealed the detailed 3D arrangement of the ODA complex and the conformational changes between ODAs in two functionally critical states without the influence of other axonemal proteins.

The elaborate interdomain linkers and nucleotide-induced conformational changes of the ODA indicate that the three head domains have different functions. The appearance of the α head domain both in the rigor and relaxed states suggests that it is more flexible than the other two head domains, which may be related to a proposed regulatory function (Sakakibara and Kamiya, 1989; Sakakibara et al., 1991; Sakato and King, 2004). The β heavy chain may be important for synchronized movement of ODAs along the axoneme by communication through the inter-ODA connections. Given that the swimming velocity of the oda-11 α heavy chain–missing mutant is slightly slower than that of the wild type (Sakakibara et al., 1991), the nucleotide-induced movement of the β head domain presumably generates major force for the sliding. The γ head domain, which showed little displacement, may serve as a stable anchor to the A tubule.
This displacement of the β head is a new observation, and it is distinct from the rotation of the head proposed by Burgess et al. (2003). Because their hypothesis is based on the negative-staining EM of isolated dynein in the absence of MTs, the movement of dynein relative to the MT cannot be observed; however, it is possible that both the rotation and the displacement occur at the same time and that they amplify the sliding speed.

Because the β head domain displaced vertically in the direction of sliding, there should be a mechanism for converting the direction of the movement. The stalk domain is the most probable candidate for mediating this conversion because it bridges the head domain and the β tubule. One possible explanation for how the stalk domain converts the movement of the head domain is as follows: if the stalk length is constant, the angle of the stalk is a function of distance between the head domain and the MT. For example, if the stalk length is 15 nm, the 3.7-nm movement of the head domain results in a 40° rotation of the stalk, which can produce a 7-nm sliding movement at the stalk head. To investigate these possibilities, the orientation of the head domain and stalk must be determined by specific labeling (for example, by using antibodies).

In the 3D reconstructions and in the aforementioned discussions, we assumed that most of the ODAs were in the relaxed state. This assumption is supported by our EM observations as well as previous biochemical data (described in this paragraph). When ODA and tubulin are copolymerized, there is more than one type of ODA–MT complex (Haimo et al., 1979). For structural analysis, we used the ODA–CB–MT complex, but it also contains ODAs complexed with single MTs as shown in Fig. 2 G. In this ODA–single MT complex, ODAs are bound to MTs by their stalks. All of these ODAs seem to dissociate from the MTs upon treatment with ATP and vanadate because we did not observe single MTs decorated by ODA via its stalk in the same grids, in which we collected cryo-EM images of the ODA–CB–MT complex (unpublished data). This demonstrates that stalk-side binding of most ODAs became weak after the ATP-vanadate treatment, which agrees with the finding of Shimizu and Johnson (1983b). It is also consistent with previous biochemical data: according to Shimizu and Johnson (1983a), 2 μM is the lowest concentration of ATP that can completely dissociate ODA from MTs within 1 s. Furthermore, according to kinetic studies of dynein, the rate constants of ATP binding, ATP hydrolysis, phosphate release, and vanadate binding are all faster than 8/s in our condition (Johnson, 1983; Porter and Johnson, 1983; Shimizu and Johnson, 1983a; Tani and Kamimura, 1999). As we incubated the ODA–CB–MT complex with ATP-vanadate for 10–15 s, <1% of ODA should have remained in the rigor state. Although the occurrence of the cross-bridging even in the relaxed ODA–CB–MT complex is perplexing, we think that the cross-bridging of the two MTs is maintained by the sum of many weak ODA–MT interactions.

