A role for the vesicle-associated tubulin binding protein ARL6 (BBS3) in flagellum extension in *Trypanosoma brucei*

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Abstract

The small GTPase Arl6 is implicated in the ciliopathic human genetic disorder Bardet–Biedl syndrome, acting at primary cilia in recruitment of the octomeric BBSome complex, which is required for specific trafficking events to and from the cilium in eukaryotes. Here we describe functional characterisation of Arl6 in the flagellated model eukaryote *Trypanosoma brucei*, which requires motility for viability. Unlike human Arl6 which has a ciliary localisation, TbARL6 is associated with electron-dense vesicles throughout the cell body following co-translational modification by N-myristoylation. Similar to the related protein ARL-3A in *T. brucei*, modulation of expression of Arl6 by RNA interference does not prevent motility but causes a significant reduction in flagellum length. Tubulin is identified as an Arl6 interacting partner, suggesting that Arl6 may act as an anchor between vesicles and cytoplasmic microtubules. We provide evidence that the interaction between Arl6 and the BBSome is conserved in unicellular eukaryotes. Overexpression of BBS1 leads to translocation of endogenous Arl6 to the site of exogenous BBS1 at the flagellar pocket. Furthermore, a combination of BBS1 overexpression and Arl6 RNAi has a synergistic inhibitory effect on cell growth. Our findings indicate that Arl6 in trypanosomes contributes to flagellum biogenesis, most likely through an interaction with the BBSome.

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1. Introduction

The Arf-like (Arl) proteins are a subfamily of the Arf small GTPases, with diverse roles in vesicle trafficking [1], tubulin folding and microtubule dynamics [2], endosome–lysosome fusion [3] and ciliogenesis [4]. While the functions and effectors of the closely related Arf proteins have been extensively characterised, much less is known about Arl protein interactions and their regulation of fundamental processes within the eukaryotic cell.

Abbreviations: Arf, ADP-ribosylation factor; Arl, ADP-ribosylation factor-like; Arl6, Arl6 interacting protein; BBS, Bardet–Biedl syndrome; BBS1, Bardet–Biedl syndrome 1 protein; BSF, bloodstream form; ConA, Concanavalin A; GEF, guanine nucleotide exchange factor; GPCR, G-protein coupled receptor; HRG4, human retinal gene 4; IFT, intraflagellar transport; ITC, isothermal titration calorimetry; MANT, N-methylanthraniloyl; MAP2, microtubule associated protein 2; NES, nuclear export signal; NLS, nuclear localisation signal; NMT, myristoyl-CoA:protein N-myristoyltransferase; PCF, procyctic form; PCMI, pericentriolar material 1; PFR, paraflagellar rod; PM, plasma membrane; RNAi, RNA interference; RP2, retinitis pigmentosa protein 2; TAP, tandem affinity purification; TiEM, transmission immuno-electron microscopy.

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subunit BBS7 localises transiently to the nucleus where it binds to the polycomb protein RNF2 and may have a direct role in transcriptional regulation [14].

The BBSome in *C. elegans* appears to mediate the IFT machinery, as loss-of-function mutations in *Bbs7* and *Bbs8* orthologues cause abnormalities in localisation and motility of two IFT proteins, OSM-5/Polaris/IFT88 and CHE-11, resulting in truncated cilia and inhibition of chemotaxis [15]. However, in the majority of systems studied to date, the BBSome is believed to be required for specific trafficking events to and from the cilia rather than for ciliogenesis. Two G-protein-coupled receptors (GPCR), somatostatin receptor 3 and melanin-concentrating hormone receptor 1, fail to localise to the cilia of hippocampal neurons in BBSome knockout strains of mice [16]. In contrast, the GPCR dopamine receptor 1, which normally shuttles to and from the cilia, can still be trafficked to neuronal cilia in *Bbs2* and *Bbs4* null mice but fails to exit the cilium in response to receptor ligand in these animals [17]. Insertional mutations of *Bbs1*, *Bbs4* or *Bbs7* in the green alga *Chlamydomonas reinhardtii* produce normal length flagella but phototaxis is disrupted due to a defect in retrograde transport from the flagellum leading to accumulation of signalling proteins [18]. No reports have been published to date on the functions of the BBSome or associated proteins in protozoan pathogeno-organisms.

Here we describe functional characterisation of Arl6 in the parasitic protozoan *Trypanosoma brucei*, as part of a comprehensive study of ARF/ARL proteins as substrates of the validated drug target myristoyl-CoA:N-myristoyltransferase (NMT) [19–22]. We provide experimental evidence that *T. brucei* Arl6 is N-myristoylated as predicted by protein sequence analysis. The trypanosome orthologue of Arl6 is found on small electron-dense vesicles throughout the parasite body. The protein is able to associate with tubulin and might therefore act as a link between the BBSome complex and the intracellular microtubule network. Knockdown of *TbARL6* is not lethal to the parasite but causes a significant decrease in flagellum length, perhaps indicative of defects in protein trafficking to the flagellum. This observation suggests a degree of evolutionary conservation in the role of this protein and possibly of the BBSome complex in trypansomes.

2. Materials and methods

2.1. DNA constructs

All primer sequences are provided in Supplementary Table 1. For protein expression in *Escherichia coli*, a fragment spanning nucleotides 1–570 of the *TbARL6* (TriTrypDB ID: Tb927.8.5060) open reading frame was amplified from genomic DNA using primers ARL6-F1 and ARL6-R1 and cloned into plasmid vector pET-YSBLIC3C (pET28a modified for ligation-independent cloning) [23,24] to produce the construct pET-HisARL6. For tetracycline-inducible overexpression of protein expression in *E. coli* BL21 star and expression of recombinant protein induced with 1 mM IPTG for 4 h at 30 °C. For large-scale protein purification, cells from 5 l of culture were resuspended in 100 ml lysis buffer (300 mM NaCl, 20 mM sodium phosphate pH 7.4, 40 mM imidazole and 1× complete protease inhibitor cocktail, Roche). Cells were lysed by three rounds of sonication then centrifuged at 50,000 g for 40 min at 15 °C. Protein purification was performed using Ni2+ -affinity/size exclusion multi-dimensional liquid chromatography on an ÄKTA Express (GE Healthcare). The clarified lysate was applied at 1 ml/min to a first stage affinity-chromatography column (1 ml Histrap FF, GE Healthcare) equilibrated in 20 mM sodium phosphate, pH 7.5, 300 mM NaCl and 20 mM imidazole (Buffer A), followed by elution with a 1 column volume isocratic gradient of Buffer A containing 0.5 M imidazole (Buffer B) and the UV280 nm absorbing material was peak-fractionated and collected into a system sample loop. The contents of the sample loop were subsequently applied at 1 ml/min to a second stage size exclusion-chromatography column (Superdex200 16/60, GE Healthcare) equilibrated in 20 mM sodium phosphate, pH 7.5, 150 mM NaCl (Buffer C) and then eluted using a 1 column volume isocratic gradient of Buffer C. Fractions (1.8 ml) of UV280 nm absorbing material were collected and analysed for specific protein content by SDS-PAGE.

