

Structural insights into the architecture and assembly of eukaryotic flagella

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ABSTRACT Cilia and flagella are slender projections found on most eukaryotic cells including unicellular organisms such as *Chlamydomonas*, *Trypanosoma* and *Tetrahymena*, where they serve motility and signaling functions. The cilium is a large molecular machine consisting of hundreds of different proteins that are trafficked into the organelle to organize a repetitive microtubule-based axoneme. Several recent studies took advantage of improved cryo-EM methodology to unravel the high-resolution structures of ciliary complexes. These include the recently reported purification and structure determination of axonemal doublet microtubules from the green alga *Chlamydomonas reinhardtii*, which allows for the modeling of more than 30 associated protein factors to provide deep molecular insight into the architecture and repetitive nature of doublet microtubules. In addition, we will review several recent contributions that dissect the structure and function of ciliary trafficking complexes that ferry structural and signaling components between the cell body and the cilium organelle.

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Abbreviations:

DMT – doublet MT,

EM – electron microscopy,

ET – electron tomography,

IDA – inner dynein arm,

IFT – intraflagellar transport,

MIP – MT inner protein,

MT – microtubules,

N-DRC – nexin-dynein regulatory complex,

ODA – outer dynein arm.

PF – protofilament,

RS – radial spoke.

INTRODUCTION

Microbial cells often utilize cellular structures known as flagella to swim in aqueous environments. Flagellum-driven motility can occur in response to chemical stimuli (chemotaxis) or in response to a light source (phototaxis) and allows cells to swim towards beneficial environments. Bacterial flagella are surface appendages that serve as helical propellers often essential for the successful infection by pathogens [1]. Although different types of bacterial flagella exist, they have a common architecture based on a rotary ATPase motor that via Basal body and Hook structures links to and propels a long filament that typically extends from the cell body [2]. Archaeal microbes can also utilize flagella for motility, but these flagella are only superficially similar to the bacterial counterparts and have evolved independently to have different structures and assembly mechanisms [2].

Flagella are also found on eukaryotic microbes but are evolutionarily and structurally distinct from their prokaryotic counterparts. The motile eukaryotic flagellum (also known as a cilium) was discovered in 1675 by Antoine von Leeuwenhoek [3] and is conserved on a wide range of unicellular eukaryotic organisms such as the green alga *Chlamydomonas reinhardtii* [4], the parasite *Trypanosoma brucei* [5, 6], and the ciliates *Paramecium* and *Tetrahymena* [7]. Yeasts are important exceptions as they lack cilia, which were presumably lost during evolution. The flagellum on eukaryotic microorganisms, like that of prokaryotes, serves motility and sensory reception functions and allow cells to swim in response to external cues [8].

In mammals, motile cilia power the sperm cells but are also present on the apical surface of epithelial cells where they generate external fluid flows [9]. This is the case for

airway cilia that create a mucus flow to clear the lungs of dust particles and pathogens, and cilia in the fallopian tubes that create a fluid flow to move the egg cell from the ovaries to the uterus [10]. Additionally, many vertebrate cells express a single primary cilium that is non-motile but serves important functions in sensory reception and signaling [11]. Primary and motile cilia share a common architecture consisting of a basal body that templates the microtubule (MT)-based axoneme, and a ciliary membrane that is continuous with the plasma membrane but compositionally distinct [8] (Figure 1a). The basal body organizes the growth of nine MT triplets [12] that transition into

doublets that constitute the structural backbone of the ciliary axoneme. A transition zone, formed by Y-shaped structures, bridges MTs and plasma membrane, separates the ciliary and cytosolic compartment and provides a selectivity barrier at the ciliary base [13–15]. In motile cilia, the nine outer doublet microtubules (DMTs) typically encircle two central MT singlets (Figure 1b). Motile cilia are also structurally different from primary cilia given that they harbour inner and outer dynein arms, which are macromolecular motor complexes that associate with DMTs and hydrolyse ATP to power the ciliary beat required for swimming [16–19] (Figure 1b). However, these macromo-