The arrangement of ODAs in the ODA–CB–MT complex also raises the concern that the sliding directions induced by the two rows of ODA are antagonistic and should cause distortion of the complex. As we show in this study, cross-bridging of the ODA–CB–MT complex was retained in the presence of ATP and vanadate probably because ODA was in the presliding state and did not exert force in this nucleotide state. It seems that phosphate release from ODA in the ADP + Pi state is required to separate the cross-bridging, as shown in Fig. 6 A. These results are...
consistent with other biochemical data showing that ATP and vanadate inhibit the MT sliding (Sale and Gibbons, 1979; Satir et al., 1981; Warner and Mitchell, 1981) and that force generation is suggested to require the release of phosphate (Johnson, 1983; Johnson et al., 1984; Kon et al., 2005). Therefore, it is reasonable to assume that our relaxed structure represents the presliding state.

In a recent study, cryo-electron tomography was applied to study axoneme structures, including ODA (Nicastro et al., 2006). Although their overall architecture of ODA in the rigor state is quite similar to our reconstruction, there are minor differences in the assignment of the molecular boundary and of the inter-ODA linker to a particular heavy chain. Nicastro et al. (2006) did not show the position of the base complex relative to the head domains in the tomograph of the C. reinhardii axoneme tomograph. In the present study, we assigned the base complex to the minus end side of the head domains (Fig. 5, green spherical objects). This assignment is based on the globular shape of ODA and the molecular boundary between adjacent molecules (Figs. 4 and 5 and Video 1) and is consistent with the QFDE-EM study (Fig. 9; Goodenough and Heuser, 1982). Future high-resolution studies on isolated ODA complexed with MTs may be needed to clarify the domain assignments.

Nicastro et al. (2006) also interpreted that ODA molecules are connected along the MT by an outer outer dynein linker, which extends diagonally from the α head domain to the neighboring γ head domain (see Fig. 5 D in Nicastro et al., 2006). In our map, the inter-ODA linker, which is functionally equivalent to outer outer dynein, seems to connect the β head domain and the neighboring base complex. A similar inter-ODA linker was also observed by QFDE-EM (see Fig. 6 in Goodenough and Heuser, 1982). The difference between the interpretations of
Materials and methods

Strains

The C. reinhardtii 137c strain and the oda-11 mutant strain, which lacks the α heavy chain and the 16kD light chain, were provided by R. Kamiya (University of Tokyo, Tokyo, Japan) and S.M. King (University of Connecticut Health Center, Farmington, CT), respectively.

Preparation of the ODA–CB-MT complex

Axonemes were obtained from wild type (137c) and the oda-11 mutant C. reinhardtii by the dibucaine method (Witman et al., 1978; Kagami and Kamiya, 1992). The axoneme extract was desalted using a Sephadex G-25 gel filtration column equilibrated in PEM buffer (100 mM Pipes-NaOH, pH 6.8, 1 mM MgCl₂, and 1 mM EGTA). The eluent was concentrated to 2–3 mg/ml using Centricon YM-100 (Millipore). Bovine tubulin was purified as described previously (Shelanski et al., 1973). MTs were polymerized in PEM buffer with 1 mM GTP in the presence of 1.3 mg/ml axonemal extract at 30°C for 1 h. The ODA–CB-MT complexes were purified by centrifugation through a 50% (wt/vol) sucrose cushion in PEM buffer at 35,000 g for 30 min at 30°C.

In vitro motility assay

ODA–CB-MT was labeled with AlexaFlour543 (Invitrogen), and the molar ratio of unlabeled to labeled tubulin was 20:1. 100 μg/ml of the ODA–CB-MT complex in HMDE buffer (10 mM Heps-NaOH, pH 7.2, 5 mM MgCl₂, 1 mM DTT, 1 mM EGTA, and 1 mM PMSF) containing 10 μM paclitaxel was treated with 20 μM ATP and attached onto the glass-bottom microwell dish (MatTek). Next, the glass surface was blocked with HMDE buffer containing 1 mg/ml casein. Plain MTs labeled with 10 μg/ml AlexaFlour488 in HMDE buffer containing 0.1% NP-40 were sheared with a 28-gauge needle and added to the dish. MT translocation was initiated by adding 100 μM ATP (final concentration). The movements of the MTs were observed using a total internal reflection fluorescence microscope (IX71; Olympus) equipped with plan Apo 100× NA 1.45 total internal reflection fluorescence microscopy oil immersion lens (Nikon). The images were projected onto a CCD camera (Cascade II; Photometrics) and were contrast enhanced with RS Image software (Roper Scientific). Experiments were performed at room temperature.

