An Essential Quality Control Mechanism at the Eukaryotic Basal Body Prior to Intraflagellar Transport

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Constructing a eukaryotic cilium/flagellum is a demanding task requiring the transport of proteins from their cytoplasmic synthesis site into a spatially and environmentally distinct cellular compartment. The clear potential hazard is that import of aberrant proteins could seriously disable cilia/flagella assembly or turnover processes. Here, we reveal that tubulin protein destined for incorporation into axonemal microtubules interacts with a tubulin cofactor C (TBCC) domain-containing protein that is specifically located at the mature basal body transitional fibres. RNA interference-mediated ablation of this protein results in axonemal microtubule defects but no effect on other microtubule populations within the cell. Bioinformatics analysis indicates that this protein belongs to a clade of flagellum-specific TBCC-like proteins that includes the human protein, XRP2, mutations which lead to certain forms of the hereditary eye disease retinitis pigmentosa. Taken with other observations regarding the role of transitional fibres in cilium/flagellum assembly, we suggest that a localized protein processing capacity embedded at transitional fibres ensures the ‘quality’ of tubulin imported into the cilium/flagellum, and further, that loss of a ciliary/flagellar quality control capability may underpin a number of human genetic disorders.

Key words: basal body, chaperone, flagellum, intraflagellar transport, retinitis pigmentosa, tubulin, tubulin cofactor C, trypanosome

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Eukaryotic cilia and flagella are evolutionarily conserved organelles that perform a diversity of biological functions, ranging from motility to sensory perception and are vital to human health. Defects in cilium/flagellum function underpin a wide range of inherited human disorders including respiratory disease, retinal degeneration as well as more complex developmental disorders such as Bardet–Biedl syndrome (BBS) and hydrocephalus (1). The construction of a flagellum/cilium, involving assembly of several hundred distinct proteins (2–5), is made more difficult by the fact that in most eukaryotes, assembly occurs in a ribosome-free cellular compartment distinct, and ultimately distant, from the normal cellular cytoplasm (6). This assembly mechanism provides additional challenges and necessitates accurate intracellular targeting, processing and transport of proteins into the cilium/flagellum.

With a few exceptions (7,8), cilium/flagellum assembly relies upon an evolutionarily conserved bi-directional transport mechanism known as intraflagellar transport (IFT) (reviewed in 9). Transitional fibres radiating from the mature basal body demarcate the boundary of the cilium/flagellum compartment and are pivotal to this process acting as both docking sites for IFT motor proteins and regulating entry of IFT particles into the cilium/flagellum (10,11). Molecular understanding of IFT has advanced rapidly in recent years, yet our knowledge of events relating to recruitment and processing of proteins prior to IFT-mediated translocation remains poor.

The major structural component of all eukaryotic cilia/flagella (whether motile or sensory) is a microtubule-based axoneme constructed from the heterodimeric protein, α/β-tubulin. The generation of a functional α/β-tubulin heterodimer depends upon a well-defined tubulin-folding pathway, in which quasi-native intermediates sequentially interact with specific tubulin cofactors (TBCC) (12–15). Functional α/β-tubulin heterodimers are ultimately generated from an interaction between TBCC and a super-complex containing α/β-tubulin monomers and tubulin-specific cofactors TBCD and TBCE. This interaction with TBCC stimulates hydrolysis of GTP by β-tubulin and releases GDP-bound α/β-tubulin heterodimer; subsequent free exchange results in the production of a GTP-bound tubulin heterodimer competent for microtubule incorporation. The necessity to hydrolyse GTP to enact heterodimer release may function as a quality control step ensuring that only GTPase competent heterodimer is released. This is an important consideration as polymerization of α/β-tubulin heterodimers into microtubules is intimately coupled to hydrolysis of the GTP nucleotide bound to β-tubulin. The question therefore arises, as to how such protein quality control processes are handled when tubulin is exported from the cytoplasm into the flagellum compartment.

We have exploited the experimentally tractable flagellated protozoan parasite Trypanosoma brucei (a pathogen of
significant medical and veterinary importance in Sub-Saharan Africa) to investigate tubulin processing requirements during cilium/flagellum formation. *T. brucei* possesses a single flagellum and forms a new flagellum during each cell division cycle. Although some ciliated/flagellated cells (e.g. the green algae *Chlamydomonas reinhardtii*) resorb cilia/flagella during the cell cycle, *T. brucei* (and many other ciliated/flagellated cells) divide with flagella present. Trypanosomes thus provide a model to compare processes operating to construct and maintain a new and old flagellum, respectively, within the same cell.