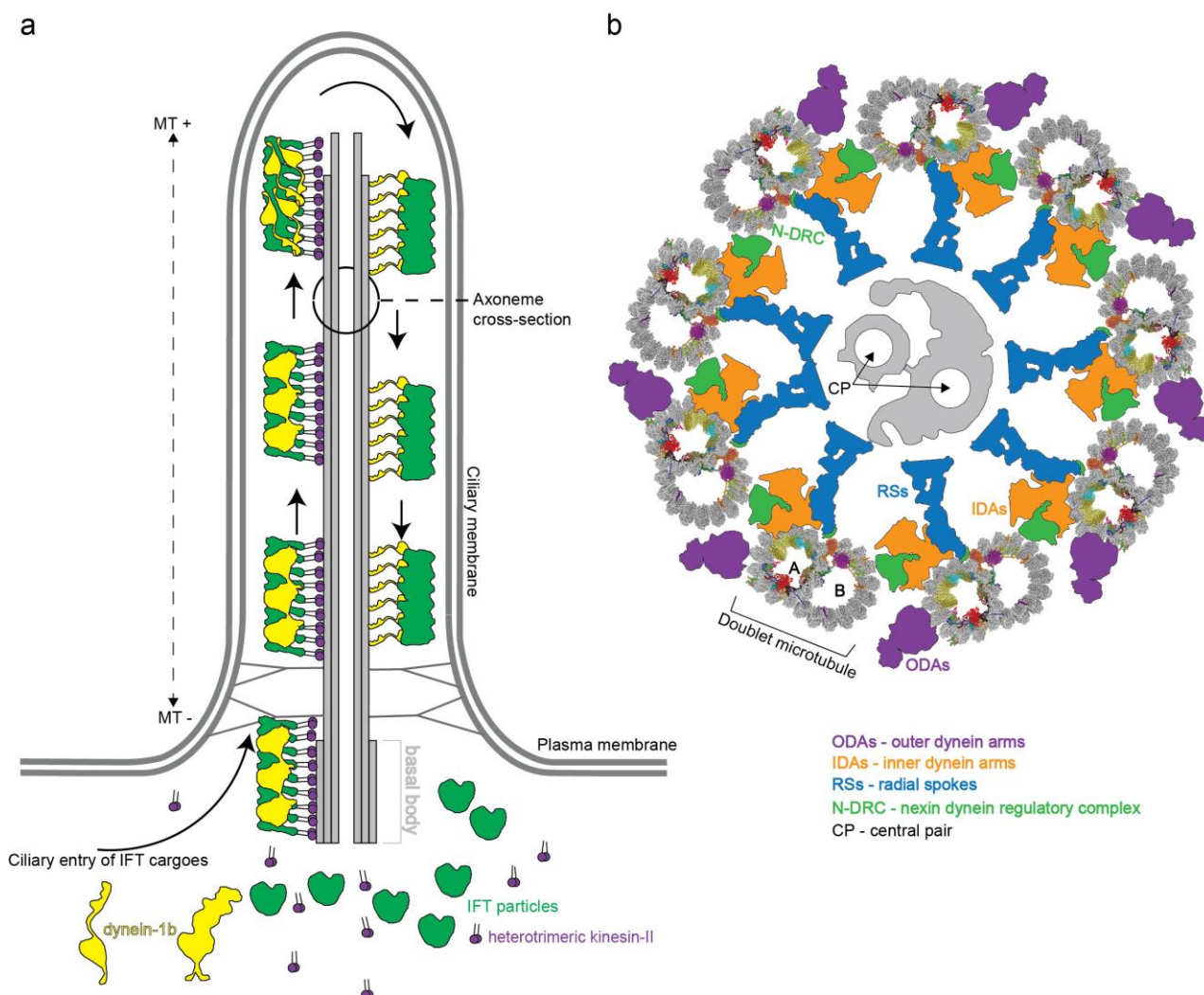


FIGURE 1: Schematic representation of a cilium from *Chlamydomonas reinhardtii*. (a) Architecture of a cilium with a simplified scheme of the bi-directional intraflagellar transport (IFT) system trafficking proteins between the cilium and the cell body. IFT complexes (green) assemble into train-like polymers powered by heterotrimeric kinesin-II (purple) in the anterograde direction (base->tip). Inactive dynein-1b (yellow) is loaded as a cargo onto anterograde IFT ‘trains’. Once assembled and loaded with ciliary cargo, these trains are driven across the transition zone and Y-links (depicted as grey connections spanning from the ciliary membrane to the axoneme at the ciliary base) to reach the ciliary tip. Upon arrival at the ciliary tip, the IFT ‘trains’ are re-modelled, turnover products are picked up and moved back again to the ciliary base by retrograde IFT powered by the now activated dynein-1b motor. (b) Schematic representation of an axoneme cross section of a motile cilium, which depicts the nine peripheral doublet microtubules (DMTs; PDB entry: 6u42) that surround a central microtubule pair (CP, shown in grey). The DMTs are connected through the nexin dynein regulatory complex (N-DRC, shown in green). Complexes responsible for ciliary beating are the inner- and outer- dynein arms (IDAs and ODAs; orange and magenta, respectively). The central pair is connected to the nine DMTs by radial spokes (RS; blue).