2.2. Production of recombinant TbARL6 in *E. coli*

The construct pET-HisARL6 was introduced into *E. coli* BL21 star and produced recombinant protein induced with 1 mM IPTG for 4 h at 30 °C. For large-scale protein purification, cells from 5 l of culture were resuspended in 100 ml lysis buffer (300 mM NaCl, 20 mM sodium phosphate pH 7.4, 40 mM imidazole and 1× complete protease inhibitor cocktail, Roche). Cells were lysed by three rounds of sonication then centrifuged at 50,000 g for 40 min at 15 °C. Protein purification was performed using Ni2+ -affinity/size exclusion multi-dimensional liquid chromatography on an ÄKTA Express (GE Healthcare). The clarified lysate was applied at 1 ml/min to a first stage affinity-chromatography column (1 ml Histrap FF, GE Healthcare) equilibrated in 20 mM sodium phosphate, pH 7.5, 300 mM NaCl and 20 mM imidazole (Buffer A), followed by elution with a 1 column volume isocratic gradient of Buffer A containing 0.5 M imidazole (Buffer B) and the UV280 nm absorbing material was peak-fractionated and collected into a system sample loop. The contents of the sample loop were subsequently applied at 1 ml/min to a second stage size exclusion-chromatography column (Superdex200 16/60, GE Healthcare) equilibrated in 20 mM sodium phosphate, pH 7.5, 150 mM NaCl (Buffer C) and then eluted using a 1 column volume isocratic gradient of Buffer C. Fractions (1.8 ml) of UV280 nm absorbing material were collected and analysed for specific protein content by SDS-PAGE.

2.3. Antibody production

Polyclonal antibodies were produced from two rabbits (Eurogentech, 87 day Classic protocol). Antibodies were purified using a 1 ml NHS-activated HP column (GE) coupled to 5 mg of recombinant *TbARL6* protein. Following column equilibration with 10 ml binding buffer (20 mM sodium phosphate pH 7 and 150 mM NaCl), 15 ml rabbit serum was loaded onto the column at 0.3 ml/min. Unbound sample was removed with a 5 ml wash with binding buffer. Elution was then performed using elution buffer at low pH (0.1 M glycine pH 2.7 and 0.5 M NaCl). Fractions of 0.5 ml were collected directly into tubes containing 50 μl 1 M Tris–HCl pH 9.0 and analysed by SDS-PAGE. Peak fractions were pooled and tested by immunoblotting.

2.4. Parasite culture

The *T. brucei brucei* BSF strain Lister 427 was maintained in vitro at 37 °C with 5% CO2 in HMI-9 medium containing 2 μg/ml Geneticin (Invitrogen) as described [26]. The *T. brucei brucei* procyclic strain 449 was maintained in vitro at 26 °C in SDM-79 medium containing 25 μg/ml phleomycin [26]. All culture media contained 10% tetracycline-free fetal bovine serum (Autogen Bioclear).

2.5. Protein analysis

Immunoblotting of total parasite lysates was performed as described previously [22,27]. Primary antibodies used were: rabbit anti-*TbARL6* (1:500 dilution); rabbit anti-*TbNMT* [19] (1:500 dilution); mouse anti-myec (1:2000 dilution, Invitrogen); and mouse monoclonal L13D6 against PFR1/2 (1:500 dilution, a gift from Keith Gull, Sir William Dunn School of Pathology, University of Oxford, UK). For subcellular fractionation, parasites were centrifuged at 800 g for 10 min at 20 °C,
then washed in PBS. For cytosolic (S100) and particulate membrane-containing (P100) fractions, cells were subjected to hypotonic lysis on ice for 1 h in 10 mM Tris pH 7.5 containing 1× Complete protease inhibitor cocktail (Roche). 7.5 μM Pepstatin A and 5 μM EDTA and 1 mM MgSO₄, pH 6.9) containing 1% Nonidet P40, 1× Complete protease inhibitor cocktail (Roche), 7.5 μM Pepstatin A and 5 μM EDTA, incubated on ice for 10 min, then centrifuged at 15,000 g for 15 min at 4 °C. Pellets were either washed twice in PEME and resuspended in Laemmli buffer (NP-40-insoluble fraction containing cytoskeleton) or further extracted in PEME containing 1 M NaCl, 200 μg/ml DNase I, 50 μg/ml RNase A and protease inhibitors as above. Samples were incubated on ice for 10 min, centrifuged as above, salt extraction repeated once, then pellets washed twice in PEME. Pellets were resuspended in Laemmli buffer (NP-40/1 M NaCl-insoluble fraction containing flagella). Total cell lysates and subcellular fractions from the equivalent of 1×10⁷ cells per sample were analysed by immunoblotting and probed with antibodies as above.

2.6. Myristate tagging and fluorescent labelling of TbARL6

Parasites were metabolically labelled by the addition of 100 μM myristic acid or the myristate analogue YnC₁₂[29] to the equivalent of 1×10⁷ cells per sample were analysed by im-

Section 2.5, 1:50), rabbit anti-TbRab5A (1:250, a gift from Mark Field, Department of Pathology, University of Cambridge, UK) and mouse anti-myc (1:250, Invitrogen) with detection using Alexa Fluor 488 and 594 conjugated secondary antibodies (Invitrogen). Staining with FM4-64FX (Invitrogen) and Alexa Fluor 488-conjugated ConA (Invitrogen) were performed as described previously [21,22]. Samples were visualised by confocal microscopy as described previously [21].

Transmission electron microscopy was performed as described [30]. For transmission immuno-electron microscopy, BSF cells in culture medium were diluted in a 1:1 ratio with 8% formaldehyde (w/v)/0.4% glutaraldehyde (w/v) in 100 mM sodium phosphate and incubated for 15 min. Following centrifugation, cells were sequentially treated with 4% formaldehyde (w/v)/0.2% glutaraldehyde (w/v) in 100 mM sodium phosphate buffer for 15 min, 0.1% glycine (w/v) in 100 mM PBS for 20 min, then in 1% gelatin in pure water for 30 min. Cells were then treated with 10% gelatin in pure water and the cell pellet spun down, cut into pieces and infused with 2.3 M sucrose in PBS overnight at 4 °C. Samples were mounted on Leica cryo-holders and frozen in liquid nitrogen before sectioning (approximately 90 nm thickness) on a Leica Ultracut with cryochamber. Sections were transferred to copper grids, then inverted onto drops of liquid on paraffin for immunogold labelling. The grids were incubated in 50 mM glycine/PBS for 5 min, in blocking solution (10% goat serum/PBS) for 20 min, then in primary antibody (rabbit anti-TbARL6 as in Section 2.3 or mouse anti-myc, Invitrogen) diluted 1:25 in blocking solution for 1 h. Samples were washed five times in blocking buffer, then incubated in 10 nM gold conjugated goat anti-rabbit or anti-mouse IgG (Agar Scientific) diluted 1:100–1:400 in blocking buffer for 45 min, then washed five times in blocking buffer. Cells were then fixed in 1% glutaraldehyde in 100 mM sodium phosphate buffer for 10 min and washed five times in pure water. Samples were incubated with 2% methyl cellulose/0.4% uranyl acetate on paraffin placed on an ice-chilled metal block in the dark for 10 min. Viewing was at 120 kV with a Technai 12 BioTwin (FEI) and images were captured with a SIS Megaview III digital camera.

2.8. Expression of myc-tagged ARL6 and BBS1

The constructs pTbARL6Myc, pTbARL6-G2AMYc, pTbARL6-T21NMYc and pTbMycBBS1 (NotI digested) were transfected into mid-log phase T. brucei BSF Lister 427 using the Nucleofector® system as described [31] and transformants selected with 10 μg/ml hygromycin. Expression of tagged protein was induced in stable cell lines by incubating parasites in 1 μg/ml tetracycline for 24 h. Parasite growth was analysed using a Beckman Coulter counter.