This study reveals that a flagellum-specific isoform of TBCC is embedded at transitional fibres radiating from the mature basal body, where it provides an essential cilium/flagellum-specific function. Detailed bioinformatics analyses show that this trypanosome protein belongs to a family of related TBCC domain-containing proteins, whose phylogenetic occurrence is restricted to eukaryotic organisms capable of forming a flagellum/cilium. This clade also contains the human protein XRP2, mutations in which lead to certain forms of the hereditary eye disease X-linked retinitis pigmentosa (RP, OMIM 312600), characterized by progressive degeneration of photoreceptor cells in the retina. Within retinal cells, a modified cilium provides an intracellular link between the functionally and morphologically distinct inner and outer photoreceptor compartments (16) and we suggest that our results may provide insight into the molecular basis for both this disease and also a number of other inherited human diseases characterized by dysfunctional cilia.

Results

**Bioinformatics analysis identifies a flagellum/cilium TBCC**

We recently completed a comprehensive proteomic analysis of the *T. brucei* flagellum resulting in the publication of the *T. brucei* flagellar proteome (TbFP) (2). Of the 331 proteins present in the TbFP, 86 are present in the genomes of at least one other flagellated/ciliated eukaryote but absent in the genomes of non-flagellates. Given the evolutionary distance between trypanosomes and other flagellated/ciliated eukaryotes, these proteins represent a highly conserved flagellum/cilium-specific cohort. Among this conserved cohort, we identified a *T. brucei* protein (GeneDB accession number Tb10.61.2870), which encodes a domain with homology to TBCC, one of the key molecular chaperones required to assemble functional \( \alpha/\beta \)-tubulin heterodimer. As TBCC is likely to have an essential role in tubulin formation in all eukaryotes, identification of this protein as flagellum/cilium specific appeared anomalous.

Using an iterative profile-based searching technique (based on a Pfam TBCC Hidden Markov model seed alignment), we interrogated a wide range of phylogenetically diverse eukaryotic organisms to identify all proteins possessing homology with the Pfam-defined TBCC domain (Figure 1). This analysis revealed that many (but intriguingly not all) eukaryotes encode several TBCC proteins that resolve into three distinct clades. Clade 1 contains TBCC proteins from a diverse range of eukaryotes and includes the canonical human and yeast cytoplasmic TBCC protein essential for de novo \( \alpha/\beta \)-tubulin heterodimer formation. However, our analysis reveals that TBCC domain-containing proteins in clade 2 are restricted to eukaryotes capable of forming a cilium/flagellum and includes human RP2 and the Tb10.61.2870 protein detected in our flagellar proteome (hereafter designated TbRP2). Clade 3 contains proteins from a range of flagellated and non-flagellated eukaryotes. However, the TBCC domain identified in this clade does not easily align with proteins from clades 1 and 2, particularly with regard to a key arginine residue previously identified as critical to TBCC protein function (17), raising questions regarding the functionality of TBCC proteins within this clade. Our analysis also reveals another intriguing anomaly. Despite constructing cilia and/or flagella, the fruit fly *Drosophila melanogaster* and the apicomplexan parasite *Plasmodium falciparum* do not encode a flagellate-specific isoform of TBCC. This may simply reflect the different way these organisms build cilia/flagella (8,18).

**RNA interference-mediated ablation of TbRP2 results in structurally aberrant flagella**

Biochemical studies have shown that human RP2 protein has partial functional overlap with canonical TBCC, in that RP2 can stimulate the GTPase activity of native tubulin heterodimers but cannot generate \( \alpha/\beta \)-tubulin heterodimer from \( \alpha/\beta \)-tubulin monomers (17). Despite this biochemical evidence, there is no direct experimental evidence demonstrating that RP2 mutations lead to microtubule abnormalities and so the in vivo function of RP2 remains a matter of debate (19). To address this question, we subjected the trypanosome protein TbRP2 to functional analysis using RNA interference (RNAi) and demonstrated that induction of RNAi in procyclic (insect stage) trypanosomes resulted in increasingly severe growth defects from 48 h post-induction (Figure 2A). Real-time reverse transcriptase-polymerase chain reaction experiments show reduction of TbRP2 messenger RNA levels to <10% of non-induced controls within 48 h of induction (data not shown).