lecular complexes are not present on all DMTs. One of the nine DMTs is lacking the outer dynein arms [20] while some inner dynein arms are missing in parts of the axoneme [21]. In addition, some of the dyneins have a preferred location at either the distal or the proximal end of the axoneme providing a longitudinal asymmetry [4, 22]. How DMTs maintain stability during ciliary beat has long been enigmatic given that cytoplasmic MTs easily break when exposed to similar forces [23–25]. It has however been evident from electron tomographic reconstructions of flagella *in situ* that the inner lumen of DMTs harbour large densities likely representing unknown protein factors [26–32]. Significant progress in understanding DMTs came with a recent publication where the authors made use of technical advances in cryo electron microscopy (cryo-EM) [33, 34] to obtain high-resolution single-particle reconstructions of DMTs, which allowed for the structural modelling of 33 MT inner proteins (MIPs) associated with the lumen of DMTs [35]. Important implications for DMT stability and periodicity gleaned from these structures will be highlighted in this review.

ARCHITECTURE OF THE AXONEME

Proteomics studies reveal that as many as 500–1000 unique proteins participate in the construction of the motile eukaryotic flagellum [36, 37]. How all these factors associate to assemble ciliary axonemes has been a major topic of structural studies by EM since the early 1950. Initial studies of negatively stained axonemes from different species revealed the 9+2 arrangement [38–40] where the two central fibrils have a different size than that of the nine outer fibrils [40]. We now know that this difference in size originates from doublet vs singlet MTs. In the early 1960s it was indeed shown that these 9+2 fibril structures consist of MTs with similar properties to MTs found in the cytoplasm of cells [41]. Both cytoplasmic and axonemal MTs are built from a protofilament (PF) of α - and β -tubulin heterodimers, which provide a polarity for MT-based axoneme with (+) at the tip and (–) at the base (Figure 1a). MTs often contain 13 PFs although the number may vary [41]. Cytoplasmic MTs are dynamic structures that undergo cycles of polymerization and depolymerization of $\alpha\beta$ -tubulin and are key players in cellular processes such as cell motility and cell division as well as intracellular transport [42].

The repetitive nature of 8.0 nm $\alpha\beta$ -heterodimeric tubulins dictates a periodic arrangement of DMT-binding proteins. This axonemal periodicity is advantageous in structural studies as it allows for averaging of computationally extracted 48 nm or 96 nm repeating units imaged by cryo electron tomography (cryo-ET), which has been a powerful technique producing reconstructions at 20–40 angstroms (Å) resolutions [26–29, 34, 43–53]. These studies have determined the arrangement and conformation of several axonemal-associated protein complexes such as outer dynein arms (ODAs) and inner dynein arms (IDAs). ODAs and IDAs are ATP-hydrolysing motor complexes that facilitates ciliary beating through a controlled microtubule sliding motion [54] (Figure 1b). Another large axonemal mac-

romolecular complex is the nexin-dynein regulatory complex (N-DRC) that bridges adjacent DMTs [28]. The N-DRC was recently shown to bind polyglutamylated tubulin via electrostatic interactions to regulate flagellar motility [55]. Additionally, the architecture of the T-shaped protein radial spoke (RS) complexes that connect DMTs to the central pair MTs has been resolved [47]. ODA, IDA, RS and N-DRC axonemal structures have been comprehensively reviewed by others [4, 56, 57] and will not be further covered here.