2.9. RNA interference of ARL6

The construct pT7TARL6 (NotI digested) was transfected into mid-log phase T. brucei BSF Lister 427 by nucleofection as in Section 2.7 and transformants selected with 2.5 μg/ml phleomycin. Production of TbARL6-specific double-stranded RNA was induced by incubating stable cell lines in 1 μg/ml tetracycline for 24 h. Knockdown of ARL6 expression was monitored by immunoblotting with anti-TbARL6 antibody. Stable ARL6 RNAi cell lines were also transfected with NotI digested pTbMycBBS1, transformants selected by the addition of 10 μg/ml hygromycin and cell lines analysed as in Section 2.7. Cell body and paraglellar rod (PFR) lengths were analysed by performing indirect immunofluorescence on cells treated with 1% Triton X-100 at room temperature for 5 min before staining with mouse monoclonal L13D6 or TAT1 (as in Section 2.6) and detection using Alexa Fluor 488-conjugated goat anti-mouse IgG and IgM (Invitrogen). Cell body (posterior end of the body to the flagellum tip) and PFR lengths (100 per sample) were measured from acquired images using LSM 510 version 3.2 software (Zeiss). Statistical analysis (1 way ANOVA) was performed using GraphPad Prism 4.
2.10. Mouse infections

Swiss outbred (CD-1) mice were infected by i.p. injection with 2 x 10^7 parasites of the parental or TbARL6 RNAi lines as in Section 2.8, grown in the absence or presence of tetracycline for 4 h prior to infection. Mice given tetracycline-treated cells were given doxycycline in drinking water. Parasitaemia was measured at 48 h and 72 h by microscopy analysis of tail-cut blood samples and the experiment terminated at the later timepoint.

2.11. Tandem affinity purification

Log phase T. brucei procyclic strain 449 cells were transfected with BsmI-digested pTbARL6<sub>PTP</sub> by nucleofection as in Section 2.7, replacing one endogenous allele of ARL6 with a C-terminal PTP tagged copy of the ARL6 open reading frame. Transformants were selected with 25 μg/ml hygromycin. Once stable cell lines had been established, tandem affinity chromatography was performed as described previously [32]. Briefly, a crude extract was prepared from 9 L of culture (approximating 1 x 10<sup>11</sup> cells) by resuspension of cells in 15 ml of lysis buffer (150 mM sucrose, 150 mM KCl, 20 mM potassium glutamate, 20 mM Hepes-KOH pH 7.7, 3 mM MgCl<sub>2</sub>, 10 μg/ml leupeptin, and 0.5% DTT) and passing through a French press at 4 °C. The lysate was subjected to IgG affinity chromatography and bound complexes were eluted by incubation with AcTEV protease. The eluate was then subjected to anti-ProTyr affinity chromatography. Proteins were eluted by the addition of EGTA/EDTA to inhibit calcium-dependent binding. The final eluate was concentrated by binding to 10 μl of hydrophobic StrataClean resin (Stratagene) then released into Laemmli SDS-PAGE loading buffer. Samples were separated by SDS-PAGE and stained with Coomassie Blue. Blots were incubated in TBS/1% Tween-20 for 5 min and probed in TBS/0.2% Tween-20 overnight at 4 °C. Blots were washed in TBS/0.2% Tween-20, blocked in 5% milk/TBS/0.2% Tween-20 and probed with mouse monoclonal antibody TAT1 (α-tubulin, 1:10,000 dilution), mouse monoclonal clone 6-11B-1 (acetylated α-tubulin, 1:10,000 dilution, Sigma) or mouse monoclonal TUB 2.1 (β-tubulin, 1:1000, Sigma).

2.12. Tubulin overlay assay

Overlay assays were performed as described [33] using TbARL6 recombinant protein (described in Section 2.2) and negative controls: recombinant human Ras (R9894, Sigma) and bovine serum albumin (New England Biolabs). Proteins (1 μg per lane) were subjected to SDS-PAGE and transferred to nitrocellulose. A duplicate gel was stained with Coomassie Blue. Protein bands were excised and identified by MS/MS analysis of trypsin digests on a 4700 Proteomics Analyser (Applied Biosystems). The mass spectral data were submitted to database searching against the NCBInr database using a locally running copy of the Mascot software (Matrix Science) through a GPS Explorer (Applied Biosystems) interface. The data shown are representative of 4 independent biological replicates.

2.13. Microtubule binding assay

In order to determine if TbARL6 was able to bind to microtubules in vitro, a Microtubule Binding Protein Spin-Down Assay kit (Cytoskeleton, Inc.) was used, following the manufacturer’s instructions. All buffers containing polymerised tubulin were supplemented with 20 μM Taxol. Briefly, 2.5 μg of pure TbARL6 protein was incubated for 30 min at room temperature with a suspension of freshly polymerised microtubules (equivalent of 2 μM of tubulin dimer) in the presence of 10 mM MgCl<sub>2</sub> and 4 mM GDP or GTP. Control samples were also prepared using 2.5 μg of MAP2 or BSA instead of TbARL6, or with the omission of either the test protein or microtubules from the reaction mix. Following incubation, samples were placed onto a layer of Cushion Buffer (80 mM PIPES, pH 7.0, 2 mM MgCl<sub>2</sub>, 1 mM EGTA, and 60% glycerol) and centrifuged at 100,000 g for 40 min. The uppermost layer of supernatant (soluble fraction) and pellet (insoluble fraction) were subjected to SDS-PAGE and stained with Coomassie Blue. In addition, the effects of TbARL6 on tubulin polymerisation in vitro were analysed by measuring turbidity as described previously [34].

2.14. Fluorescence spectroscopy

All samples were diluted in 50 mM KH<sub>2</sub>PO<sub>4</sub> (pH 7.4), with 10 mM MgCl<sub>2</sub> and 12.5 μM unlabelled GDP/GTP (Sigma) or 5 μM N-methylanthranilic acid (mant) labelled GDP/GTP (Invitrogen). Measurements were recorded on a Fluoromax-4 spectrophotometer (Horiba Jobin-Yvon Ltd, Stanmore, UK). Trypophan fluorescence spectra (5 μM protein) were measured at an excitation of 278 nm and emission range of 300–450 nm. Mant fluorescence spectra were measured at an excitation of 350 nm and emission range of 350–550 nm. Kinetic data were collected at an emission of 447 nm (GDP) or 445 nm (GTP) using 5 μM of mant labelled nucleotide and a range of TbARL6 concentrations between 0.2 and 14 μM. Nucleotide exchange was studied by loading 5 μM of protein with 5 μM mant labelled nucleotide for 5 min, then adding a 100-fold excess of unlabelled nucleotide and collecting fluorescence data every 10 s for a further 20 min. K<sub>f</sub> and K<sub>off</sub> values were calculated by nonlinear regression using GraphPad Prism 4. Values shown represent the mean from three independent experiments.

3. Results

3.1. T. brucei ARL6 orthologue

We first identified the T. brucei ARL6 orthologue in a bioinformatic search for downstream targets of the validated drug target myristoyl-CoA:protein N-myristoyltransferase (NMT) [19,22,35]. The constitutively expressed TbARL6 gene encodes a protein of 21 kDa, which is predicted to be N-myristoylated by virtue of the well-conserved N-terminal consensus for N-myristoylation [36]. This is in contrast to ARL6 protein sequences from other species including humans [37] which have an N-terminal glycine residue but are not predicted by available algorithms to be N-myristoylated. Similar to human BBS7, the TbARL6 protein does not contain a nuclear localisation signal (NLS) but does have a leucine-rich nuclear export signal (NES) motif (residues 139–147), using the prediction software programs NLStradamus and NetNES 1.1 [38,39]. Using the same programs, human Arl6 also has a predicted NES (residues 9–13) but no NLS.

