Eukaryotic Cell	A SAS-6-Like Protein Suggests that the Toxoplasma Conoid Complex Evolved from Flagellar Components
	Jessica Cruz de Leon, Nicole Scheumann, Wandy Beatty, Josh R. Beck, Johnson Q. Tran, Candace Yau, Peter J. Bradley, Keith Gull, Bill Wickstead and Naomi S. Morrissette <i>Eukaryotic Cell</i> 2013, 12(7):1009. DOI: 10.1128/EC.00096-13. Published Ahead of Print 17 May 2013.
	Updated information and services can be found at: http://ec.asm.org/content/12/7/1009
	These include:
SUPPLEMENTAL MATERIAL	Supplemental material
REFERENCES	This article cites 87 articles, 33 of which can be accessed free at: http://ec.asm.org/content/12/7/1009#ref-list-1
CONTENT ALERTS	Receive: RSS Feeds, eTOCs, free email alerts (when new articles cite this article), more»

Information about commercial reprint orders: http://journals.asm.org/site/misc/reprints.xhtml To subscribe to to another ASM Journal go to: http://journals.asm.org/site/subscriptions/







A SAS-6-Like Protein Suggests that the *Toxoplasma* Conoid Complex Evolved from Flagellar Components

Jessica Cruz de Leon,^a Nicole Scheumann,^b Wandy Beatty,^c Josh R. Beck,^d Johnson Q. Tran,^a Candace Yau,^a Peter J. Bradley,^d Keith Gull,^b Bill Wickstead,^{b,e} Naomi S. Morrissette^a

Department of Molecular Biology and Biochemistry, University of California, Irvine, Irvine, California, USA^a; Sir William Dunn School of Pathology, University of Oxford, Oxford, United Kingdom^b; Department of Molecular Microbiology, Washington University School of Medicine, St. Louis, Missouri, USA^c; Department of Microbiology, Immunology and Molecular Genetics, University of California, Los Angeles, Los Angeles, California, USA^d; Centre for Genetics and Genomics, University of Nottingham, Nottingham, United Kingdom^e

SAS-6 is required for centriole biogenesis in diverse eukaryotes. Here, we describe a novel family of SAS-6-like (SAS6L) proteins that share an N-terminal domain with SAS-6 but lack coiled-coil tails. SAS6L proteins are found in a subset of eukaryotes that contain SAS-6, including diverse protozoa and green algae. In the apicomplexan parasite *Toxoplasma gondii*, SAS-6 localizes to the centriole but SAS6L is found above the conoid, an enigmatic tubulin-containing structure found at the apex of a subset of alveolate organisms. Loss of SAS6L causes reduced fitness in *Toxoplasma*. The *Trypanosoma brucei* homolog of SAS6L localizes to the basal-plate region, the site in the axoneme where the central-pair microtubules are nucleated. When endogenous SAS6L is overexpressed in *Toxoplasma* tachyzoites or *Trypanosoma* trypomastigotes, it forms prominent filaments that extend through the cell cytoplasm, indicating that it retains a capacity to form higher-order structures despite lacking a coiled-coil domain. We conclude that although SAS6L proteins share a conserved domain with SAS-6, they are a functionally distinct family that predates the last common ancestor of eukaryotes. Moreover, the distinct localization of the SAS6L protein in *Trypanosoma* and *Toxoplasma* adds weight to the hypothesis that the conoid complex evolved from flagellar components.

entrioles and basal bodies are microtubule-based structures found in many eukaryotic lineages, including animals, lower plants, and diverse unicellular organisms (1-3). Basal bodies are distinguished from centrioles by association with a flagellar axoneme that is templated from an extension to the centriole known as a transition zone (1, 4). The widespread occurrence of centrioles and basal bodies shows that they have an ancient origin in eukaryotes (5-8), but lineages such as flowering plants and most fungi have lost the ability to build these structures. Centrioles and basal bodies typically contain nine triplet microtubules organized with radial symmetry, although they can rarely be built of nine doublet or singlet microtubules (e.g., those in Drosophila embryos and Caenorhabditis elegans testes, respectively) or have likely 6-fold symmetry (9, 10). In spite of their presence at spindle poles, centrioles are dispensable for bipolar spindle formation in several lineages (11-15). However, all species that build centrioles/basal bodies have flagella at some stage of their life cycle; this observation likely underlies the true evolutionary imperative of these structures (2, 3).

Many protozoan organisms, including apicomplexan and kinetoplastid parasites, use spatially separated and morphologically distinct microtubule-organizing centers (MTOCs) to organize individual microtubule populations (16). Kinetoplastid parasites of the genera *Trypanosoma* and *Leishmania* cause human infectious diseases, including African sleeping sickness, Chagas disease, and kala azar (17). The shape of these organisms is maintained by an array of ~100 densely packed corset microtubules that underlie the plasma membrane (18). The single flagellum is nucleated by a membrane-docked basal body that is distant from the nucleus (18–22). Flagella are not disassembled during division, and in replicating cells, basal bodies do not contribute to the organization of the poles of the intranuclear spindle (23). Apicomplexan parasites also cause a variety of medically significant diseases, including

malaria, toxoplasmosis, and cryptosporidiosis (17). Apicomplexans typically have a complex life cycle that involves both asexual and sexual replication, and they alter microtubule architecture between the sexual and asexual life cycle stages. Microgamete motility is required for fertilization of macrogametes and is powered by flagella that originate from apical basal bodies (24). Asexual stages lack flagella and use a characteristic actin-and-myosinbased gliding motility to invade host cells (25, 26). These asexual forms (e.g., merozoites, tachyzoites) have two microtubule populations; spindle microtubules coordinate chromosome segregation during mitosis, and subpellicular microtubules subtend the pellicle to impose an elongated cell shape and cell polarity (27-29). Each microtubule population is associated with a distinct MTOC; subpellicular microtubules radiate from the apical polar ring (APR), an MTOC unique to apicomplexan organisms (28, 30, 31), whereas spindle microtubules originate near a specialized region of the nuclear envelope termed the centrocone (32–35).

Toxoplasma gondii is a member of the Coccidia—a subclass of apicomplexan parasites that build two tubulin-based structures in addition to the spindle and subpellicular microtubules, i.e., the

Received 15 April 2013 Accepted 8 May 2013 Published ahead of print 17 May 2013 Address correspondence to Naomi Morrissette, nmorriss@uci.edu, or Bill Wickstead, bill.wickstead@nottingham.ac.uk. J.C.d.L. and N.S. are joint first authors. Supplemental material for this article may be found at http://dx doi.org/

Supplemental material for this article may be found at http://dx.doi.org/10.1128 /EC.00096-13.

Copyright © 2013 de Leon et al. This is an open-access article distributed under the terms of the Creative Commons Attribution 3.0 Unported license. doi:10.1128/EC.00096-13 conoid and centrioles. The conoid is an apical organelle constructed of comma-shaped tubulin sheets that spiral to form a cone-shaped structure (36-38). Two preconoidal rings surmount the conoid, and when extended, the conoid and preconoidal ring complex reside above the APR (28, 39). The conoid and preconoidal rings can also retract through the APR to be surrounded by the subpellicular microtubules. The conoid is permanently retracted in intracellular parasites, but extracellular tachyzoites extend and retract this structure, a probing behavior that is believed to facilitate host cell invasion (36-38). Two short, closely apposed microtubules are located at the center of the conoid. In contrast to the apical conoid, Toxoplasma centrioles are found in the cytoplasm proximal to the nucleus and duplicated centrioles are found at the spindle poles in the region of the centrocone structures. Coccidian centrioles are composed of nine singlet microtubules and a central tubule and are organized in a parallel rather than an orthogonal configuration (27, 33, 40).

SAS-6 is required for centriole biogenesis in eukaryotes ranging from protozoa to vertebrates (41-44). Chlamydomonas reinhardtii mutants deficient in SAS-6 (bld12) produce basal bodies with variable numbers of triplet blades (45). Studies of nematodes and Drosophila indicate that SAS-6 is needed to form the "cartwheel hub" in nascent centrioles (45-49). The structural properties of SAS-6 elegantly dovetail with its ability to template a centriole (50, 51). SAS-6 assembles into homodimers that contain a globular head domain at the N terminus and an extended coiledcoil rod. Remarkably, SAS-6 homodimers can self-assemble into a structure with a central hub and nine rods, akin to the cartwheel observed in structural studies of centrioles. In this paper, we describe for the first time the properties of a family of SAS-6-like proteins that are related to SAS-6 but structurally distinct. We investigate the function of SAS-6-like proteins in two distantly related protozoa and discuss their evolutionary relationship to conventional SAS-6.