DMTs are highly conserved structures built from an incomplete B-tubule fused onto a complete A-tubule composed of 13 tubulin PFs [24, 58] (Figure 2). Structural studies of axonemal MTs have a long history with the first three-dimensional models published as early as 1974 based on negatively stained electron micrographs, which clearly resolved the 8 nm repeat distance between $\alpha\beta$ -tubulin heterodimers [59]. Analysis of these early negative stain EM pictures could however not clearly resolve if the B-tubule of DMTs consists of ten or eleven PFs [60]. However, with the improved resolution of cryo-ET reconstructions of *Chlamydomonas* and sea urchin sperm flagella it was clearly shown that the B-tubule consists of ten PFs [27, 30]. Extra density that could be mistaken for an 11th PF instead corresponds to an inner junction protein complex, which attaches the B-tubule onto the A-tubule [30], and is composed of the FAP20/PACRG complex [61] (Figure 2). A subsequent study of DMTs used both cryo-ET and single particle cryo-EM to reach an improved resolution of 19 Å, which allowed for the exact assignment of the structurally similar α - and β -tubulin subunits within the DMTs based on EM data alone [51]. This study revealed that within the DMTs, the lattice pattern of tubulin isoforms is a B-lattice characterized by a left-handed helical arrangement of the PFs with a 0.92 nm stagger between α - α and β - β tubulins. Along the A-tubule, the 13 PFs are forming a discontinuity or a “seam” that lies between the PFs nine and ten [51, 59, 62].

MOLECULAR STRUCTURE OF CILIARY DOUBLET MICROTUBULES AND ASSOCIATED PROTEIN FACTORS

Earlier studies have identified a class of MIPs associated with the lumen of *C. reinhardtii*, *T. thermophila* and sea urchin sperm DMTs [27, 30, 63, 64]. The periodicity of MIPs with respect to the $\alpha\beta$ -tubulin lattices has also been resolved [51] but the identity, structure and function of these MIPs remained enigmatic. A recent study pushed the boundaries of axonemal structural biology significantly by imaging extracted and purified *Chlamydomonas* axonemal DMTs by cryo-EM to obtain single-particle reconstructions of unprecedented resolution [35]. 3.6 Å resolution maps was obtained for the computationally extracted 96 nm DMT repeat whereas a somewhat better resolution of 3.4 Å was obtained for the 48 nm repeat. At this resolution, α - and β -tubulins as well as the respective nucleotide states (GDP vs GTP) could be distinguished in the cryo-EM maps [35]. These maps also allowed for the fold recognition and in most cases molecular modelling of 38 MT associated proteins of which 33 are MIPs associated with the lumen of