3.2. TbARL6 associates with vesicles throughout the parasite body

In order to investigate subcellular localisation in T. brucei, we raised a polyclonal antibody against full length recombinant protein. TbARL6 fused to a C-terminal His tag was readily expressed as a soluble protein in E. coli and purified by affinity purification (Supplementary Fig. 1A). Serum from a rabbit inoculated with recombinant TbARL6 detected a band of approximately 22 kDa on immunoblot of total T. brucei lysates (Supplementary Fig. 1B). Further, only the 22 kDa band was detected in cell lysates by affinity purified antibody (Supplementary Fig. 1B). Although we cannot rule out the possibility that other ARF family proteins could be detected by the antibody, the most closely related protein to ARL6 encoded by the T. brucei genome is the Golgi protein ARL1, which shares only 37% identity [22], while TbARF1 shares less than 30% identity with TbARL6.

Immunofluorescence assays using the antibody showed that ARL6 is present in a punctate distribution throughout the cell body in both bloodstream form (BSF) and insect procyclic form (PCF) T. brucei and has the same localisation in non-dividing and dividing cells (Fig. 1A).
Some staining was observed at the periphery of the nucleus, consistent with the presence of a NES motif in the protein. There was no obvious co-localisation between TbARL6 and the early endosomes, as visualised using an antibody against T. brucei Rab5 (Supplementary Fig. 2B). This may represent either mislocalisation due to the epitope tag or slower trafficking of the over-expressed protein. TiEM images show that the tagged protein is associated with circular structures (Supplementary Fig. 3) similar in morphology to the 200 nm diameter structures found adjacent to a minor population of the endogenous protein (Fig. 1D and data not shown). In contrast, a G2A mutant form of the protein lacking the N-terminal glycine residue (the myristate acceptor) mis-localised to the plasma membrane (Supplementary Fig. 2B). This mutant isoform was readily expressed even in the absence of tetracycline, showing some leakiness of the inducible promoter system. Unlike the wild-type, the G2A mutant protein appeared to be a target for proteolysis at its new site in the cell, as shown by immunoblotting (Supplementary Fig. 2C). Another mutant form of TbARL6, T21N (a critical residue for nucleotide binding found at the P-loop) was found to be unstable and could not be captured by TiEM. This mutant isoform was readily expressed even in the absence of tetracycline, showing some leakiness of the inducible promoter system. Unlike the wild-type, the T21N mutant protein appeared to be a target for proteolysis at its new site in the cell, as shown by immunoblotting (Supplementary Fig. 2C). Another mutant form of TbARL6, T21N (a critical residue for nucleotide binding found at the P-loop) localised partially at the plasma membrane and partially in a punctate distribution, a subpopulation of which co-localises with Rab5 (Supplementary Fig. 2B).

3.3. Evidence that TbARL6 is N-myristoylated

We tested the ability of TbARL6 to be N-myristoylated by metabolic labelling of parasites with the alkynyl-myristate YnC12 [40], which acts as a myristic acid analogue in cells and can be captured by copper-catalysed click chemistry in combination with fluorescent probes [29]. TbARL6 was captured from parasite lysates by immunoprecipitation and labelling of YnC12-tagged proteins was carried out on-bead via a click chemistry reaction between the alkynyl functionality and fluorescent rebinding (Supplementary Fig. 2A). A number of background bands were observed, particularly in the PCF samples. However, specific bands were observed for TbARL6 in YnC12-labelled PCF and BSF lysates following immunoprecipitation: the 21 kDa band and an additional higher molecular weight band which likely corresponds to the protein carrying the Az-TB label (mw of ~1000), as shown in

Based cytoskeleton but further extraction with salt solubilises the subpellicular microtubules, therefore the NP40/1M NaCl-insoluble fraction contains only the highly insoluble flagella. TbARL6 protein was found in the membrane-containing P100 fraction of BSF cell lysate but not in the cytosolic S100 fraction, NP40-insoluble or NP40/1M NaCl-insoluble fractions (Fig. 1B). This suggests that the protein associates with cellular membranes but is not an integral part of the cytoskeleton, consistent with its reported absence from the flagellar proteome [28]. Intracellular localisation was studied at high resolution by transmission immuno-electron microscopy (TiEM) (Fig. 1C–F). Endogenous TbARL6 was detected throughout the cell, particularly in association with electron-dense vesicles of less than 100 nm diameter (Fig. 1C, E, and F) and to a lesser extent with larger structures of approximately 200 nm diameter (Fig. 1D). Negligible levels of gold labelling were detected in negative-control samples without primary antibody (Supplementary Fig. 3A).

When TbARL6 was overexpressed with a C-terminal myc tag, a punctate pattern was also observed by immunofluorescence, although the foci were larger for endogenous TbARL6 (Supplementary Fig. 2B, top panel). Unlike endogenous TbARL6, the myc-tagged protein showed some co-localisation with the early endosomes as visualised using an antibody against T. brucei Rab5 (Supplementary Fig. 2B). This may represent either mislocalisation due to the epitope tag or slower trafficking of the over-expressed protein. TiEM images show that the tagged protein is associated with circular structures (Supplementary Fig. 3) similar in morphology to the 200 nm diameter structures found adjacent to a minor population of the endogenous protein (Fig. 1D and data not shown). In contrast, a G2A mutant form of the protein lacking the N-terminal glycine residue (the myristate acceptor) mis-localised to the plasma membrane (Supplementary Fig. 2B). This mutant isoform was readily expressed even in the absence of tetracycline, showing some leakiness of the inducible promoter system. Unlike the wild-type, the G2A mutant protein appeared to be a target for proteolysis at its new site in the cell, as shown by immunoblotting (Supplementary Fig. 2C). Another mutant form of TbARL6, T21N (a critical residue for nucleotide binding found at the P-loop) localised partially at the plasma membrane and partially in a punctate distribution, a subpopulation of which co-localises with Rab5 (Supplementary Fig. 2B).
previous analysis of cholesterol-modified Sonic hedgehog protein [41]. Comparison of the relative intensities of the two bands gives an approximate tagging/labelling efficiency for TbARL6 of 50% using this methodology.

![Image](image_url)

**Fig. 2.** Myristate tagging and fluorescent labelling to show N-myristoylation of TbARL6. (A–C) T. brucei insect procyclic form (PCF) line 449 and bloodstream form (BSF) line Lister 427 were metabolically labelled with myristic acid or the myristate analogue YnC12. Immunoprecipitation was performed on cell lysates by the addition of rabbit anti-TbARL6 and Protein A Sepharose beads, and captured proteins were subjected to on-bead click labelling with the fluorophore TAMRA. Samples were eluted, separated by SDS-PAGE, transferred to PVDF membrane and fluorescence was detected using an Ettan DIGE scanner (GE Healthcare). In-gel (A) and in-blot (B) fluorescence are shown for PCF (left panels) and BSF (right panels) lysates. (C) Immunoblot was probed with anti-TbARL6. 1, 2, Total lysate from myristic acid-labelled and YnC12-labelled cells respectively (100%), 3, 4, Supernatant following immunoprecipitation for myristic-acid labelled and YnC12-labelled cells respectively (20%).
by electron microscopy at 24 h post-induction [Supplementary Fig. 4], except in a small minority of cells (<5%) which displayed axoneme defects and a swollen flagellar pocket [Supplementary Fig. 4C] compared to an estimated <1% in uninduced cells.