Trypanosomes possess a single flagellum and construct a new flagellum during each cell cycle. As the new flagellum is invariably positioned posterior to the old, it is possible to unambiguously distinguish new and old flagella in dividing trypanosomes. Observation of TbRP2-RNAi induced trypanosomes reveals that the new flagellum is abnormally short (compare the non-induced cell in Figure 2B with the TbRP2-ablated cell in Figure 2C). Confirmation that the length of this new flagellum is aberrant comes from comparison with cell-cycle event markers
such as mitosis. In control trypanosomes, mitosis initiates when the new flagellum length is approximately 10 μm (Figure 2D), and during the subsequent M phase, the new flagellum continues to elongate to over 15 μm as the mitotic spindle elongates (to approximately 8 μm). In contrast, although mitotic spindle microtubules elongate...
as normal in TbRP2-RNAi induced trypanosomes, the new flagellum remains abnormally short (Figure 2E).

This lack of a mitotic defect is significant, as it indicates that spindle microtubule formation is not adversely affected by TbRP2 ablation. Careful examination of the trypanosome cytoskeleton shown in Figure 2C also demonstrates the integrity of sub-pellicular microtubules in the cell body is unaffected. However, a detailed and extensive examination of axonemal ultrastructure in these TbRP2-ablated cells reveals that approximately 36% of axonemes display a range of structural abnormalities; including missing or mis-aligned central pair microtubules and/or disruptions to outer doublet microtubules (Figure 2F). In contrast, a canonical 9+2 axonemal microtubule configuration was observed in non-induced trypanosomes (Figure 2G). These results demonstrate that TbRP2 ablation specifically compromises axonemal microtubule formation in the flagellum, without affecting other microtubule populations within the cell; a finding fully consistent with our bioinformatics predictions. It is significant that the ultrastructure of the paraflagellar rod (PFR) is unaffected in these flagella (the PFR is an extra-axonemal structure within the trypanosome flagellum [20]), as this establishes that TbRP2 ablation has specific effects on axonemal microtubule formation rather than a more generalized effect on IFT processes.

**TbRP2 specifically interacts with α-tubulin at transitional fibres radiating from the mature basal body**

To address the question of how TbRP2 fulfils this flagellum-specific role, we generated trypanosome cell lines expressing a TbRP2:green fluorescent protein (GFP) fusion protein. These studies demonstrated specific localization of TbRP2:GFP at the mature basal body subtending the flagellum (Figure 3A). Importantly, this localization was maintained on detergent-extracted cytoskeletons, indicating that TbRP2 protein is tightly associated with the basal body. At this locality, TbRP2 would be ideally positioned to act in a flagellum-specific capacity, assessing the GTPase competency of native tubulin heterodimer prior to allowing its IFT-mediated transport into the cilium/flagellum compartment.

To provide experimental evidence in support of this proposal, we asked whether it was possible to visualize a specific interaction between TbRP2 and tubulin protein at the basal body. The antibody YL1/2 [specific for a carboxyl-tyrosinated form of α-tubulin (21)], proved informative in this regard. In many eukaryotes, α-tubulin is synthesized with a C-terminal tyrosine residue, but this tyrosine residue is removed following microtubule assembly (reviewed in 22). As new microtubules are tyrosinated but become progressively de tyrosinated over time, YL1/2 is regarded as a specific marker for new microtubules. A highly characteristic labelling pattern has previously been reported for YL1/2 on trypanosome cells; tyrosinated α-tubulin is restricted to microtubules within the posterior third of the cell body, the axoneme of the new flagellum and the basal bodies subtending new and old flagella (23,24). However, labelling of the trypanosome basal bodies is rather puzzling because basal body microtubules will have been formed at least within the previous cell cycle and in many cases, several generations before that.