MATERIALS AND METHODS

Culture of parasites. *Toxoplasma* lines, including the previously described RNG1-yellow fluorescent protein (YFP) line (31, 52), were grown in confluent monolayers of human foreskin fibroblast (HFF) cells as previously described (53). *Trypanosoma brucei* procyclic forms were cultured at 28°C in SDM-79 medium supplemented with 10% fetal calf serum (54).

Generation of fusion protein lines. To generate in-frame C-terminal chimeras with proteins of interest, regions of TGGT1_040920 (TgSAS-6), TGGT1_059860 (TgSAS6L), Tb927.10.7920 (TbSAS6L), and Tb11.02.5550 (TbWDR16) were amplified and cloned into LIC (52), pLew-MH-TAP-eYFP (55), and pEnT6P-Y (56) vectors by standard methods. TgSAS-6 and TgSAS6L constructs in the pYFP.LIC.D-HFR vector were transfected into ku80-null Toxoplasma parasites as previously described, and stable lines were isolated by selection in 1 μ M pyrimethamine and single cell cloned (52). The coding sequence of TgSAS6L was validated by amplification of cDNA encoding the complete open reading frame with primers described in Table 1. A pre-existing GRASP55-YFP construct (57) was modified to replace the GRASP55 coding sequence with SAS6L to drive the expression of TgSAS6L-YFP from the α 1-tubulin promoter, and stable lines were isolated by selection for chloramphenicol resistance. Linearized plasmid DNA was used to transfect exponentially growing cultures of procyclic-form T. brucei by electroporation (three 100-µs pulses of 4.25 kV/cm). Transfected cells were selected by the addition of 1 µg/ml puromycin (pEnT6P-TbSAS6L1-Y) or 10 µg/ml blasticidin (pEnT6B-WDR16-mCherry). Inducible overexpression of TbSAS6L was achieved by cloning TbSAS6L into the pLew-MH-TAP-eYFP vector with primers

study	
Primer	Sequence
TbSAS6L CDS pEnT6P 5'	GTTGCGGCCGCAAAGAACCCCCAACGCTTTTT
TbSAS6L CDS pEnT6P 3'	GTCTACTAGTTCGCGTTTTTCCAAGTGTG
TbSAS6L UTR pEnT6P 5'	GTTAAGCTTGAGTGCAAATTCTTCAGTGCG
TbSAS6L UTR pEnT6P 3'	GTTGCGGCCGCCGACATTAGCTCCGGTTTCT
WDR16 CDS pEnT6P 5'	CGTTCTAGAGCAGCGAAGGAGTACGAAA
WDR16 CDS pEnT6P 3'	TCGAGCACGGCCAGGTAGCTGAT
WDR16 UTR pEnT6P 5'	GCTCGAGCTTTCGCAAACCCATTCG
WDR16 UTR pEnT6P 3'	GTAGGATCCAACGGAAGGGTGGCAGTT
5′ TbSAS6L HindIII pLew	ATCAAGCTTATGGATCGCATAGAAATATACTACCAG
3′ TbSAS6L1 XhoI pLew	GATCTCGAGTCGCGTTTTTCCAAGTGTGACAGTAGC
SAS6L LIC vector 5'	TACTTCCAATCCAATTTAATGCACAGACGGAAATG CTCTCC
SAS6L LIC vector 3'	TCCTCCACTTCCAATTTTAGCGAGGAACCGAGTG GATGC
SAS6L MBP plasmid 5'	ATGGCGACAAACTTCGGCTTTGG
SAS6L MBP plasmid 3'	TATAAAGCTTTCAGAGGAACCGAGTGGATGCGCC
ptub-SAS6L-YFP 5'	CAAAGATCTATGGCGACAAACTTCGGCTTTGG
ptub-SAS6L-YFP 3'	TATACCTAGGGAGGAACCGAGTGGATGCGCC
SAS6L KO 5' flank 5'	AAGGTACCCCCTCTCTCACAGTCGAAGACC
SAS6L KO 5' flank 3'	TTGGGCCCGTTATTCTGTTCGAACCCGGGG
SAS6L KO 3' flank 5' SAS6L KO 3' flank 3'	AAACTAGTCGGTGGGAGGTCTCAAGCG GCGCGGCCGCCACGTAATCACACAATCCGAGCGT ATAG

Downloaded from http://ec.asm.org/ on April 29, 2014 by Univ of Nottingham

described in Table 1. Transfectants were selected in 5 μ g/ml phleomycin, and overexpression of TbSAS6L1-YFP was induced by the addition of 1 μ g/ml doxycycline to the culture medium.

Toxoplasma SAS6L gene knockout. We constructed a knockout vector with the pmini.GFP.HPT plasmid (58). A 4.9-kb region upstream of the TGGT1_059860 coding sequence and a 5-kb downstream region were amplified with primers listed in Table 1. These inserts were cloned upstream and downstream of the hypoxanthine xanthine guanine phosphoribosyl transferase (HPT) gene. We electroporated $ku80\Delta$ hpt mutant RH tachyzoites with the construct and selected for transformants in 50 µg/ml mycophenolic acid and 50 µg/ml xanthine. Single-knockout clones were isolated from two independent transfections by single-cell cloning. We also isolated green fluorescent protein (GFP)-expressing transformants with a nonhomologous integration of the knockout vector. These were used in competition assays with the knockout line (see below).

Escherichia coli expression and purification of TgSAS6L. The cDNA sequence of TgSAS6L was inserted into the c2x pMAL vector (NEB) in order to express a maltose-binding protein (MBP)-TgSAS6L fusion in *E. coli*. After induction for 2 h with 300 μ M isopropyl- β -D-thiogalactopyranoside (IPTG), the protein was purified on an amylose column and cleaved away from MBP with Factor Xa. Dialyzed TgSAS6L was used as an immunogen to generate mouse polyclonal antiserum.

Toxoplasma conoid extrusion. Conoid extrusion was induced in freshly lysed extracellular parasites by treatment with HEPES-buffered saline supplemented with 5 μ M ionomycin (Sigma) and 5 mM CaCl₂ (Sigma) (36–38), and samples were fixed as previously described (31).

Immunofluorescence staining and fluorescence microscopy. YFPor mCherry-tagged intracellular *Toxoplasma* parasites were fixed, permeabilized, and stained as previously described (27). Extracellular parasites were filtered through a 3- μ m polycarbonate filter (GE Water & Process Technologies), centrifuged at 1,000 × g for 20 min at 4°C, and suspended in a small volume of phosphate-buffered saline (PBS). Detergent-extracted parasites were generated and stained as previously described (31). The antibodies used for immunofluorescence assays included a mouse polyclonal serum against TgSAS6L, a mouse monoclonal antibody against GFP (Roche), and a rabbit *Toxoplasma*-specific tubulin serum (27) detected with Alexa 594-, Alexa 488-, and Cascade blue-conjugated secondary antibodies (Invitrogen). DNA was visualized by 4',6-diamidino-2-phenylindole (DAPI) staining. *Toxoplasma* samples were imaged on a Zeiss Axiovert 200M microscope with the AxioVision system. Trypanosomes were settled onto glass slides and either fixed with 1% formaldehyde for whole-cell preparations or extracted with 1% Nonidet P-40 in PEME [100 mM 100 mM piperazine-*N*,*N'*-bis(2-ethanesulfonic acid) (PIPES, pH 6.9), 2 mM EGTA, 1 mM MgSO₄, 0.1 mM EDTA] for cytoskeleton preparations. Cells were labeled with BBA4 or YL1/2 antibodies, which were visualized with anti-mouse IgM–Cy5 and anti-rat IgG–tetramethyl rhodamine isothiocyanate (Jackson ImmunoResearch). Cells were mounted in 90% glycerol–50 mM sodium phosphate (pH 8) supplemented with 1% 1,4-diazabicyclo[2.2.2]octane and 0.4 μ g ml⁻¹ DAPI.

Immunoblot assays. Protein lysates containing $\sim 5 \times 10^6$ *Toxoplasma* tachyzoites per lane were resolved by 15% SDS-acrylamide gel electrophoresis and transferred to nitrocellulose. After blocking for 1 h in 5% milk in PBS-Tween, the blot was probed with primary antibodies (1:300 anti-TgSAS6L and 1:750 anti-ISP3 in 5% milk) for 1 h, washed in PBS-Tween, probed with a secondary antibody (1:2,000 horseradish peroxidase-conjugated goat anti-mouse antibody; Invitrogen) for 1 h, and washed in PBS-Tween prior to detection by chemiluminescence assay.