The trypanosome flagellum is composed of a canonical 9 + 2 microtubular axoneme and the paraflagellar rod (PFR), a filamentous structure which runs parallel to the axoneme [43]. In order to analyse the effects of TbARL6 knockdown on the flagellum, the length of the PFR was measured in non-dividing cells (containing one nucleus and one kinetoplast, 1N1K) stained with DAPI and anti-PFR antibody prior to and following the induction of RNAi. The PFR extends to the distal tip of the flagellum in induced cells and therefore can be used as a reliable marker of flagellum length (Fig. 3C). Body length was also measured in non-dividing cells stained with DAPI and anti-α tubulin (Fig. 3D). There was a modest but highly significant decrease in PFR length at 24 and 48 h post-induction compared to wild-type and to uninduced cells (p = <0.001) (Fig. 3E). Cells with shorter flagella remained motile, which is consistent with the observed normal growth described above. Motility is known to be essential for viability of this parasite stage [28] and any defects in this process would have resulted in cell death. We also observed a significant decrease in PFR length in uninduced RNAi cells compared to the parental line (p = <0.001), which may be due to leaky expression of double-stranded RNA in the absence of tetracycline, as described in previous studies [44]. The mean PFR length was restored at later time points, again suggesting the emergence of escape mutants. Conversely, body length (including the region of the flagellum extending beyond the anterior tip of the body) was only marginally decreased in induced cells, with statistical significance at 48h compared to wild-type cells (p = <0.01) (Fig. 3F). Our data therefore suggest that a moderate reduction in TbARL6 protein level resulted in a significant decrease in flagellum length but otherwise did not seem to affect cell growth or morphology.

We also tested the ability of BSF cells to establish an infection in a mouse in vivo model (data not shown). TbARL6 RNAi was induced with tetracycline for 4 h before infection and levels of parasitaemia checked at 48 and 72 h post-infection. Parasite concentrations in the blood were greater than 5×10⁷/ml in mice infected with either the parental or RNAi line by 72 h, indicating that parasites with a modest reduction in ARL6 due to RNAi retain the ability to mount an infection in vivo.

Fig. 4. Identification of tubulin as an interacting partner of T. brucei ARL6. (A) PTP purification of TbARL6. Samples were separated by SDS-PAGE and stained with Coomassie. 1, Input material (0.05% of fraction loaded), 2, 3, 4, Flowthrough and washes of IgG Sepharose column (0.05%, 0.2%, and 0.13%), 5 and 8, TEV protease eluate (1%), 6, Flowthrough of the anti-ProtC matrix (0.3%), 7, Flowthrough of StrataClean resin (2.9%), 9, Final eluate (100%). The final eluted fraction contains three major protein bands (marked by asterisks) which were identified as protein C-tagged TbARL6 (23 kDa), α/β-tubulin (50 kDa) and a keratin contaminant (~65 kDa). (B) Tubulin overlay assay. Purified recombinant proteins (1 μg per lane) were separated by SDS-PAGE and either stained with Coomassie (Total) or transferred to PVDF (remaining panels). 1, Human Ras (R9894, Sigma), 2, TbARL6, 3, bovine serum albumin (New England Biolabs). Blots were incubated in tubulin solution overnight, then probed with mouse monoclonal antibody TAT1 against α-tubulin (α-Tub), monoclonal antibody clone 6-11B-1 which specifically recognises acetylated α-tubulin (Acet α-Tub) or monoclonal antibody TUB 2.1 against β-tubulin (β-Tub). Control immunoblots were processed with TAT1 antibody but without tubulin (−Tub + Ab) or without primary antibody (+Tub − Ab). (C) Microtubule binding protein assay. Purified proteins were incubated with a microtubule preparation before ultracentrifugation. Soluble (S) and pellet (P) fractions were subjected to SDS-polyacrylamide electrophoresis and stained with Coomassie. MT, microtubules. MAP2, Microtubule associated protein 2 (280 kDa, positive control), BSA, bovine serum albumin (68 kDa, negative control). Corresponding protein marker positions are shown (kDa).
3.5. Attempts to make a TbARL6 knockout line

As RNAi knockdown was incomplete, we attempted to generate a BSF knockout line in which both alleles of the ARL6 gene were replaced by antibiotic resistance selection markers. Parasite lines with one deleted allele were readily produced but no double knockout clones could be generated despite repeated transfections. Of a potential 192 clones (8 × 24 well plates) only two clones were successfully cultivated, a 50-fold reduction in efficiency compared to transfections to delete the first allele of the gene. Both of these clones encoded the

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**Fig. 5. T. brucei ARL6 nucleotide interactions.** Purified recombinant TbARL6 was used at 5 μM in 50 mM phosphate buffer unless otherwise stated. (A) Tryptophan fluorescence emission spectra of TbARL6 in the presence and absence of 6 M guanidine hydrochloride. (B) Tryptophan fluorescence emission spectra of TbARL6 in phosphate buffer and 10 mM MgCl2 in the presence and absence of 12.5 μM unlabelled GDP or GTP. (C, D) Mant fluorescence emission spectra of mantGDP (C) and mantGTP (D) in the presence and absence of TbARL6 protein. (E, F) Dissociation constants (KD) for TbARL6 and mantGDP (E) or mantGTP (F) were calculated by non-linear regression (one site specific binding) from data obtained at a single emission wavelength (GDP, 445 nm, GTP, 447 nm) with a fixed concentration of nucleotide and a range of protein concentrations. (G, H) Guanine nucleotide exchange by TbARL6. Protein was prebound to mantGDP (G) or mantGTP (H), then mixed at 300 s with a large excess of unlabelled nucleotide. The decrease in fluorescence was recorded over time and dissociation rate constants (k_diss) calculated by non-linear regression (one phase exponential decay). Controls represent protein and mant-labelled nucleotide without the addition of unlabelled nucleotide. All data are representative of at least 3 independent experiments.
two antibiotic resistance genes but also retained a copy of the ARL6 gene, suggesting gene amplification events had occurred in these cells. These results are in direct contrast to those obtained in our recent study on a unrelated trypanosome gene DIP13. Both alleles of this gene were successfully deleted from the conditional knockout line in which both alleles of the cies observed for the second and second transfections and no growth defects was found in DIP13 null cells [45].

As an alternative approach, attempts were made to produce a conditional knockout line in which both alleles of the ARL6 gene were deleted in cells expressing TbARL6MYC under the control of a tetracycline-inducible promoter. As before, single allele deletion was achieved but no double knockout clones could be produced. Southern blotting and PCR showed that the TbARL6 locus was still present following correct integration of two antibiotic resistance cassettes in 8 independent clones (data not shown), again indicating that gene amplification had occurred in these parasites. Therefore, while moderate knockdown of the protein has no effect on cell growth, attempts to delete both alleles of the gene have so far been unsuccessful, resulting in either transfection failure or genetic rearrangements. When put into context with current dogma in kinetoplastid research, this suggests that the TbARL6 gene could in fact be essential for viability and that the myc-tagged protein used for complementation (which localises differently in the cell) is not sufficient to rescue cells. Further studies are therefore required to determine whether TbARL6 is essential for viability in the parasite.