We therefore re-investigated this and determined that whilst the YL1/2 labelling presented as two discrete punctuate dots, these did not represent the basal body and probasal body but were specifically associated with the mature basal body (Figure 3B). In colocalization studies, we demonstrate that this YL1/2 labelling actually colocalizes with TbRP2:GFP in a distinct ring-shaped structure around the mature basal body (Figure 3C–E). The ‘two-dot’ pattern emerges when the ring is visualized from the side. Subsequent immunogold electron microscopy experiments confirmed this view and revealed that YL1/2 specifically labels transitional fibres radiating from the mature basal body (Figure 3F). In Figure 3G, colloidal gold particles are clearly evident on transitional fibres radiating from the mature basal body subtending the flagellum, while the immature probasal body (lying orthogonal to the mature basal body in this figure) is YL1/2 negative. Later in the trypanosome cell cycle when both basal bodies are mature (as evidenced by their possession of a transition zone, basal plate and axoneme nucleation), both are YL1/2 positive (Figure 3H). The acquisition of YL1/2 reactivity through the trypanosome cell cycle can be also visualized by immunofluorescence microscopy (Figure 3I–K). Figure 3I and J likely represent intermediate stages in the cell cycle, in which acquisition of YL1/2 reactivity is associated with probasal body elongation and nucleation of the new flagellum. This is an important observation as it indicates that an aggregation of tyrosinated α-tubulin forms around the basal body as it matures to nucleate the formation of a new flagellum. It should also be noted that the presence of YL1/2 labelling on basal bodies subtending both the old and new flagella suggests a continuing need for tubulin turnover within the microtubule axoneme.

It should be emphasized that we are not suggesting that transitional fibres are constructed from α-tubulin; rather that YL1/2 detects a pool of unpolymerized tyrosinated α-tubulin recruited to the basal body and awaiting transport into the flagellar compartment by IFT. However, as YL1/2 labelling is retained on whole cells, cytoskeletons and isolated flagella, this pool of unpolymerized tyrosinated α-tubulin must be stably associated with the transitional fibres. Significantly following ablation of TbRP2, YL1/2 staining of the basal body is lost (compare basal body staining in Figure 3L and M) suggesting not only that TbRP2 interacts with tyrosinated α-tubulin, but also that it plays a critical role in the recruitment and/or retention of tubulin at the transitional fibres. In contrast, YL1/2 labelling of the cell body remains intense, confirming that the incorporation of tyrosinated α-tubulin into sub-pellicular microtubules is unaffected in TbRP2-ablated cells.
**Discussion**

Our study resolves a long-standing conundrum as to why YL1/2 (a reliable and documented marker for new microtubules) apparently labeled basal bodies given their microtubule maturation cycle (25). Our studies demonstrate that YL1/2 detects an aggregation of unpolymerized tubulin rather than the basal body microtubules themselves.

In most ciliatedflagellated eukaryotic cells, the formation of a ciliumflagellum is dependent upon IFT, an evolutionarily conserved transport mechanism that moves proteins bi-directionally along the length of the flagellum. The basal body is pivotal to IFT, with transitional fibres radiating from the mature basal body acting as the docking site for IFT motor proteins at the base of the ciliumflagellum (reviewed in 26). These transitional fibres provide a critical
interface between cytoplasmic and cilium/flagellum compartments and are proposed to regulate IFT particle entry into the cilium/flagellum (11). However, data from this current study further extends the role of the basal body in cilium/flagellum assembly and suggests that a specific isoform of TBCC embedded at these transitional fibres acts to influence the 'quality' of tubulin heterodimers destined for axonemal incorporation.

Previous biochemical studies suggest functional overlap between RP2 and TBCC (17), whereby RP2 acts to stimulate the GTPase activity of native tubulin heterodimers but does not function in the generation of α/β-tubulin heterodimer from tubulin monomers (17). This biochemical evidence indicates that RP2 is not part of a canonical tubulin-folding pathway but rather provides specific functions related to assessing GTPase activity of tubulin heterodimer. Our bioinformatics analysis further reveals that RP2-like proteins are phylogenetically restricted to ciliated/flagellated organisms and suggests that this protein has evolved to deal with the specific demands of tubulin provision in a cilium/flagellum context; a proposal strongly supported by our experimental RNAi studies.

Despite clear biochemical evidence indicating a role for RP2 in microtubule dynamics (17), the role of RP2 in vivo remains a matter of debate because it has recently been suggested that RP2 exhibits 3' to 5' exonuclease activity and relocates to the nucleus in response to DNA damage (19). While our study does not address this proposed exonuclease role, it provides direct in vivo evidence that axonemal microtubule formation is defective following ablation of TbRP2. This result is consistent with previously published in vitro biochemical evidence indicating a role for RP2 in tubulin processing.