Electron microscopy (EM). Freshly lysed *Toxoplasma* tachyzoites from stable TgSAS6L-YFP lines were isolated by filtration. Parasites were fixed and cryosectioned as previously described (31), with an anti-GFP antibody (Abcam). Trypanosomes were settled onto Formvar-coated nickel finder grids and extracted with 1% NP-40–PEME. For Ca²⁺ preparations, cells were additionally incubated with 65 mM Ca²⁺–PIPES, pH 6.9. Preparations were fixed with 2.5% glutaraldehyde–PEME for 10 min and then neutralized with 1% glycine–PEME. Grids were mounted in H₂O, and native fluorescence was analyzed with a Leica DM5500 microscope. Grids were negatively staining with 1% aurothioglucose (USP) and viewed on an FEI Tecnai-F12 electron microscope.

Toxoplasma competition assays. T_{25} flasks with confluent HFF cells were inoculated with a 1:1 ratio of *sas6l*-null parasites and GFP-expressing clones from the same transfection that harbor a nonhomologous integration of the knockout vector (10^7 parasites of each line) as described in a previously established competition assay (59). The relative numbers of GFP-positive (control) and GFP-negative (*sas6l*-null) parasites were determined by flow cytometry (Apogee Flow Systems).

Alignments and phylogenetic analysis. SAS6 and SAS-6-like proteins form a natural set on the basis of a BLASTp clustering approach (60). This clustering results from the presence of a conserved domain of \sim 200 aa covered by a Pfam-B domain (Pfam_B_2528) of unknown function. To define the family of SAS6 and SAS-6-like proteins, the hidden Markov model defining Pfam_B_2528 was used to search the complete predicted proteomes of 44 diverse eukaryotes (HMMER3.0; hmmer.janelia.org). All 69 proteins containing the domain (e-value, <0.001) were extracted, trimmed to the conserved domain, and aligned by MAFFT v6.811b by adopting the E-INS-i strategy (61). Bayesian phylogenies were inferred from four runs of the metropolis-coupled Markov chain Monte Carlo method implemented in the program MrBayes3.1.2 (62) (WAG substitution matrix; $GTR + \Gamma + I$ with Γ distribution approximated to four discrete categories and shape parameters estimated from the data). Maximumlikelihood support for the inferred topology was generated from 100 bootstrap replicates of the data with PhyML3.0 (63) (LG matrix; $GTR+\Gamma+I$, as described above).

RESULTS

SAS-6 and SAS6L localize to distinct structures in *Toxoplasma* **tachyzoites.** SAS-6 is an essential and universal component of centrioles that creates the "cartwheel hub" found at the proximal end of the centriole barrel (41, 42, 45). We came upon the hypothetical protein TGGT1_059860 during a genome database search

(www.ToxoDB.org) for the Toxoplasma homolog of SAS-6. Since TGGT1_059860 has similarity to SAS-6, we named it SAS-6-like (SAS6L). When the Toxoplasma SAS-6 homolog (TGGT1_ 040920) was tagged by fusion of the C terminus to YFP, the label localized exclusively to the centriole, coincident with the centriolar marker centrin (Fig. 1A), as is expected from the localization of SAS-6 in other systems. In contrast, TgSAS6L localizes exclusively to a region at the apex of the parasites (Fig. 1B). Since TgSAS6L labeled a small spot in the apex of Toxoplasma tachyzoites, we induced conoid extrusion in extracellular parasites and used a marker for the APR (RNG1) (31) to localize TgSAS6L more accurately. TgSAS6L is found at the apical tip of extended conoids, above the RNG1 signal (Fig. 1C). In parasites with retracted conoids, labeling is coincident with the RNG1 marker (Fig. 1D). TgSAS6L remains associated with the cytoskeleton following treatment with deoxycholate (DOC), which extracts much of the inner membrane complex-associated cytoskeleton and frees the subpellicular microtubules. In DOC-extracted samples, TgSAS6L labeling is at the apical tip of the conoid (Fig. 1E). Immunogold labeling of TgSAS6L-YFP parasites with an anti-GFP antibody indicates that the protein is located at the apex of the conoid, in a region consistent with the preconoidal rings (Fig. 1F).

SAS6L-null *Toxoplasma* are less fit. We deleted the SAS6L gene from *Toxoplasma* tachyzoites by targeted integration of the HPT gene (Fig. 2A) and confirmed the knockout by the absence of SAS6L protein (Fig. 2B and C). Since there are no specific markers for the conoid fibers, we induced conoid extrusion and stained with a tubulin antibody to ascertain that the conoid is still present in null parasites (not shown). TgICMAP1 is a microtubule-associated protein that specifically localizes to the two intraconoid microtubules that run down the center of the conoid (64). We used an established expression construct (pmin-eGFP-TgICMAP1 [64]) to verify that the eGFP-TgICMAP1 fusion protein correctly localizes to the region of the intraconoid microtubules in the absence of SAS6L (not shown).

During creation of the sas6l Δ mutant lines, we also cloned lines that had nonhomologous integration of the knockout vector for use as a matched control in growth competition assays. We coinfected host cells with an equivalent number of each clonal line and followed the relative rates of replication and growth over serial passage by flow cytometry (Fig. 2D). Two independent knockout lines were outcompeted by a matched control at a rate of 0.07 day⁻¹, demonstrating a reduction in fitness. The reduced sas6l Δ mutant parasite growth observed may reflect a specific defect in a number of cellular processes required for invasion and intracellular growth (for example, motility, attachment, invasion, doubling time, replication fidelity, egress, and extracellular viability). In order to assess specific defects in these processes, we measured gliding motility, performed plaque formation assays (for invasion and extracellular survival), assessed Ca²⁺ ionophore-mediated egress, and quantified replication rates and the frequency of replication defects. Within the resolution of these assays, no significant differences between two independent sas6l-null lines and the matched control were detected (data not shown). However, the expected size of a specific defect in assays such as the plaque assay $(\sim 10\%)$, given the rate of outcompetition) was below the assay's resolving power because of the relative variation between biological replicates. Hence, it is likely that the direct fitness assay detected a defect that was below the sensitivity of the other assays in our experiments.

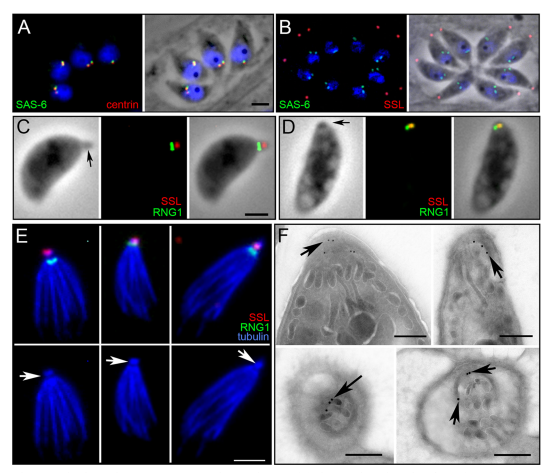


FIG 1 TgSAS6 and TgSAS6L localize to distinct regions in *Toxoplasma* tachyzoites. (A) The *Toxoplasma* homolog of SAS6 (green) localizes to centrioles along with the centriole marker centrin (red). The overlap between these proteins is not complete, which likely reflects distinct localization within centrioles; while centrin is typically located throughout the centriole, SAS-6 is a marker of the proximal end of centrioles. (B) TgSAS6L (red, anti-SAS6L antiserum) is found at the apex of *Toxoplasma* tachyzoites and does not colocalize with SAS-6 in the centrioles (green, YFP tag). (C) Ionomycin triggers conoid extrusion in most extracellular parasites. In these cells, SAS6L-mCherry labeling (red) is above the APR (RNG1-YFP, green). (D) When the conoid is retracted, TgSAS6L and RNG1-YFP partially colocalize (the preconoidal region is smaller than the APR). (E) Extraction of extracellular tachyzoites with DOC frees the subpellicular microtubules and conoid (blue) and APR (RNG1-YFP, green) from other cellular material. TgSAS6L is located at the apical end of the conoid, and RNG1-YFP is beneath the extended conoid at the APR. The tubulin channel alone permits identification of the conoid (arrows) above the subpellicular microtubules. (F) Immunogold labeling indicates that SAS6L-YFP localizes to the extreme apical region of *Toxoplasma* tachyzoites, above the conoid in both longitudinal (arrows, top) and transverse (arrows, bottom) sections of the apical region. Scale bars, 2 µm (A to E) or 250 nm (F).