3.6. TbARL6 binds to tubulin

In order to find the binding partners of TbARL6, we used a modified version of tandem affinity purification (TAP), which involves fusion of the protein to Protein A and Protein C epitopes (termed a TAP tag) [32] to enable highly-specific purification of protein complexes from crude cell lysate. A construct encoding C-terminal TAP-tagged ARL6 was inserted into the endogenous ARL6 locus of T. brucei procyclic parasites to produce cells expressing tagged protein at endogenous levels. This reduced the possibility of erroneous interactions due to overexpression but necessitated the use of large volumes of cells in order to produce detectable amounts of final product. Following a series of affinity purification steps, three proteins were detected in the final eluate (Fig. 4A). These were identified by mass spectrometry as TbARL6, α/β-tubulin and a keratin contaminant, with identical results produced in four independent biological replicates. As tubulin is highly abundant in trypanosomes and often found as a contaminant in protein interaction studies, additional studies were required to validate this protein as a binding partner of TbARL6.

This was investigated further using a tubulin overlay assay (Fig. 4B). Immunoblots of N-terminal His-tagged recombinant ARL6 (which cannot be N-myristoylated) and controls were incubated in porcine tubulin solution before probing with anti-tubulin antibodies. The presence of a 22 kDa band in the His/ARL6 lane (Fig. 4B) clearly showed binding of this protein to α-tubulin in vitro, whereas no signal was produced in lanes containing either BSA or the small GTPase Ras. The same band was also detected when blots were probed with anti-tubulin antibodies which cannot be

By directly binding to the BBS1 subunit[11,53]. To investigate whether this interaction is conserved in kinetoplastids, an N-terminal myc

To summarise these findings, our data indicate that TbARL6 is a tubulin-binding protein (with evidence from a range of techniques) and has the ability to bind both unpolymerised tubulin and assembled microtubules (albeit transiently or at lower affinity than the MAP2 positive control). Further, these interactions are not dependent on N-myristoylation or on specific nucleotide binding conformation of the small GTPase.

We also tested the ability of TbARL6 to alter the in vitro polymerisation of tubulin in the presence of taxol, monitoring the increase in turbidity as microtubules are produced, as described previously [34]. No significant differences were seen in the presence or absence of recombinant TbARL6 (data not shown). Thus, while our data strongly suggest interaction between TbARL6 and tubulin, the functional relevance of this interaction in vivo will require further investigation.

3.7. TbARL6 nucleotide interactions

In addition to identifying protein binding partners of TbARL6, we studied the nucleotide interactions of recombinant protein. To test the folding integrity of purified protein, emission spectra were recorded for TbARL6-His either in phosphate buffer or following denaturation in 6 M guanidinium hydrochloride. Denaturation caused an increase in tryptophan fluorescence and a red shift of the emission maximum from 350 nm to 356 nm (Fig. 5A). This suggests that the protein has a hydrophobic core and is capable of folding in phosphate buffer in the absence of Mg2+ or nucleotide. TbARL6 is therefore likely to be stable in a nucleotide-free state. Intensity of the intrinsic fluorescence of TbARL6-His decreased in the presence of GDP/GTP and 10 mM MgCl2 (Fig. 5B), with GTP exerting the greatest effect. No change in fluorescence was seen in the absence of magnesium (data not shown), confirming that nucleotide binding, as for other Arf GTPases, is Mg2+-dependent.

TbARL6-His showed rapid binding with mant-labelled GDP and GTP, causing an increase in the fluorescence intensity of the nucleotides (Fig. 5C and D). Fluorescence reached a plateau within 3 min (data not shown) and remained relatively constant when measured at 10 s intervals for 20 min (control samples in Fig. 5G and H). Equilibrium binding constants were determined by titrating mant-labelled nucleotides with increasing concentrations of protein and plotting the change in fluorescence (Fig. 5E and F). The dissociation constant (Kd) value of each interaction was calculated by non-linear regression (one site specific binding) and compared with known values for other small GTPases (Table 1) [46–52]. The Kd of TbARL6-His with GDP is 4.35 μM (± 1.12) and with GTP is 13.71 μM (± 3.52). We attempted to corroborate these data with unlabelled nucleotides by isothermal titration calorimetry (ITC) but the protein precipitated in response to the vigorous agitation required.

Dissociation rate constants (κdiss) were measured by loading TbARL6-His with mant-labelled nucleotide, then adding a 100-fold excess of unlabelled nucleotide and recording the decrease in fluorescence as the mant-labelled molecules were displaced (Fig. 5G and H). For both GDP and GTP, the resultant curves fitted one-phase rather than two-phase exponential decay, indicating a single dissociation event. Similar values were produced when displacing the mant nucleotides with either GDP or GTP (Fig. 5G and H and Table 1). The κdiss for mantGDP was found to be 0.0076 s−1 (± 0.0014) when displaced with unlabelled GDP and 0.0093 s−1 (± 0.0013) when displaced with unlabelled GTP. The equivalent values for mantGTP were 0.0047 (± 0.0006) with GDP and 0.0041 (± 0.0007) with GTP.

3.8. Interaction between TbARL6 and the BBSome

Human Arl6 is now believed to interact with the BBSome complex by directly binding to the BBS1 subunit[11,53]. To investigate whether this interaction is conserved in kinetoplastids, an N-terminal myc
Table 1

Guanine nucleotide binding and exchange kinetics of TbARL6. Dissociation constants (Kd) and dissociation rate constants (κdiss) for the binding of N-terminal His-tagged T. brucei ARL6 to mant guanine nucleotides (mGDP and mGTP), compared to values obtained previously for other small GTPases using either mant or radioactive nucleotides (shown in separate columns).

| Protein     | Kd (μM)  | κdiss (s⁻¹) |
|-------------|----------|-------------|
| TbARL6      | 4.354    | 13.710      |
| Arf2        | 0.492    | 0.173       |
| Arf3        | 0.240    | 48.000      |
| Dynamin     | 100.000  | 2.500       |
| Rab3A       | 0.063    | 0.406       |
| Cdc42       | 0.590    | 0.600       |
| RhoA        | 0.480    | 0.400       |
| Rac1        | 0.420    | 0.200       |

Fig. 6. Association of TbARL6 with the BBSome. (A) Immunofluorescence analysis of T. brucei BSF transfected line 427/pMYCBBS1 grown in the presence of tetracycline for 24 h. Cells were incubated on ice with the lipophilic dye FM4-64 (red) to stain the flagellar pocket prior to fixation, then probed with mouse anti-myc (green) and co-stained with DAPI (blue). Bar, 5 μm. (B) Cells as in (A) above were grown in the absence (−Tet) or presence (+Tet) of tetracycline for 72 h, then probed with mouse anti-myc (green) and rabbit anti-ARL6 (red) and co-stained with DAPI. Bar, 5 μm. (C) Total cell lysates (1 × 10⁶ cells/lane) from BSF parental line Lister 427 (wt) and cells 427/pMYCBBS1 and 427/p2T7ARL6/pMYCBBS1 in the absence (−) or presence (+) of tetracycline were immunoblotted and probed with mouse anti-myc and anti-NMT to monitor equal sample loading. M, (Ref.) Myc-cyclic GMP phosphodiesterase, J Biol Chem, 269 (1994) 28130–28135. **R. Price et al. / Biochimica et Biophysica Acta 1823 (2012) 1178–1191**
on nucleotide binding status. As TbARL6 does not contain the equivalent of the highly conserved Cln71 (discussed in Section 4.6), further investigation of relevant residues will be required before analysis of a GTP-locked mutant is feasible. A recent study suggested that the interaction between Arl6 and BBS1 may be regulated by the Arl6 N-terminus [53]. Further studies are required to determine if N-myristoylated TbARL6 retains the ability to bind BBS1. Moreover, the full complement of Arl6 and TbARL6 interacting proteins and their roles in protein trafficking remain to be fully characterised. An earlier study used yeast two-hybrid screening to identify six interacting partners of human Arl6, including SEC61beta, a subunit of the heterotrimeric protein conducting channel SEC61p and a molecule of unknown function, Arl6 interacting protein 1 (Arl6ip) [54]. However, there are no clear orthologues of any of these interacting proteins in trypanosomes that are identifiable by computational analysis.