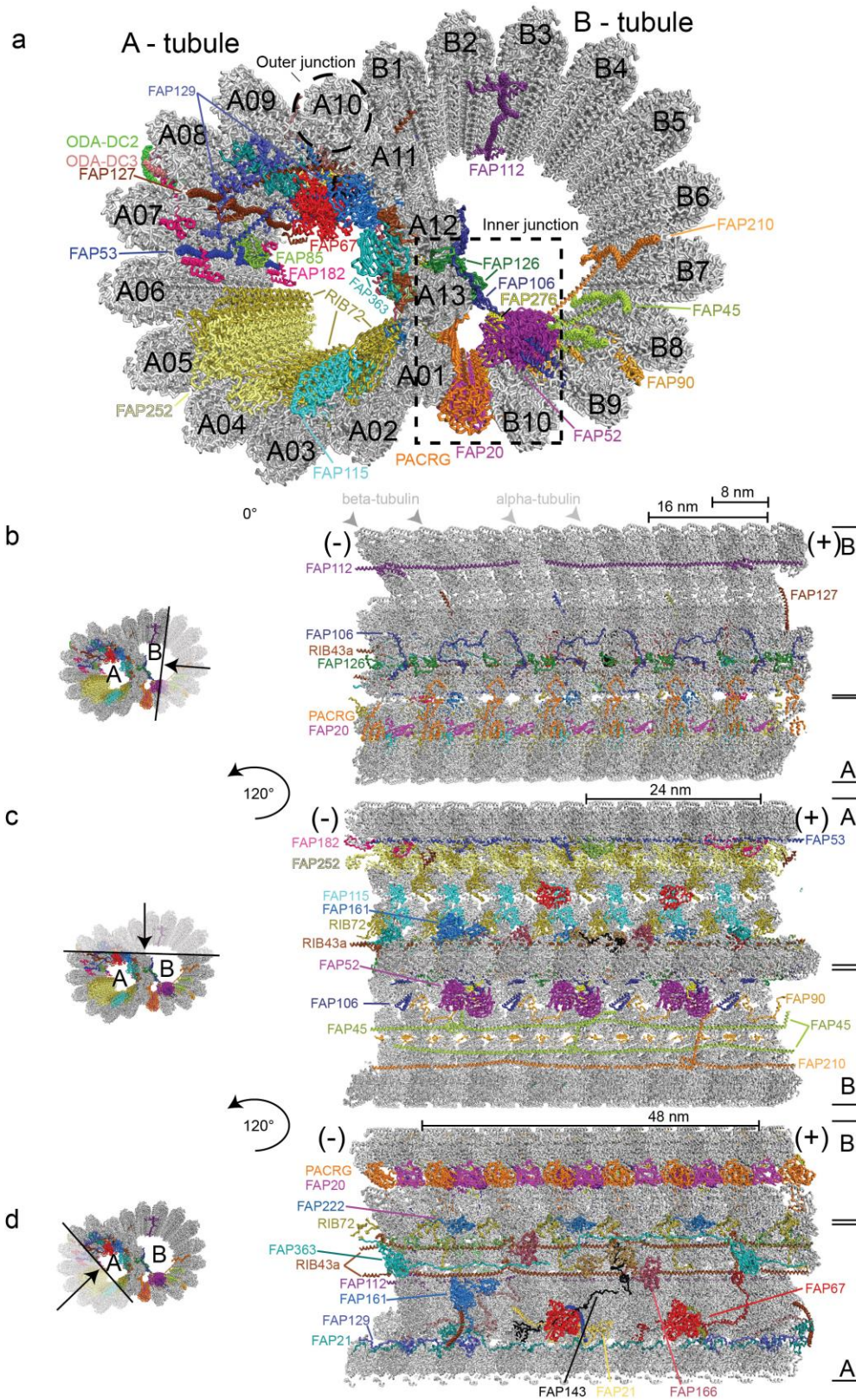


FIGURE 2: Structure of the 48nm ciliary doublet microtubule repeat (pdb entry: 6u42). **(a)** Cross section through a ciliary DMT. The outer dynein arms docking complexes 2 and 3 serve as docking sites for ODAs on the exterior of the DMTs and are labelled ODA-DC2 and ODA-DC3. Inner microtubule proteins (MIPs) are highlighted in different colours and labelled according to protein name. **(b-d)** Depiction of microtubule inner proteins, their interaction network and periodicity visualized within the confinements of the doublet microtubule from the minus (-) to the plus (+) end. **(b)** Lateral view of MIPs decorating the lumen of B-tubule as seen after a 10 nm deep slice facing the viewer was removed. The black arrow in the transparent region on the left representation indicates the region that has been removed for better clarity. The ODA-DC complex is left out as part of the removed section. **(c)** A lateral view displaying MIPs decorating both the A- and B-tubules. A 10 nm deep slice facing the viewer was removed as shown by the black arrow in the transparent region on the left representation. MIPs are visualized after rotation of the DMTs of 120° along the longitudinal axis relative to (b). **(d)** Visualisation of MIPs decorating the lumen of the A-tubule after removal of a 10 nm deep slice facing the viewer and rotation of the doublet microtubule with 120° along its longitudinal axis relative to (c). In this representation the alternating PCRG/FAP20 complex is observed from the exterior of the B-tubule.

DMTs [35]. Proteins seen to associate with the outer surface of DMTs include the ODA docking complex (ODA-DC), which is located between A07 and A08 PFs (A07-A08), has 24 nm periodicity and is required for the proper attachment of ODAs to the DMTs (**Figure 2a**). Furthermore, a 96

nm long coiled-coil segment was identified as Ccdc39/Ccdc40 [35] supporting the notion that this complex acts as a molecular ruler defining the 96 nm periodicity of the axoneme [65].

Cryo-EM reconstruction of the 48 nm repeat allows structural modelling of 33 inner microtubule binding proteins