SAS6L is an ancient protein found in diverse simple eukaryotes. Similarity searches suggested the presence of SAS6L homologs in diverse eukaryotes. All SAS6L proteins contain an ~200-amino-acid (aa) conserved domain that is shared with SAS-6 homologs (see Fig. S1 in the supplemental material) and includes the ~50-aa PISA motif previously described in SAS-6 (42). This domain is described by the automatically generated Pfam-B domain, Pfam_B_2528. We used Pfam_B_2528 to identify proteins of the SAS-6/SAS6L family encoded by the genomes of 29 of 44 diverse eukaryotes (Fig. 3A). These proteins share an average of 24% identity (46% similarity) across the conserved domain. A similar set was defined by an alternative approach based on clustering by BLASTp score (data not shown).

Phylogenetic analysis based on the conserved domain robustly supported the existence of two subfamilies comprising SAS-6 and SAS6L (Fig. 3B). Genes encoding SAS-6 are found in the genomes of eukaryotes that are known to build a centriole or basal body and also in the microsporidian *Encephalitozoon cuniculi* and *Ostreo*- *coccus tauri* as previously described (42–45, 60). With the exception of *Giardia lamblia*, which lacks an identifiable SAS-6 ortholog (60), SAS6L is found only in lineages that also contain SAS-6 (Fig. 3A). In addition to *Toxoplasma*, SAS6L is found in multiple protistan lineages (including other apicomplexans, such as *Plasmo-dium* and *Cryptosporidium* species). However, SAS6L appears to be absent from metazoan and choanoflagellate genomes (Holozoa). The presence of SAS6L in a basal flagellate fungus (*Batrachochytrium dendrobatidis*) indicates that the absence of SAS6L from animals is likely the result of a specific loss near the root of Holozoa.

Although only the conserved domain was used to define the phylogenetic families, the SAS-6 and SAS6L proteins have clearly distinct architectures (Fig. 3B). SAS-6 sequences are generally \sim 600 aa in length and comprise the conserved domain near the N terminus, followed by \sim 300 aa of sequence rich in predicted coiled coils. In contrast, SAS6L sequences contain little sequence outside the conserved domain and appear to lack the C-terminal

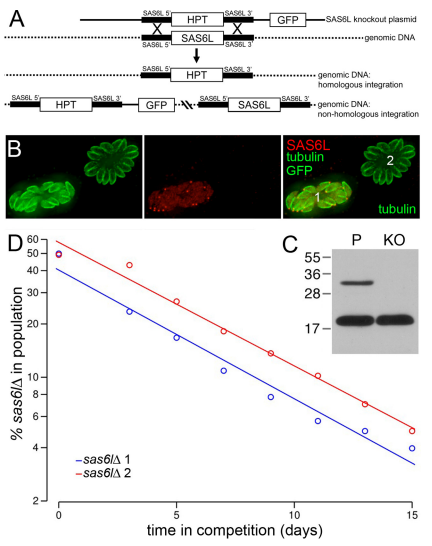


FIG 2 Loss of TgSAS6L reduces *Toxoplasma* fitness. (A) Knockouts were generated by homologous recombination of the HPT gene into the TgSAS6L locus. In the case that the vector integrates nonhomologously, both the GFP and HPT genes are integrated, generating fluorescent parasites. (B) *sas6l* Δ mutants were confirmed by loss of immunofluorescence labeling for TgSAS6L (red). All parasites (1 and 2) are labeled for tubulin (green). Nonhomologous integration of the knockout vector (parasite 1) leads to parasites that retain TgSAS6L and also have the GFP signal. (C) Immunoblotting with both anti-TgSAS6L and anti-ISP3 antisera also demonstrates the complete absence of TgSAS6L from *sas6l* Δ mutant knockout (KO) cells relative to the parental line (P). The values to the left are molecular sizes in kilodaltons. (D) Two independently derived *sas6l*-null parasite lines have reduced growth relative to that of GFP-expressing clones with a nonhomologous integration of the knockout vector.

tail entirely. The conserved domain essentially matches the globular head domain of SAS-6, the structure of which has been solved (50, 51). Homology modeling suggests that SAS6L proteins can adopt structures very similar to those seen in the head of SAS-6 (see Fig. S2 in the supplemental material). The apicomplexan SAS-6 proteins are unusual in possessing long N-terminal extensions before the conserved domain (Fig. 3B).

TbSAS6L localizes to the axonemal basal plate in trypanosomes. Antiserum against TgSAS6L labels a spot at the apex of the related parasite *Neospora caninum* (not shown). Unfortunately, our antiserum does not label SAS6L proteins in more distantly related organisms, such as *Tetrahymena thermophila* (data not shown). Since the majority of SAS6L-producing eukaryotes do not have a conoid, we tested the localization of a SAS6L ortholog in a distantly related organism. In *Trypanosoma brucei* trypomastigotes constitutively expressing TbSAS6L-YFP from the endogenous locus, TbSAS6L-YFP localizes near the base of the flagellum (Fig. 4A to C). Label appears as one or two prominent dots, depending on the cell cycle stage, and there is also weaker labeling in the proximal region of the axoneme, in the region of the flagellar pocket collar. TbSAS6L-YFP labeling is distal to markers of the proximal (BBA4), middle (tyrosinated α -tubulin, YL1/2), and distal (WDR16) parts of the basal body. Quantification of the location of TbSAS6L-YFP relative to these markers indicates that this protein is located in a fixed position within the axoneme (Fig. 4D), in a region consistent with the location of the basal plate (the area where the central-pair microtubules of a flagellar axoneme are nucleated). Ablation of TbSAS6L by RNA interference does not alter the overall morphology of the flagellum, as viewed by EM, suggesting that SAS6L has a nonessential role in flagellum

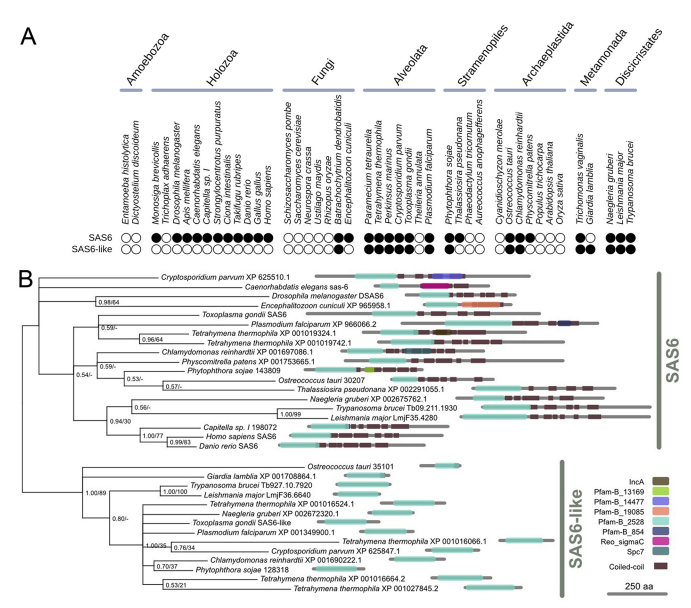


FIG 3 SAS-6 and SAS6L proteins are found in a subset of organisms that form cilia or flagella. (A) Presence (filled circles) or absence (empty circles) of identifiable homologues of SAS-6 and SAS6L in the predicted proteomes of 44 diverse eukaryotes. SAS6L protein is present in most organisms that also possess SAS-6 but absent from the Holozoa. (B) The SAS-6 and SAS6L protein families share a conserved domain of \sim 200 as but can be robustly separated by phylogenetic analysis and have distinct architectures. A Bayesian phylogeny of the conserved domain is shown with topology support from Bayesian posterior probabilities and maximum likelihood bootstrap replicates (PP/BS). Architectures show predictions of Pfam-A and -B domains (89) and coiled-coil regions (90).

formation (not shown). This is not entirely surprising, given the plasticity of flagellum formation, but it is also, of course, extremely difficult to detect subtle alterations in this complex structure.