4.2. Loss of function of proteins associated with Bardet–Biedl syndrome

A range of model systems have been used to study loss of function of the Bardet–Biedl associated proteins. With few exceptions, disruption of cilium function is the consistent phenotype regardless of which BBS gene is targeted, leading to specific trafficking or signalling defects and downstream events depending on the model organism studied. Unlike many of the IFT components, loss of BBS gene function does not appear to be lethal [18,55]. Also, the assembly of cilia/flagella is still able to occur in the absence of specific subunits of the BBSome complex in all eukaryotic systems studied to date, with the notable exception of flagellum production in the spermatooza of knockout mice [18,53,56–60]. Abnormalities in rhodopsin trafficking in BBS knockout mice leads to degeneration of photoreceptor cells [56,61,62], a phenotype which can be prevented in Bbs4 knockout mice by adeno-viral delivery of the Bbs4 gene [63]. Bbs3 (Arl6) knockout mice display many of the characteristic features of Bardet–Biedl syndrome, including retinal degeneration and male infertility but also have unique phenotypes such as severe hydrocephalus, suggesting that the protein may have roles in addition to BBSome recruitment [53]. Knockdown of BBS genes (including BBS3) in zebrafish causes delayed retrograde melanosome transport and disruption of the Kupffer’s Vesicle, a fluid-filled ciliated epithelial sac found in embryonic fish [58–60]. Changes in Sonic Hedgehog (shh) expression are also observed, leading to alterations in fin bud patterning [59]. Conversely, specific knockdown of an alternative transcript of BBS3 termed BBS3L does not produce this phenotype, instead resulting in retinal defects [64], therefore suggesting that these two Arl6 variants have distinct non-overlapping functions. BBSome insertional mutants in Chlamydomonas reinhardtii can assemble normal length motile flagella and have normal IFT particle structure and movement but display a defect in phototaxis. This is due to an intraflagellar accumulation of four putative signalling proteins, indicating that the BBSome has a role in export of signalling proteins from the flagellum [18].

In trypanosomes, we found that a modest knockdown of ARL6 does not kill the parasite but causes a significant decrease in flagellum length, suggesting some evolutionary conservation in the function of this protein. Flagellum motility and maintenance of flagellar pocket integrity are essential for viability of the host bloodstream form (BSF) of the parasite and disruption of these by RNA interference invariably leads to cell death [28,65,66]. Partial knockdown of TbARL6 expression did not cause widespread cell death, unlike RNAi knockdown of the related small GTPases ARL1 and ARF1 in T. brucei [21,22]. Nevertheless, despite repeated attempts, we were unable to generate a double knock out null line and further studies will be required to confirm the full effects of ARL6 loss of function in this organism. Further, we found that overexpressed BBS1 in T. brucei localises to the flagellar pocket and results in a translocation of endogenous TbARL6 to this site. A combination of BBS1 overexpression and knockdown of ARL6 has a synergistic inhibitory effect on cell growth, indicating a close link between ARL6 and the BBSome in trypanosomes.

4.3. The BBSome and Rab8

BBS1 is able to directly bind Rab8 [12] and a model has been proposed in which the BBSome may be able to regulate activation of the multifunctional small GTPase Rab8, leading to the sorting and transport of cargo to the cilium [12,67] and potentially across the periciliary barrier for conveyance to the IFT system [11]. However, while Rab8 is found in almost all eukaryotes [68], the BBSome is restricted to ciliated/flagellated organisms [9]. Moreover, flagellated protozoan parasites have conserved BBSome subunits but lack a member of the Rab8/Rab10/Rab13 clade [9,69], indicating that there is only a partial overlap in functional mechanism between the BBSome and Rab8, and that Rab8-independent BBSome processes must exist, at least in protozoa.

4.4. Localisation of ARL6 and the BBSome

Human Arl6 localises to the distal end of basal bodies and to primary cilium membranes [11,70] and the orthologue of this protein has been detected in the flagellar proteome of C. reinhardtii [71] but not in similar studies of T. brucei procyclic and BSF parasites [28,45]. Human BBSome components have been detected in two pools, localising to centriolar satellites and inside the primary cilium. Affinity purification techniques have shown that the subunit BBS4 interacts directly with PCM1, the core component of the centriolar satellites, which may act as a chaperone to prevent coat assembly before it is required. In contrast, the BBSome found within the primary cilium is dissociated from PCM1 and associates instead with GTP-bound Rab8 [11,12]. PCM1 and centriolar satellites are not conserved in protozoans, therefore a different mechanism may be required for safe transport of the BBSome components from their site of synthesis (as yet unidentified) to the flagellar pocket or other locations in the cell. Our data suggest that endogenous TbARL6 is associated with small vesicles throughout the parasite body and we speculate that TbARL6 could mediate delivery of the BBSome components on these vesicles.

4.5. TbARL6 localisation and tubulin association

Our protein interaction studies identified tubulin as a potential binding partner of TbARL6, independent of N-myristoylation or nucleotide binding status. This suggests a possible association of ARL6-positive vesicles with cytoplasmic microtubules to facilitate transport to the flagellar pocket, although these structures have not yet been visualised in T. brucei [72]. No significant co-localisation was found between endogenous TbARL6 and the endosomes but myc-tagged ARL6 partially co-localises with the early endosomal marker Rab5 in parasites, signifying either mislocalisation due to the epitope tag or abnormal accumulation of overexpressed protein at this stage due to saturation of the transport mechanism. Although the N-terminus is known to be critical for N-myristoylated Arf/Arl function, blocking the C-terminus by epitope-tagging may also be detrimental to protein function [27,73]. It is interesting to note that the G2A mutant of TbARL6 shows a subcellular localisation consistent with binding to the subpellicular microtubules at the plasma membrane. We hypothesise that N-myristoylation could provide the switch required for additional interactions to occur which limit potential association with tubulin and thus enable TbARL6 to reach its correct vesicular localisation. In the absence of this modification, TbARL6 may instead accumulate at a “default” localisation in the region with the highest tubulin concentration, that of the subpellicular cortex underlying the plasma membrane [74]. Endogenous TbARL6 has not been detected in detergent-insoluble cell extracts (which contain the cytoskeleton), showing that any association with microtubules is probably transient. An intriguing alternative hypothesis to direct microtubule binding is that tubulin may be a cargo molecule of ARL6-positive vesicles. It has
been hypothesised that all proteins destined for the ciliary membrane and axoneme, including tubulin, may be transported via association with Golgi-derived vesicles which exocytose with the basal body and then the ciliary membrane [75]. Tubulin has been identified in vesicles destined for the cilium and may in fact peripherally associate with the outside of vesicles prior to exocytosis with the ciliary membrane [75], in a similar mechanism to that described for F-actin delivery on endosome-derived vesicles to ingressing cleavage furrows during cytokinesis [76].