In the Ma *et al.* study, the 48 nm repeat reconstruction of DMTs displayed the highest resolution and allowed for the modelling the atomic structures of 33 MIPs (Figure 2). Molecular models were obtained by a combination of automatic fold recognition and chain tracing using the cryo-EM map [35]. To validate the correct assignment of proteins to their respective map densities, the structures of axonemal DMTs of mutants missing either the 72 kDa ribbon-associated protein RIB72 or the flagellar associated protein 166 (FAP166) were determined and the lack of density at the respective positions confirmed. Two isoforms of RIB72 were recently identified in *Tetrahymena* as MIPs associated with the lumen of the A-tubule and were shown to be important for ciliary assembly and motility [64]. Many more MIPs are found to associate with the lumen of the A-tubule than the B-tubule (Figure 2). An intricate network of protein-protein interactions connects different PFs with MIPs both within the respective tubules but also between A- and B-tubules [35] (Figure 2). These interactions likely account for the increased stability and longevity of axonemal DMTs compared to cytoplasmic MTs [23, 24]. Indeed, a recent study convincingly demonstrated using high-speed atomic force microscopy that FAP45 and FAP52, both proteins found in the lumen of DMTs, stabilize ciliary axonemes [25].

The 33 MIPs can be divided into different groups depending on their periodicity along the axoneme. MIPs with a periodicity of 8 nm such as FAP115, FAP252 and the C-terminal domains of RIB72 follow the 8 nm repeat distance of $\alpha\beta$ -tubulin heterodimers and bind to protofilament A01-A05 (Figure 2a and 2c). The PACRG/FAP20 complex that forms the inner junction between PF B10 of the B-tubule and A01 of the A-tubule also adheres to the 8 nm repeat although the ladder of PACRG/FAP20 is not completely continuous as one copy of PACRG is missing for every 96 nm repeat [30, 35]. *Chlamydomonas* mutants lacking FAP20 display split axonemes consistent with the role of PACRG/FAP20 in stabilizing the junction between A- and B-tubules [61]. The PACRG-FAP20 complex could be mistaken for a MT PF at lower resolution although this complex clearly has smaller overall dimensions than $\alpha\beta$ -tubulin.

Interestingly, the N-terminal portion of RIB72 follows a 16 nm periodicity and interacts with other 16 nm repeats MIPs including FAP106, FAP126 and FAP52, which constitute a direct link between PFs of the A- and B-tubules (Figure 2a and 2b). FAP52 makes extensive contacts to FAP276 that also follows 16 nm periodicity [66] (Figure 2a). Both FAP52 and FAP276 display an overall Y-shape and connect perpendicularly on the MTDs with the inner junction formed by the PACRG/FAP20 complex, the groove between microtubules B09 - B10 and FAP106 (Figure 2a, [35, 66]). Furthermore, both the N- and C-terminus of FAP276 contact the tubulin lattice [66], which may provide additional stability in gluing the A- and B-tubules together at the inner junction. Several MIPs extend from the internal lumen to the external surface of the

DMTs where they may facilitate inside-outside communication and create new binding sites on the outside of DMTs. This is the case for the MIPs FAP85, FAP129 and FAP182 (48 nm periodicity) that bind in the cleft between PFs A07-A08 and contact ODA-DC on the exterior of DMTs (Figure 2a, [35]). FAP129 interacts with FAP127 on the inside and with ODA-DC on the exterior side, which may provide a molecular rationale for why FAP127 mutants show defects in ODA-DC assembly [67, 68]. The structure of DMTs presented by Ma *et al.* thus demonstrates that the distinction between inner and outer DMT-binding proteins is not so clear-cut and that MIPs may indeed protrude through 'windows' between tubulin subunits to affect the exterior of DMTs [69].

Several MIPs polymerize along the axoneme via end-to-end interactions to establish periodicity. This is the case for the coiled-coil protein RIB43a, where two copies occupy the spaces between PFs A11-A12 and A12-A13 (Figure 2d). RIB43a self-associates along the DMTs through N-C termini interactions and the two copies of RIB43a are bridged by one copy of the protein FAP363 via a globular HSP70-like domain (Figure 2a and 2d). FAP363 also self-assembles via N-C termini interactions. Many of the MIPs thus have the propensity for periodic assembly onto DMTs encoded by their amino acid sequence and thus do not rely solely on the repetitive nature of MTs to establish periodicity along the axoneme. These end-to-end MIPs also help define the periodicity of other MIPs by creating new binding sites at the lumen of DMTs [35]. Interestingly, these end-to-end MIPs are some of the first axonemal subunits to be expressed upon flagellum assembly [70], likely because of their important role in establishing the axonemal periodicity required for the attachment of other axonemal-associated proteins.