Overexpression of SAS6L-YFP causes ectopic filament formation. To investigate the *in vivo* properties of SAS6L, we expressed tagged SAS6L in both *Toxoplasma* and *Trypanosoma* over endogenous protein. In *Toxoplasma*, when TgSAS6L-YFP expression was driven from the α 1-tubulin promoter, excess protein assembled into filaments in a subset of the sibling *Toxoplasma* parasites within a single parasitophorous vacuole (Fig. 5A). Filaments sometimes deformed the parasite's shape and may interfere with cell division, as in the parasites shown in Fig. 5B, where a large filament extends between daughters that have begun a new round of replication without completing scission. TgSAS6L-YFP filaments form when intracellular parasites are treated with oryzalin, which inhibits microtubule polymerization (not shown). Ectopic TgSAS6L-YFP filaments are retained after extraction with 1% Triton X-100 but are sensitive to 10 mg/ml DOC, which removed TgSAS6L-YFP filaments without disrupting the apical TgSAS6L signal (Fig. 5C). Immunogold staining of EM cryosections of whole cells showed that TgSAS6L-YFP was associated with continuous electron-dense structures in the cytoplasm (Fig. 5D).

Overexpression of TbSAS6L-YFP in trypanosomes also resulted in the accumulation of ectopic filaments (Fig. 5E). The formation of these structures has no effect on the morphology or

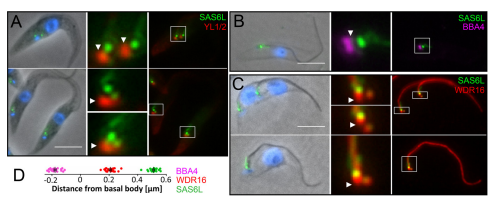


FIG 4 TbSAS6L-YFP localizes to a prominent focus near the base of the flagellum in *T. brucei*. (A) TbSAS6L-YFP (green) is distal to the basal body, which is marked by tyrosinated α -tubulin (YL1/2, red, arrowheads). A lower-intensity SAS6L-YFP signal is also associated with the developing probasal body. SAS6L-YFP is also distal to the basal-body markers BBA4 (B) and WDR16 (C) (arrowheads). (D) Distance measurements in relation to YL1/2 indicate that SAS6L-YFP localizes to a region near the axoneme basal plate (20 measurements for each marker). Scale bars, 5 μ m.

duplication time of the cells (data not shown). The filaments are insensitive to extraction with nonionic detergent (Nonidet P-40) and are not labeled with anti-tubulin antibodies (not shown), indicating that they are independent of microtubules. TbSAS6L-YFP filaments are insensitive to treatment with Ca^{2+} , which depolymerizes *T. brucei* subpellicular microtubules (Fig. 5G). Correlative microscopy of detergent-extracted trypomastigotes was used to unambiguously determine the ultrastructure of TbSAS6L-YFP filaments by first identifying filaments by fluorescence microscopy and subsequently imaging them by EM (Fig. 5F and G). TbSAS6L filaments are \sim 20 nm wide but have a structure distinct from that of microtubules (Fig. 5G).

DISCUSSION

We describe a novel conserved protein that is related to the centriole protein SAS-6, which we have hence named the SAS-6-like (SAS6L) protein. The existence of SAS6L produced by diverse organisms suggests that this protein appeared early in eukaryote

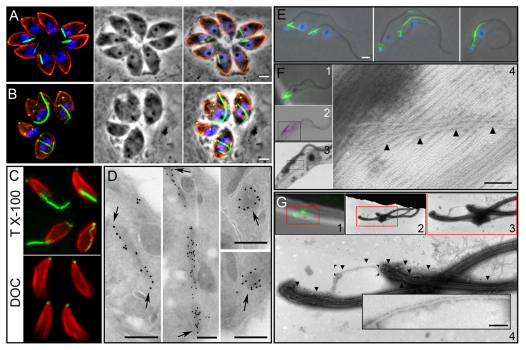


FIG 5 Overexpression of SAS6L-YFP leads to filament formation. (A) Intracellular *Toxoplasma* tachyzoites show prominent TgSAS6L-YFP filaments as follows: *Toxoplasma* tubulin, red; SAS6L-YFP, green; DNA, blue. (B) TgSAS6L-YFP filaments appear to interfere with the completion of division by some tachyzoites. (C) SAS6L-YFP filaments are insensitive to 1% Triton X-100 (TX-100), but 10 mg/ml DOC causes filament depolymerization although apical SAS6L is retained. (D) EM of immunogold-labeled cryosections of *Toxoplasma* overexpressing TgSAS6L-YFP show electron-dense structures of variable width within the cytoplasm. (E) Overexpression of the TbSAS6L protein in trypanosomes also leads to ectopic filaments. (F) Correlative light microscopy and EM of TbSAS6L-YFP filaments in trypanosomes (arrowheads). The same cell cytoskeleton is shown by phase-contrast microscopy (F1) and negatively stained transmission EM (F2 to -4). (G) Correlative microscopy of Ca²⁺-treated cytoskeleton preparations shows that TbSAS6L-YFP filaments are resistant to Ca²⁺ (arrows). Scale bars: A and B, 2 μm; D, 250 nm; E, 2 μm; F4, 200 nm; G4 inset, 200 nm.

evolution and most likely before the last common ancestor of all extant eukaryotes (65). SAS6L is found in simple eukaryotes that contain SAS-6 and centrioles/basal bodies, but the protein has been lost from the holozoan lineage. SAS6L proteins consist of a conserved domain that also comprises the N-terminal head domain of SAS-6. Recent data have shown that SAS-6 forms homodimers through interactions between their coiled-coil tail domains and that these homodimers interact via other parts of the head domain to form higher-order structures (in particular, rings) (50, 51). Since SAS6L proteins lack a SAS-6 tail, they cannot dimerize by this means. However, the opposite face of the protein is available for interaction and is predicted to have a structure very similar to that of SAS-6. The observation that overexpression of SAS6L proteins induces SAS6L filament formation indicates that SAS6L proteins do, indeed, have the ability to form higher-order structures. However, the distinct localization of SAS-6 and SAS6L proteins in Toxoplasma and Trypanosoma suggests that these proteins do not form heterodimers in vivo. The decreased stability of the ectopic SAS6L filaments after DOC extraction suggests that additional interactions stabilize the native preconoidal ring structure and do not occur in the cytoplasmic filaments.

Consistent with its location in other diverse eukaryotes, SAS-6 localizes to the basal body in Trypanosoma and the centrille Toxoplasma tachyzoites. In contrast, SAS6L localizes to the region of the basal plate in Trypanosoma brucei-the electron-dense region at the distal end of the transition zone from which the central-pair microtubules are nucleated. Unlike T. brucei (where flagella are present in all life cycle forms), only the microgamete forms of Toxoplasma or other apicomplexan organisms build flagella (24, 32, 66-69). Comparative genomics indicate that the Toxoplasma and Plasmodium genomes lack a surprising number of conserved basal-body components (60), and while studies of Plasmodium microgamete development show de novo formation of basal bodies (66), it is unclear whether Toxoplasma microgamete basal bodies form *de novo* or if the atypical centriole found in the asexual tachyzoite stage serves as a template for basal-body formation. We anticipate that SAS6L will also localize to the basal plate region in Toxoplasma microgametes, but because of the difficulty in obtaining this stage from cat intestinal epithelium, we cannot verify this prediction. In agreement with this hypothesis, there is evidence of upregulation of the Plasmodium falciparum ortholog of SAS6L (PF3D7_1316400) in gametocyte development (70).

The Toxoplasma tachyzoite stage causes most of the pathology associated with parasite infection. Despite lacking flagella, tachyzoites are motile and employ an unusual substrate-dependent gliding motility that requires actin and myosin (71–73). The apical region of tachyzoites is vital to host cell invasion and contains specialized secretory organelles (micronemes and rhoptries) and a complex and unusual organization of the microtubule cytoskeleton (28, 39, 74, 75). The elongated cell shape of tachyzoites is maintained by a corset of 22 subpellicular microtubules that radiate out of the APR. An additional tubulin-based structure, the conoid, can be extended through the APR or retracted into it. In intracellular parasites, the conoid is retracted and immotile, while in extracellular parasites, the conoid can extend and retract as tachyzoites glide and invade host cells (36). Extension of the conoid is associated with the elongation and narrowing of the apical end of tachyzoites, and it is believed that the conoid has a mechanical role in parasite invasion of host cells.