However, in this processing context, it will be important to determine whether RP2 acts solely upon newly translated tubulin heterodimers or also upon re-recycling tubulin heterodimers previously assembled into other microtubule structures. Elegant experimental studies carried out on the biflagellate algal cell Chlamydomonas, suggest that reutilization of tubulin heterodimer may occur in flagellated eukaryotic cells (27). These studies demonstrated that following removal of one of its two flagella, Chlamydomonas shortens the remaining flagellum and simultaneously regenerates the amputated flagellum. This pattern of flagellum elongation and shortening was not affected by cycloheximide (a treatment that blocks new protein synthesis). In addition to accessing pre-existing cytoplasmic pools of tubulin, it is possible that these cells also re-utilize tubulin heterodimers, previously incorporated in the axoneme of the shortening flagellum. Our bioinformatics analysis reveals that Chlamydomonas encodes two RP2-like proteins, one of which at least is present in the Chlamydomonas flagellar proteome (5). We suggest that elucidation of the role of these Chlamydomonas RP2-like proteins in tubulin recycling will be informative.

As ablation of TbRP2 results exclusively in axonemal microtubule defects, our data provide strong evidence that ciliary defects observed in X-linked RP2 arise from aberrant tubulin processing. Interestingly, the C. elegans protein K08D12.2 (a RP2-like protein also found in clade 2) also localizes to the basal body of ciliated neuronal cells; indeed, the ring-like localization pattern reported for K08D12.2 (28) is remarkably consistent with our localization of TbRP2:GFP.

However, there is an important caveat to this argument; as the human RP2 has not been reported to localize to the basal body but rather to the plasma membrane because of dual acyl modification of the N-terminus of the protein (29). If RP2 function in sensory neurons relates solely to tubulin processing, then one has to rehearse the argument that N-acylated RP2 proteins can fulfill this protein-processing function from the plasma membrane. A clue to how this might occur comes from studies on human retinal cells which show that membrane bound RP2 interacts with microtubule associated proteins such as ADP-ribosylation factor-like 3 (ARL3). It is suggested that this interaction links the membrane and microtubule cytoskeleton and acts to regulate membrane traffic and/or cell signalling (47).

In this study, we demonstrate that TbRP2 (a flagellum-specific isoform of TBCC) is exclusively localized at transitional fibres radiating from the trypanosome basal body. As RNAi-mediated ablation of TbRP2 results in the specific loss of tyrosinated α-tubulin from the basal body, we suggest a specific interaction occurs between these two proteins. As transitional fibres demarcate the boundary between the cytoplasm and the environmentally distinct flagellum compartment, we propose that TbRP2 establishes a 'quality control gateway' to assess the GTPase proficiency of tubulin heterodimer prior to IFT-mediated transport into the cilium/flagellum.

One can also rehearse another role for a tubulin quality control gateway at the basal body, namely that it may prevent or select for entry of post-translationally modified tubulin dimers into the flagellum compartment. If this is true, then enzymatic functions associated with the reversible tubulin post-translational modifications (tyrosination, acetylation and glutamylation) might also be expected to localize to the basal body. In trypanosomes, recycled tubulin would need to be retysorinated through the action of a tubulin tyrosine ligase (TTL) (24,30). Can the basal body also act as a localized factory to retysorinate α-tubulin previously incorporated into microtubules? Several TTL-like proteins and other tubulin modification enzymes (such as polyglutamylases) are encoded within the trypanosome genome (31) and one might predict that these proteins may also be localized to the trypanosome basal body.

Finally, we suggest that aberrant protein processing at the cilium/flagellum basal body may underpin a wide-range of
human ciliopathies, as we note that mutations in BBS6 and BBS10 (proteins with homology to α and ζ subunits of cytosolic chaperonin) also result in the development of Bardet–Biedl syndrome (a condition characterized by genital, renal and retinal abnormalities resulting from abnormal basal body function and cilium formation) (32,33). The establishment of a ‘quality control’ gateway at the basal body may therefore represent a widespread mechanism by which ciliated/flagellated cells ensure the fidelity of the microtubule axoneme.

Materials and Methods

Trypanosome culture and generation of T. brucei RNAi cell lines
Procytic T. brucei cells were routinely grown in semi-defined medium-79 media supplemented with 15% fetal calf serum as previously described (34). Trypanosome cell growth was monitored using a CASY1® cell counter and analyser system (Scharfe System GmbH) with cultures diluted on a daily basis to maintain a density of between 1 x 10^6 and 8 x 10^6 cells/mL. A 577 nucleotide fragment of the T. brucei, Tb10.61.2870 coding sequence was amplified by polymerase chain reaction (PCR) using gene-specific primers

\[ 5' - GCAGCTCGAGATAACCCTGCACATCTTCGC - 3' \]