The role of tubulin post-translational modifications as binding sites for MIPs

MTs undergo a wide range of post-translational modifications that constitute a tubulin code with important implications for MT function [71, 72]. These include (poly)glutamylation and (poly)glycylation added to α - and β -tubulin C-terminal tails found on the exterior of MTs. In the DMT structure presented by Ma *et al.* the C-terminal tubulin tails are mostly invisible in the cryo-EM map because of flexibility or proteolytic removal by the subtilisin enzyme used in DMT preparation. However, on PF A01, ten residues of the C-terminal tail of β -tubulin is visibly engaged in contacts with the PACRG/FAP20 complex and presumably strengthen this interaction, although it is not clear if post-translational modification plays a role here. Another long-studied MT modification is acetylation of the ϵ -group of the side-chain of lysine 40 (K40) in α -tubulin found on the luminal side of MTs [73, 74]. K40 acetylation is catalysed by α -tubulin acetyltransferase (α TAT1) [75–77] and occurs on polymerized MTs, which requires that α TAT1 gains access to the MT lumen either through openings at the MT ends or through tubulin lattice defects along the MTs [78–80]. It is a long-known fact that K40 acetylation correlates with a subset of long-lived MTs in cells but is has

been a conundrum if K40 acetylation infers MT stability or if this modification is simply a mark that accumulates on 'old' MTs [74, 81, 82]. It was shown that K40 acetylation does indeed provide mechanical stabilization of MTs [83]. Structural studies of acetylated MTs did, however, not provide a clear answer to the molecular basis given that the K40 loop region was disordered, although modelling did suggest that acetylation did reduce lateral contacts [84, 85]. Interestingly, in the structure of DMTs, the K40 loop is ordered on about 10% of the B-tubule and about 40% of the A-tubule where it participates in several interactions with various MIPs [35]. Acetylated K40-loops of α -tubulin thus serve as binding sites for MIPs and may indeed help organize the intricate network of MIPs bound at the lumen of DMTs. The structure thus suggests that K40 acetylation stabilize DMTs indirectly through the interactions with MIPs that in turn bridge PFs of both A- and B-tubules to create long-lived stable DMTs [35]. However, it is currently not known if K40 acetylation is required to recruit MIPs or if MIP-recruitment is a requirement for K40 acetylation.

INTRAFLAGELLAR TRANSPORT ON DOUBLET MICROTUBULES

DMTs do not only serve as the structural backbone of flagella but also function as 'tracks' for transport and delivery of protein cargoes. This intracellular trafficking system, known as intraflagellar transport (IFT), has evolved to ferry both structural and signalling components between the cell body and the cilium [86]. IFT was discovered in 1993 by Kozminski and Rosenbaum as a bi-directional transport process in the flagella of *Chlamydomonas* [87]. In *Chlamydomonas*, the molecular motor known as kinesin-II walks along DMTs towards the (+) end at the ciliary tip thus powering IFT in the anterograde direction [88–90]. The retrograde motor dynein1b or dynein-2 in mammals walks along DMTs towards the (–) of MTs bring IFT material to the base of the cilium [88, 91, 92]. IFT also relies on the large multi-subunit IFT particle for association with ciliary cargos such as tubulin [93–96] or ODAs [97–102] and utilize an octameric complex called the BBSome to couple the IFT system to various membrane associated signalling components for ciliary exit [103–105]. High-resolution structures are available for several IFT sub-complexes [106–110] and lower resolution structures of complete IFT trains were determined *in situ* by cryo-ET [111]. Furthermore, several recent structural cryo-EM studies of mammalian BBSome complexes were recently published [112–115]. The IFT and BBSome complexes were extensively reviewed previously and will not be further covered here [105, 116].

Given that kinesin and dynein motors both travel on DMTs, but in opposite directions, the question arises of why collisions between anterograde and retrograde IFT 'trains' are not observed. At least two different models could allow for this: 1) Anterograde and retrograde trafficking could occur on different DMTs or 2) occur on different tubules of the same DMT, thus making the DMT a bi-

directional double track for IFT [117]. An elegant study used correlative light and electron microscopy to show that hypothesis 2) is correct in *Chlamydomonas* where the kinesin-2 motors travel on B-tubules and the dynein-2 on A-tubules for retrograde transport [118]. Furthermore, in *Chlamydomonas*, all nine DMTs are actively used for IFT [118]. This situation is different in the unicellular parasite *Trypanosoma brucei* where only doublets 3-4 and 7-8 are utilized for IFT [117]. However, IFT was also observed to travel bi-directionally on each set of doublets in *Trypanosomes* suggesting that the double track mechanism may be evolutionarily conserved [117]. From the DMT structure, it is clear that IFT 'trains' have limited physical space to access the MT PFs because of numerous other associated protein complexes (Figure 1b, [35]). This is the case for the A-tubule where only PFs A08-A10 are accessible to the IFT dynein motor. Kinesin-driven anterograde IFT 'trains' have more space on the B-tubule but the fact that these IFT 'trains' also contact the ciliary membrane suggests that they are restricted to PFs B01-04 [35, 118].