EM

The MT pellet was resuspended in HMDE buffer and absorbed to holey carbon grids. After washing with HMDETN buffer (HMDE buffer containing 10% trehalose and 0.2% NP-40, the grids were blotted for 5 s and plunged into liquid ethane (−180°C). Images of the ODA–CB-MT complex were recorded on SO-163 film under low-dose conditions with an electron microscope (JEM-2200FS; JEOL) at a nominal magnification of 40,000× with a 15,000–35,000 Å defocus. For QFDE-EM, the MT pellet was resuspended in PEM buffer without sucrose and centrifuged again. Quick-freeze replica of the MTs were prepared as described previously (Hirokawa and Heuser, 1981). For negative-staining EM, samples were fixed in 0.5% glutaraldehyde and stained with 5% uranyl acetate.

Relaxed state

To prepare the ODA–CB-MT complex in the relaxed state, we treated the complex in the rigor state with 10 μM vanadate from the sucrose cushion step and added 2 μM ATP 10–15 s before plunge freezing. This concentration of ATP is shown to be enough for relaxing the axoneme (Okuno, 1980).

Image processing

ODA–CB-MT complexes that have MTs with 14 protofilaments/three-start helix were selected (Ray et al., 1993), and their polarity was determined based on the moire pattern in the image (Sosa and Chretien, 1998). Selected micrographs were digitized with a scanner (LeafScan 45; Scitex) at a pixel size of 10 μm, which corresponds to 2.5 Å in the specimen. Each filament was straightened by fitting the longitudinal axis to a cubic spline curve. The straightened filaments were divided into 48-nm-long segments.

For the initial angle assignment for each segment, we assumed that the distance between the two MTs is constant and used a simple relationship between the diameter of the complex and the rotation angle around the axis: $\alpha = \cos^{-1} \left( \frac{d}{d_{\text{max}}} \right)$, where $d$ is the distance between the centers of the two MTs, and $d_{\text{max}}$ is the maximum of the distance determined from two representative ODA–CB-MT complexes.

For the translational alignment along the long axis of the complex, we generated a model reference composed of two MTs twisted over each other and spherical densities aligned along the MT with a periodicity of 24 nm. We filtered the Fourier transformed image and the reference segments, leaving 6-, 12-, and 24-nm layer lines and calculated the cross-correlation between the image and the reference. The hand of the helix was determined to be left by the images of QFDE and negative-staining EM with tilting.

Initial 3D reconstruction was generated using back projection based on the rotation and translation determined as described in the previous two paragraphs. We imposed twofold helical symmetry on the reconstruction and applied solvent flattening with a five-pixel Gaussian mask (pixel size = 4.872 Å).
After each refinement, we replaced the reference with the symmetry-imposed and masked reconstruction. We repeated this cycle until the phase residual values converged.

The numbers of ODA molecules and filaments used for each of the reconstructions were as follows: 3,737 molecules from 31 filaments for the rigor state, 3,872 molecules from 38 filaments for the relaxed state, and 1,350 molecules from 11 filaments for the oda-11 mutant. The effective resolution was determined to be 2.7 nm, 2.7 nm, and 3.5 nm for the rigor state, the relaxed state, and the oda-11 mutant, respectively, from the Fourier shell correlation between two independent datasets using a 0.3 cut off. Image analysis was performed using Ruby-Helix scripts (Metlagel et al., 2007) and Frealign (Grigorieff, 1998). Surface renderings were performed with UCSF Chimera (Pettersen et al., 2004), Pymol (Delano Scientific), or the AVS software package (Advanced Visual Systems).

References