Two other proteins in the Arf family, Arl7 and Gic/Arl8, have been shown to date to bind directly with tubulin [77,78]. Arl7 binds to α-tubulin [77], shows upregulated expression in response to cholesterol loading and has been implicated in transport of apolipoprotein A-dependent cholesterol export [77,79]. Gic/Arl8 is essential for equal segregation of chromosomes in mammalian cells and Drosophila [78]. Like T. brucei Arl6, Gic is able to bind to both α- and β-tubulin, independent of the nucleotide-binding status of the GTPase [78]. The ability to bind tubulin is retained in mutant forms of Gic1 (ArlB8) lacking the Switch 1 or Switch II effector binding domains, suggesting that tubulin is a binding partner rather than an effector or regulator molecule. Other members of the Arf family, Arl2 and Arl3, have roles in microtubule-related processes but these are not believed to require a direct interaction with tubulin. GDP-bound Arl2 binds to tubulin cofactor D, preventing the disassembly of microtubules [2]. The orthologue of Arl2 in T. brucei is required for cleavage furrow formation during cytokinesis and knockdown causes the production of multinucleated cells [27]. Arl3 is closely related to Arl2 and is able to bind all Arf2 effector proteins with the exception of cofactor D [80]. Arl3 has an additional set of interacting partners, human retinal gene 4 (HRG4/UNC119) and retinitis pigmentosa 2 (RP2), with which it can form a ternary complex [81]. Similar to Arl6, Arl3 has been detected on primary cilium in human photoreceptor cells [82,83] and Arl3 knockout mice develop photoreceptor degeneration [84]. Kinetoplastids have three putative Arl3 orthologues [22], of which one, termed ARL-3A, has been studied in detail. Disruption of ARL-3A function in Leishmania by expression of either GFP-locked or GDP-locked forms and in T. brucei by RNAi knockdown causes significant shortening of flagella without loss of motility [85–87], as found here with knockdown of T. brucei ARL6. ARL-3A in Leishmania amazonensis is found in a punctate pattern [87] and may therefore have a vesicular localisation similar to that for TbARL6. Recent studies have shown that Arf proteins can act in synergy to produce an activation cascade, as shown previously for Rab8 [88], with a minor population of GTP-bound activated Arf6 or Arl4 binding to the PH domain of the Arno GEF to enhance binding of Arf1 to the Sec7 domain, thus triggering Arf1 activation [89]. Our data suggest that T. brucei ARL6 has spatial and functional similarities to ARL-3A and there may conceivably be an association between the two proteins, either in function or in triggering activation.

4.6. Guanine nucleotide binding properties of TbARL6

As a small GTPase, the nucleotide binding properties of Arf6 will almost certainly be critical for its function. As a reflection of this, many of the Arf6 mutations described in BBS patients are missense mutations of the highly conserved amino acids required for guanine nucleotide association [90]. Expression of GTP- and GDP-locked variants of Arf6 in a human cell line does not alter cell growth or gross morphology but significantly changes the proportion of cells possessing cilia and the length of assembled cilia. Overexpression of wild-type Arf6 causes an increase in Wnt signalling, as measured by the level of β-catenin transcriptional activity, but expression of nucleotide-locked variants of Arf6 has no effect on this response [70]. The amino acid residues Thr 31 and Gln 71 (Arf1 numbering) are highly conserved in nearly all Ras-like small GTPases, with critical roles in GTP binding and hydrolysis, respectively. Sequence alignment of putative Arf6 orthologues (Supplementary Fig. 5) shows that while Thr 31 is conserved in the kinetoplastid orthologues (except in T. cruzi), the equivalent residue of Gln 71 in Arf1, is substituted by alanine in the parasite Arf6 sequences (residue 63). Gln 71 is also absent from the larger GTase dynamin, which has the ability to self-assemble into higher order structures, resulting in active-site conformational changes which in turn stimulate GTP hydrolysis [91].

In light of this information, we investigated the ability of recombinant His-tagged TbARL6 to bind nucleotides using tryptophan and fluorescence spectroscopy. There are two tryptophan residues in TbARL6, one of which resides in the Switch II effector binding domain. Our data indicate that nucleotide binding causes a local conformational change in this region which partially quenches fluorescence, as shown for Rab5 [92]. Previous studies have shown that Arf1 requires the presence of lipids and Mg<sup>2+</sup> for nucleotide binding and is unstable in vitro in a nucleotide-free state. Arf1 has a very high affinity for guanine nucleotides (nanomolar range) and the exchange rate is extremely slow [48]. In contrast, the related protein Arl2 is stable in a nucleotide-free state and has low affinity for nucleotides due partly to a rapid exchange rate [46]. We conclude from our data that TbARL6—like protein is stable in the absence of nucleotides and is able to bind guanine nucleotides with low affinity in the absence of lipids, with relatively rapid exchange rates. Therefore, the protein appears to be similar in terms of nucleotide binding characteristics to Arf2. Both TbARL6 and dynamin lack the equivalent residue of Gln 71 in Arf1, an amino acid shown to be crucial for GTP hydrolysis. As both of these proteins also exhibit low affinity for GTP, it is possible that this residue may have a role in modulating nucleotide affinity in small GTPases. Alternatively, the His tag may exert some inhibitory effects or TbARL6 may require other factors such as membrane lipids for optimal nucleotide association.

4.7. Concluding remarks

In conclusion, we have investigated the functions of the Arf6 orthologue in T. brucei, a divergent lower eukaryote which encodes the BBSome components but lacks a Rab8 orthologue. It is yet to be proven that the BBSome has a role in transporting cargo to the flagellum in kinetoplastid parasites but the functional analysis of T. brucei ARL6 presented here suggests that this mechanism is fundamentally conserved and required for flagellum extension. We propose that there is a conserved functional association between the BBSome and ARL6 in T. brucei which is required for flagellar extension in a Rab8 null background. Remarkable recent evidence has shown that the scope of the intraflagellar transport system is not restricted to that of transport within the flagellum. One of the IFT components, IFT20, has a dynamic localisation in photoreceptor cells, moving between the Golgi and the primary cilium, therefore providing a potential transport link between the endomembrane system and the IFT machinery [93]. Considering the widespread distribution of Arf6 in trypanosomes, we cannot rule out the possibility that this protein may also have a role linking transport between multiple sites in the parasite. Further studies will be required to elucidate the exact molecular mechanisms of Arf6 and related BBSome functions in trypanosomes and other eukaryotic cells.

Supplementary data related to this article can be found online at http://dx.doi.org/10.1016/j.bbamcr.2012.05.007.

Acknowledgements

We acknowledge contributions from the following colleagues: Sam Alsford and David Horn for myc-tag vectors; Arthur Günzl for the modified TAP vector; George Cross and Doug LaCount for parasite strains and the p2T7TI vector; Mark Field and Keith Gull for primary antibodies. Recombinant His-tagged TbARL6 protein was expressed and purified by the HiTel laboratory in the Department of Biology.
Technology Facility at the University of York, UK. We also thank Mark Field, Glen Hemsworth, Karen Hodgkinson, Andrew Leech, Lorna MacLean, Graeme Park, Paul Robinson, Joel Rosenbaum, Berni Strongtharm and Daniel Ungar for technical advice and helpful discussions. This research was funded by the Wellcome Trust (grant no. 077503).

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