During anterograde IFT, inactive dynein is loaded onto IFT trains as a cargo and delivered to the ciliary tip [111, 119, 120]. Recent cryo-EM studies elucidated the structure of human dynein-2 at a resolution of 4.4 Å [120] and revealed a dual mechanism of inactivation during anterograde transport. Firstly, dynein-2 motor domains have an intrinsic propensity to stack against each other in an inactive dimeric conformation where the speed of dynein-2 on MTs is decreased to 140 nm/s [120]. Contrary, a dynein mutant that prevents dimerization walks along MTs with a velocity of 530 nm/s, which is in agreement with the speed of purified monomeric dynein-2 motor alone [119] and with the speed of retrograde IFT in mammalian cells [104]. Interestingly, the inactive conformation of dynein-2 dimers provided a good fit to the density in electron tomograms of IFT 'trains' suggesting that this is indeed the state adopted by dynein-2 during anterograde IFT [111, 120]. Secondly, when the auto-inhibited dynein-2 dimer is loaded onto IFT trains, the MT-binding domains are oriented so that they point away from the MTs, which prevents dynein-2 in engaging the DMT tracks during anterograde IFT [111, 120].

Fluorescent labelling revealed that the anterograde IFT 'trains' are fragmented at the ciliary tip, which suggest that a re-modeling event takes place [121–123]. Arrival of IFT 'trains' at the ciliary tip allows for activation of dynein-2 [122, 124], engagement on the microtubule tracks for the retrograde IFT (Figure 1a) as well as kinesin-II detachment and diffusion or transport back to the cell body [121]. The re-configuration of dynein-2 at the ciliary tip is supported by cryo-EM studies that suggest an open conformation of dynein-2 motor domains [111] (Figure 1a). It is unclear how the inactive dynein-2 is converted at the tip to the active form that drives the retrograde IFT. Because no additional factors such as dynactin were found to modulated the activity of dynein-2 [125–127], it is possible that components of the IFT complex are responsible. There are three GTPases (IFT22, IFT27 and RabL2) embedded in the IFT 'trains' travelling at different stages of anterograde IFT [106, 110, 128, 129]. Of these, IFT22 and IFT27 accompa-

nies the IFT ‘trains’ all the way to the tip where they may participate in remodelling of the IFT ‘trains’. Together with IFT25, IFT27 was shown to be required for the exit of certain GPCRs from cilia [130, 131] most likely indirectly by controlling ciliary trafficking of the BBSome. Interestingly, IFT27 seems to specifically sense ubiquitinated GPCRs and trigger their ciliary removal [132, 133]. Additionally, it was shown that the ubiquitination machinery is present in the flagella of *Chlamydomonas* [134] and that components of the IFT machinery interact with ubiquitinated proteins such as α -tubulin during cilia disassembly [135]. Furthermore, molecular genetics studies suggest that the membrane-associated IFT subunit IFT172 is also important for dynein-2 targeting or turnaround at the ciliary tip by an unknown mechanism that likely involves the *Chlamydomonas* microtubule end binding protein 1 [136–139]. Interestingly, IFT172 predominantly co-immunoprecipitates a version of the dynein-2 heavy chain that is somewhat larger in molecular mass suggesting that it could be modified, perhaps by ubiquitination [124]. Taken together, these data raise the possibility that ubiquitination plays important roles in IFT tip turnaround and retrograde transport. Another mechanism that can contribute to ciliary turnaround at the tip is phosphorylation. Specific kinases were discovered to localize at the tip of cilia both in *Chlamydomonas* and in mammalian cells targeting the kinesin-II motor and disrupting its interaction with IFT-B proteins [140, 141]. The com-

prehensive molecular mechanisms of dynein-2 activation and remodelling of anterograde to retrograde IFT ‘trains’ remains to be elucidated.

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

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