Apicomplexans share ancestry with ciliates and dinoflagellates;

a link between these organelles. blin) in Toxoplasma. SFA proteins were first described in green

such as Colpodella vorax and Rastrimonas subtilis (previously known as Cryptophagus subtilis) have "pseudoconoid" or "incomplete conoid" structures consisting of a set of 5 to 14 interlinked microtubules located at the cell apex but lack the adjacent APR structure observed in coccidians (78-81). Colpodella and Rastrimonas also contain apical secretory organelles reminiscent of apicomplexan micronemes and rhoptries that are used along with a pseudoconoid to partially or completely invade other unicellular protists. Colpodella attaches to and aspirates the cytoplasmic contents of unicellular flagellates (76, 78, 80, 81), while Rastrimonas is an intracellular parasite of unicellular algae (79, 80). The precise phylogenetic position of Colpodella and Rastrimonas is unclear, as these alveolates have been variously described as dinoflagellates, as early-branching apicomplexans, as Perkinsozoa, or within sister lineages of dinoflagellates or apicomplexans (76, 78, 80, 81). What is clear is that these organisms are poised between predator and parasite states and provide tantalizing clues to the evolution of the intracellular apicomplexan lifestyle. Both Colpodella and Rastrimonas simultaneously build a pseudoconoid structure and adjacent flagella (Fig. 6A and B). In contrast, the flagellar axoneme and "true" conoid are mutually exclusive structures in coccidian parasites; microgametes have apically located basal bodies that template flagellar axonemes, while asexual proliferative-stage coccidians have juxtanuclear centrioles and apical conoid structures (Fig. 6A and B). This study describes SAS6L, a novel conserved protein that is located in the region of the basal plate of the axoneme in *T. brucei* and above the conoid in *Toxoplasma* tachyzoites, a life cycle stage that lacks flagella (Fig. 6C). Rastrimonas absorbs its flagella during intracellular growth, and this feature may have become more pronounced during the evolution of apicomplexan parasites, ultimately leading to the loss of flagella in all stages other than microgametes. Our results emphasize this relationship between the conoid and flagellum, providing molecular evidence of The observations presented here are reinforced by recent work on the basal-body-associated protein SFA (striated fiber assem-

collectively, these organisms are classified as alveolates (76, 77). It

is likely that the last common ancestor of apicomplexans and dinoflagellates had an open-sided "incomplete" conoid that was

modified into a closed conoid in apicomplexans and lost from

dinoflagellates (77, 78). Within the Apicomplexa, the conoid was

again lost from noncoccidian lineages. Although "true" conoid

structures are found only in coccidian apicomplexans, alveolates

algae, where flagellar basal bodies are embedded in a "cage" of flagellar rootlet fibers. Rootlets consist of distinct populations of centrin-containing contractile fibers (82), sinister fibers (83), and SFA-containing noncontractile striated fibers (84, 85). Genes for SFA homologs are found in apicomplexan genomes (86, 87). Remarkably, two SFA proteins (TgSFA2 and TgSFA3) form a filament that links the upper edge of the conoid to the juxtanuclear centrioles during tachyzoite replication (88). The SFAs appear shortly after centriole duplication (87), colocalizing in a fiber-like structure that emerges from the centrioles and extends apically to tether the conoid and APR to the centrioles (88). Loss of either SFA protein inhibits parasite replication by blocking the formation of daughter APRs and the subsequent construction of daughter buds. Nuclear division is not inhibited, and multiple nuclei accumulate in these aberrant parasites. These results suggest that although the asexual stages of apicomplexans have dispensed with

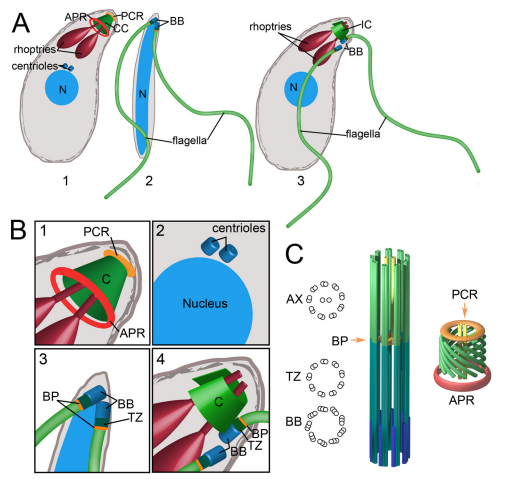


FIG 6 The Toxoplasma conoid incorporates flagellar components, and its appearance and that of flagella are mutually exclusive during the parasite life cycle. (A) Toxoplasma tachyzoites (part 1) have juxtanuclear centrioles and a specialized apical region used to invade host cells. Rhoptries (red) secrete components used during invasion, while the conoid (green) can extend from the APR to cause elongation of this region. Toxoplasma tachyzoites have a closed conoid (CC) that is topped by preconoidal rings (PCR), the location of SAS6L (orange). Toxoplasma microgametes (part 2) lack a conoid but have apically located basal bodies (BB) that template two flagella that extend toward the posterior. Alveolate organisms such as Colpodella and Rastrimonas (part 3) contain rhoptry-like secretory organelles (red) and an incomplete conoid (IC) structure consisting of a set of interlinked microtubules at the cell apex. They build flagella from adjacent basal bodies (BB) but lack the conoid-associated APR structure observed in coccidians. (B) The Toxoplasma tachyzoite conoid consists of 10 to 14 tubulin-containing filaments that spiral to form a closed cone that is narrower at the apex than at the base (box 1). The conoid may be extended from or retracted into the APR, which also serves to nucleate 22 subpellicular microtubules (not shown). The SAS6L protein is located at the extreme tip of the conoid (orange) in the region of the PCR. The juxtanuclear centrioles (box 2) are located at a distance from the conoid in tachyzoites. Toxoplasma microgametes (box 3) have apical centrioles that build a transition zone (TZ) and template flagellar axonemes. We predict that SAS6L localizes to the basal plate (BP) in this stage (orange). Alveolates such as Colpodella (box 4) have apical secretory organelles (red) and an incomplete conoid (IC) structure (green) that does not fully encircle the rhoptry necks and may be as simple as a set of four to six interlinked microtubules. Adjacent apical centrioles build a transition zone (TZ) to template flagellar axonemes with a basal-plate (BP) structure. These organisms lack the conoid-associated APR structure observed in Toxoplasma. (C, left) The basal body (BB) templates the flagellar axoneme (AX). The transition from the triplet microtubule organization of the basal body to the doublet microtubule organization of the axoneme occurs at the basal plate (BP), which is also where the central-pair microtubules of the axoneme begin. SAS6L in trypanosomes localizes to the region of the flagellar plate (orange). (C, right) The Toxoplasma conoid is formed from 10 to 14 C-shaped and spiraling tubulin filaments. Two 13-protofilament microtubules extend through the center of the conoid, reminiscent of the central-pair microtubules of the axoneme. The conoid ends in two preconoidal rings, and Toxoplasma SAS6L localizes to this region (orange). The conoid can be extended beyond or retracted into the APR (red), which serves as an MTOC for the subpellicular microtubules.

building flagella, they retain SFAs to ensure the reliable inheritance of apical organelles.

Given the observation that the SFAs and SAS6L are retained in aflagellate tachyzoites, an intriguing possibility is that SAS6L is one of the components that tether the SFA filament to the conoid. SAS6L cannot be the sole factor anchoring SFA to the apex, as our results indicate that SAS6L is not essential (although its loss is associated with a fitness defect) while loss of either SFA2 or SFA3 leads to a lethal arrest at the time of daughter cell budding. In light of our data on SAS6L and the observations on SFA2 and SFA3, we propose that the complete conoid evolved from an ancestral pseudoconoid in part by the incorporation of components from a vestigial apical flagellar apparatus. These components may have been from flagellum-associated structures, such as flagellar rootlets (as seen for the small portion of *Trypanosoma* SAS6L that is associated with the flagellar collar region), or could represent the direct subsumption of the flagellum into the conoid. The former premise suggests that flagellar components are retained because they are required for an essential cellular process, such as replication. In this context, SAS6L may have been retained along with the

basal-body-associated SFA proteins in order to coordinate organelle inheritance in aflagellar tachyzoites (88). The latter hypothesis implies direct homology between the central-pair microtubules of the axoneme and the microtubule pair at the center of the conoid and suggests that other orthologs of basal-plate and central-apparatus components may play a role in invasion or replication by coccidia.

ACKNOWLEDGMENTS

We thank Catherine Li and Ni Yao for technical assistance, David Coder for guidance with flow cytometry, and David Krueger for advice on Illustrator. Ke Hu (Indiana University) kindly provided us with the pmineGFP-TgICMAP1 construct. David Ferguson (Oxford) was a critically important sounding board for ideas about the apicomplexan basal body, and Susan Dutcher (Washington University) critically commented on the manuscript.