and 50

\[ 5' - GCAGGGATCCGAGAATCGTTCTCGTCCTCG - 3' \]

amplified by PCR using gene-specific primers 2870RNAiF (5'- GAACGCCTGCAATGCTCCAAGTTCCTTC-3') and 2870-RNAiR (5'- CGAGGGATCCGAGAATCGTTCTCGTCCTCG-3'), and cloned into the T. brucei RNAi vector, p2T7-177 (55). The resulting construct was linearized with NotI, transformed into the procyclic 29-13 cell line and positive transformants were selected using 2.5 μg/mL hygromycin, 15 μg/mL G418 and 50 μg/mL hygromycin as previously described (36).

Induction of RNAi and phenotypic analysis
Trypanosome cultures (grown in drug-free media for 24 h prior to initiation of RNAi studies) were diluted to 1 x 10^6 and 8 x 10^6 cells/mL and doxycycline was added to a final concentration of 1 μg/mL. Every 24 h, cell cultures were counted and diluted back to a density of 1 x 10^6 cells/mL with fresh doxycycline being added to the culture medium.

Green fluorescent protein tagging of the endogenous Tb10.61.2870 open reading frame
DNA fragments from the 3' end of the Tb10.61.2870 open reading frame (ORF) sequence (minus the TAG stop codon) and the immediate 3'-untranslated region (UTR) downstream of the TAG stop codon were amplified by PCR using the primer combinations Tb10.61.2870/orfF-Tb10.61.2870/orfR and Tb10.61.2870/orfF-Tb10.61.2870/orfR, respectively (primer sequences available on request). Resultant DNA fragments were simultaneously ligated into pET-HYG-GFP (Devaux et al., 2007). The resultant pET-GFP:2870 vector was subsequently linearized with Xhol and transfected into procyclic T. brucei 427 strain cells. Positive cells were selected using media containing 20 μg/mL hygromycin as described (37).

Immunofluorescence studies
Procytic trypanosomes (both wild type and cells expressing GFP fusion proteins) were prepared for microscopy essentially as previously described (38). Slides were incubated with selected primary antibodies YL1/2 (39), LBC4 (40) or BBA4 (41). Slides were examined either on a DeltaVision RT microscope equipped with a Hamamatsu CCD camera and images processed in SoFtWoRx (Applied Precision) or on a Zeiss Axioplan 2 microscope equipped with a CCD camera controlled by METAMORPH software (Universal Imaging) and processed in METAMORPH. All images were subsequently processed using Adobe Photoshop (Adobe).

Protein Quality Control at the Eukaryotic Basal Body

Electron microscopy and immunogold labelling
Trypanosomes were harvested by centrifugation (800 x g, 10 min), fixed in 2% paraformaldehyde, 2% glutaraldehyde and 0.2% picric acid in 100 mM phosphate buffer (pH 7.2), post-fixed and processed for transmission electron microscopy as described previously (42). Preparation of negatively stained whole mount cytoskeletons for transmission electron microscopy was carried out using previously established methods (42,43). For pre-embedding, immunogold labelling cells were fixed with 8% formaldehyde in 100 mM phosphate buffer (pH 7.0) with 0.5% Triton-X-100 for ~2 min. The cells were gently pelleted and briefly quenched (2 x 30 seconds) with PBS containing 1% BSA, 50 mM glycine and 0.1% Tween-20 followed by a brief wash in PBS blocking solution (PBS containing 1% BSA and 0.1% Tween-20). Samples were incubated with YL1/2 (diluted 1:2 in PBS blocking buffer) for 2 h at room temperature. After washing with blocking buffer (3 x 10 min), the samples were incubated with a goat anti-rat antibody conjugated with 5 nm gold (1:50 dilution with PBS blocking buffer), washed with blocker (2 x 10 min) and the samples were fixed and processed as described above. Ultra-thin (~70 nm thick) sections were examined unstained in a Philips 400 electron microscope.

Bioinformatics analyses
Tubulin cofactor C homologues in diverse eukaryotic organisms were identified by an iterative profile-based searching technique using the Pfam TBCC domain seed alignment (44). A Hidden Markov Model (HMM) was used to search the genome databases of a range of eukaryotic organisms (Table S1) essentially as detailed in (45).

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Supplementary Material

Table S1: Sources and versions of genomic data used in this paper

Supplemental materials are available as part of the online article at http://www.blackwell-synergy.com

References


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