N.S. was supported by a graduate studentship from the EPA Trust, and research was funded by NIH grant AI067981 to N.S.M., a Wellcome Trust grant to K.G., and BBSRC New Investigator grant BB/J01477X/1 to B.W.

REFERENCES

- Dutcher SK. 2004. Dissection of basal body and centriole function in the unicellular green alga Chlamydomonas reinhardtii, p 71–92. *In* Nigg EA (ed), Centrosomes in development and disease. Wiley-VCH, Weinheim, Germany.
- Azimzadeh J, Marshall WF. 2010. Building the centriole. Curr. Biol. 20:R816–R825.
- 3. Marshall WF. 2009. Centriole evolution. Curr. Opin. Cell Biol. 21:14-19.
- Kobayashi T, Dynlacht BD. 2011. Regulating the transition from centriole to basal body. J. Cell Biol. 193:435–444.
- Azimzadeh J, Bornens M. 2007. Structure and duplication of the centrosome. J. Cell Sci. 120:2139–2142.
- Bornens M. 2008. Organelle positioning and cell polarity. Nat. Rev. Mol. Cell Biol. 9:874–886.
- 7. Paoletti A, Bornens M. 1997. Organisation and functional regulation of the centrosome in animal cells. Prog. Cell Cycle Res. **3:**285–299.
- 8. Chapman MJ, Dolan MF, Margulis L. 2000. Centrioles and kinetosomes: form, function, and evolution. Q. Rev. Biol. 75:409–429.
- 9. Schrevel J, Besse C. 1975. A functional flagella with a 6 + 0 pattern. J. Cell Biol. 66:492–507.
- Preble AM, Giddings TM, Jr, Dutcher SK. 2000. Basal bodies and centrioles: their function and structure. Curr. Top. Dev. Biol. 49:207–233.
- 11. Waters JC, Salmon E. 1997. Pathways of spindle assembly. Curr. Opin. Cell Biol. 9:37–43.
- Marshall WF, Vucica Y, Rosenbaum JL. 2001. Kinetics and regulation of de novo centriole assembly. Implications for the mechanism of centriole duplication. Curr. Biol. 11:308–317.
- Basto R, Lau J, Vinogradova T, Gardiol A, Woods CG, Khodjakov A, Raff JW. 2006. Flies without centrioles. Cell 125:1375–1386.
- Hinchcliffe EH, Miller FJ, Cham M, Khodjakov A, Sluder G. 2001. Requirement of a centrosomal activity for cell cycle progression through G₁ into S phase. Science 291:1547–1550.
- Piel M, Nordberg J, Euteneuer U, Bornens M. 2001. Centrosomedependent exit of cytokinesis in animal cells. Science 291:1550–1553.
- Mignot J-P. 1996. The centrosomal big bang: from a unique central organelle towards a constellation of MTOCs. Biol. Cell 86:81–91.
- 17. Despommier D, RG, Hotez P, Knirsch C. 2005. Parasitic diseases, fifth edition. Apple Trees Productions, LLC, New York, NY.
- Kohl L, Gull K. 1998. Molecular architecture of the trypanosome cytoskeleton. Mol. Biochem. Parasitol. 93:1–9.
- 19. Briggs LJ, Davidge JA, Wickstead B, Ginger ML, Gull K. 2004. More than one way to build a flagellum: comparative genomics of parasitic protozoa. Curr. Biol. 14:R611–R612.
- Gull K, Briggs L, Vaughan S. 2004. Basal bodies and microtubule organization in pathogenic protozoa, p 401–423. *In* Nigg EA (ed), Centrosomes in development and disease. Wiley-VCH, Weinheim, Germany.
- Gull K. 1999. The cytoskeleton of trypanosomatid parasites. Annu. Rev. Microbiol. 53:629–655.
- Seeback T, Hemphill A, Lawson D. 1990. The cytoskeleton of trypanosomes. Parasitol. Today 6:49–52.

- Scott V, Sherwin T, Gull K. 1997. Gamma-tubulin in trypanosomes: molecular characterisation and localisation to multiple and diverse microtubule organising centres. J. Cell Sci. 110(Pt 2):157–168.
- Ferguson DJP, Hutchison WM, Dunachie JF, Siim JC. 1974. Ultrastructural study of early stages of asexual multiplication and microgametogony of *Toxoplasma gondii* in the small intestine of the cat. Acta Pathol. Microbiol. Scand. Sect. B Microbiol. Immunol. 82B:167–181.
- Frixione E, Mondragon R, Meza I. 1996. Kinematic analysis of *Toxoplasma gondii* motility. Cell Motil. Cytoskeleton 34:152–163.
- Håkansson S, Morisaki H, Heuser J, Sibley LD. 1999. Time-lapse video microscopy of gliding motility in *Toxoplasma gondii* reveals a novel, biphasic mechanism of cell locomotion. Mol. Biol. Cell 10:3539–3547.
- Morrissette NS, Sibley LD. 2002. Disruption of microtubules uncouples budding and nuclear division in *Toxoplasma gondii*. J. Cell Sci. 115:1017– 1025.
- Nichols BA, Chiappino ML. 1987. Cytoskeleton of *Toxoplasma gondii*. J. Protozool. 34:217–226.
- 29. Shaw MK, Compton HL, Roos DS, Tilney LG. 2000. Microtubules, but not actin filaments, drive daughter cell budding and cell division in *Toxoplasma gondii*. J. Cell Sci. 113:1241–1254.
- Russell DG, Burns RG. 1984. The polar ring of coccidian sporozoites: a unique microtubule-organizing centre. J. Cell Sci. 65:193–207.
- Tran JQ, de Leon JC, Li C, Huynh MH, Beatty W, Morrissette NS. 2010. RNG1 is a late marker of the apical polar ring in *Toxoplasma gondii*. Cytoskeleton (Hoboken) 67:586–598.
- Sibert GJ, Speer CA. 1981. Fine structure of nuclear division and microgametogony of *Eimeria nieschulzi* Dieben, 1924. Z. Parasitenkd. 66: 179–189.
- Dubremetz JF, Elsner YY. 1979. Ultrastructural study of schizogony of Eimeria bovis in cell cultures. J. Protozool. 26:367–376.
- Gubbels MJ, Vaishnava S, Boot N, Dubremetz JF, Striepen B. 2006. A MORN-repeat protein is a dynamic component of the *Toxoplasma gondii* cell division apparatus. J. Cell Sci. 119:2236–2245.
- Brooks CF, Francia ME, Gissot M, Croken MM, Kim K, Striepen B. 2011. *Toxoplasma gondii* sequesters centromeres to a specific nuclear region throughout the cell cycle. Proc. Natl. Acad. Sci. U. S. A. 108:3767– 3772.
- Del Carmen MG, Mondragon M, Gonzalez S, Mondragon R. 2009. Induction and regulation of conoid extrusion in *Toxoplasma gondii*. Cell. Microbiol. 11:967–982.
- Mondragon R, Frixione E. 1996. Ca(2+)-dependence of conoid extrusion in *Toxoplasma gondii* tachyzoites. J. Eukaryot. Microbiol. 43:120– 127.
- Monteiro VG, de Melo EJ, Attias M, de Souza W. 2001. Morphological changes during conoid extrusion in *Toxoplasma gondii* tachyzoites treated with calcium ionophore. J. Struct. Biol. 136:181–189.
- Morrissette NS, Murray JM, Roos DS. 1997. Subpellicular microtubules associate with an intramembranous particle lattice in the protozoan parasite Toxoplasma gondii. J. Cell Sci. 110(Pt 1):35–42.
- Sheffield HG. 1966. Electron microscope study of the proliferative form of Besnoitia jellisoni. J. Parasitol. 52:583–594.
- Culver BP, Meehl JB, Giddings TH, Jr, Winey M. 2009. The two SAS-6 homologs in *Tetrahymena thermophila* have distinct functions in basal body assembly. Mol. Biol. Cell 20:1865–1877.
- 42. Leidel S, Delattre M, Cerutti L, Baumer K, Gonczy P. 2005. SAS-6 defines a protein family required for centrosome duplication in *C. elegans* and in human cells. Nat. Cell Biol. 7:115–125.
- Pelletier L, O'Toole E, Schwager A, Hyman AA, Muller-Reichert T. 2006. Centriole assembly in *Caenorhabditis elegans*. Nature 444:619–623.
- Dammermann A, Muller-Reichert T, Pelletier L, Habermann B, Desai A, Oegema K. 2004. Centriole assembly requires both centriolar and pericentriolar material proteins. Dev. Cell 7:815–829.
- 45. Nakazawa Y, Hiraki M, Kamiya R, Hirono M. 2007. SAS-6 is a cartwheel protein that establishes the 9-fold symmetry of the centriole. Curr. Biol. 17:2169–2174.
- Stevens NR, Roque H, Raff JW. 2010. DSas-6 and Ana2 coassemble into tubules to promote centriole duplication and engagement. Dev. Cell 19: 913–919.
- Blachon S, Cai X, Roberts KA, Yang K, Polyanovsky A, Church A, Avidor-Reiss T. 2009. A proximal centriole-like structure is present in *Drosophila* spermatids and can serve as a model to study centriole duplication. Genetics 182:133–144.
- 48. Rodrigues-Martins A, Riparbelli M, Callaini G, Glover DM, Betten-

court-Dias M. 2008. From centriole biogenesis to cellular function: centrioles are essential for cell division at critical developmental stages. Cell Cycle 7:11–16.

- Rodrigues-Martins A, Bettencourt-Dias M, Riparbelli M, Ferreira C, Ferreira I, Callaini G, Glover DM. 2007. DSAS-6 organizes a tube-like centriole precursor, and its absence suggests modularity in centriole assembly. Curr. Biol. 17:1465–1472.
- Kitagawa D, Vakonakis I, Olieric N, Hilbert M, Keller D, Olieric V, Bortfeld M, Erat MC, Fluckiger I, Gonczy P, Steinmetz MO. 2011. Structural basis of the 9-fold symmetry of centrioles. Cell 144:364–375.
- 51. van Breugel M, Hirono M, Andreeva A, Yanagisawa HA, Yamaguchi S, Nakazawa Y, Morgner N, Petrovich M, Ebong IO, Robinson CV, Johnson CM, Veprintsev D, Zuber B. 2011. Structures of SAS-6 suggest its organization in centrioles. Science 331:1196–1199.
- Huynh MH, Carruthers VB. 2009. Tagging of endogenous genes in a Toxoplasma gondii strain lacking Ku80. Eukaryot. Cell 8:530-539.
- Roos DS, Donald RG, Morrissette NS, Moulton AL. 1994. Molecular tools for genetic dissection of the protozoan parasite *Toxoplasma gondii*. Methods Cell Biol. 45:27–63.
- Brun R, Schonenberger. 1979. Cultivation and in vitro cloning or procyclic culture forms of *Trypanosoma brucei* in a semi-defined medium. Short communication. Acta Trop. 36:289–292.
- Wirtz E, Leal S, Ochatt C, Cross GA. 1999. A tightly regulated inducible expression system for conditional gene knock-outs and dominantnegative genetics in *Trypanosoma brucei*. Mol. Biochem. Parasitol. 99:89– 101.
- Kelly S, Reed J, Kramer S, Ellis L, Webb H, Sunter J, Salje J, Marinsek N, Gull K, Wickstead B, Carrington M. 2007. Functional genomics in *Trypanosoma brucei*: a collection of vectors for the expression of tagged proteins from endogenous and ectopic gene loci. Mol. Biochem. Parasitol. 154:103–109.
- Hartmann J, Hu K, He CY, Pelletier L, Roos DS, Warren G. 2006. Golgi and centrosome cycles in *Toxoplasma gondii*. Mol. Biochem. Parasitol. 145:125–127.
- Gilbert LA, Ravindran S, Turetzky JM, Boothroyd JC, Bradley PJ. 2007. Toxoplasma gondii targets a protein phosphatase 2C to the nuclei of infected host cells. Eukaryot. Cell 6:73–83.
- Ma C, Tran J, Li C, Ganesan L, Wood D, Morrissette N. 2008. Secondary mutations correct fitness defects in *Toxoplasma gondii* with dinitroaniline resistance mutations. Genetics 180:845–856.
- Hodges ME, Scheumann N, Wickstead B, Langdale JA, Gull K. 2010. Reconstructing the evolutionary history of the centriole from protein components. J. Cell Sci. 123:1407–1413.
- Katoh K, Kuma K, Toh H, Miyata T. 2005. MAFFT version 5: improvement in accuracy of multiple sequence alignment. Nucleic Acids Res. 33: 511–518.
- Ronquist F, Huelsenbeck JP. 2003. MrBayes 3: Bayesian phylogenetic inference under mixed models. Bioinformatics 19:1572–1574.
- 63. Guindon S, Gascuel O. 2003. A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. Syst. Biol. 52:696–704.
- Heaslip AT, Ems-McClung SC, Hu K. 2009. TgICMAP1 is a novel microtubule binding protein in *Toxoplasma gondii*. PLoS One 4:e7406. doi:10.1371/journal.pone.0007406.
- Stechmann A, Cavalier-Smith T. 2002. Rooting the eukaryote tree by using a derived gene fusion. Science 297:89–91.
- 66. Sinden R, Talman A, Marques S, Wass M, Sternberg M. 2010. The flagellum in malarial parasites. Curr. Opin. Microbiol. 13:491–500.
- 67. Mueller BE, Desser SS, Haberkorn A. 1981. Ultrastructure of developing gamonts of *Eimeria contorta* Haberkorn, 1971 (Protozoa, Sporozoa) with emphasis on the host-parasite interface. J. Parasitol. **67**:487–495.
- Ferguson DJP, Birch-Andersen A, Hutchison WM, Siim JC. 1980. Ultrastructural observations on microgametogenesis and the structure of the microgamete of *Isospora felis*. Acta Pathol. Microbiol. Scand. Sect. B Microbiol. 88B:151–159.
- 69. Scholtyseck E, Mehlhorn H, Hammond DM. 1972. Electron microscope

studies of microgametogenesis in coccidia and related groups. Z Parasitenkd. 38:95–131.

- 70. Young JA, Fivelman QL, Blair PL, de la Vega P, Le Roch KG, Zhou Y, Carucci DJ, Baker DA, Winzeler EA. 2005. The *Plasmodium falciparum* sexual development transcriptome: a microarray analysis using ontologybased pattern identification. Mol. Biochem. Parasitol. 143:67–79.
- Keeley A, Soldati D. 2004. The glideosome: a molecular machine powering motility and host-cell invasion by Apicomplexa. Trends Cell Biol. 14:528–532.
- 72. Sibley LD. 2004. Intracellular parasite invasion strategies. Science 304: 248–253.
- Baum J, Richard D, Healer J, Rug M, Krnajski Z, Gilberger TW, Green JL, Holder AA, Cowman AF. 2006. A conserved molecular motor drives cell invasion and gliding motility across malaria life cycle stages and other apicomplexan parasites. J. Biol. Chem. 281:5197–5208.
- Morrissette NS, Sibley LD. 2002. Cytoskeleton of apicomplexan parasites. Microbiol. Mol. Biol. Rev. 66:21–38.
- Hu K, Roos DS, Murray JM. 2002. A novel polymer of tubulin forms the conoid of *Toxoplasma gondii*. J. Cell Biol. 156:1039–1050.
- Cavalier-Smith T, Chao EE. 2004. Protalveolate phylogeny and systematics and the origins of Sporozoa and dinoflagellates (phylum Myzozoa nom. nov.). Eur. J. Protistol. 40:185–212.
- Leander BS, Keeling PJ. 2003. Morphostasis in alveolate evolution. Trends Ecol. Evol. 18:395–402.
- Leander BS, Kuvardina ON, Aleshin VV, Mylnikov AP, Keeling PJ. 2003. Molecular phylogeny and surface morphology of *Colpodella edax* (Alveolata): insights into the phagotrophic ancestry of apicomplexans. J. Eukaryot. Microbiol. 50:334–340.
- 79. **Brugerolle G.** 2002. *Cryptophagus subtilis*: a new parasite of cryptophytes affiliated with the Perkinsozoa lineage. Eur. J. Protistol. **37**:379–390.
- Brugerolle G. 2002. Colpodella vorax: ultrastructure, predation, life-cycle, mitosis, and phylogenetic relationships. Eur. J. Protistol. 38:113–125.
- Simpson A, Patterson D. 1996. Ultrastructure and identification of the predatory flagellate *Colpodella pugnax* Cienkowski (Apicomplexa) with a description of Colpodella turpis n. sp. and a review of the genus. Syst. Parasitol. 33:187–198.
- Salisbury JL. 1995. Centrin, centrosomes, and mitotic spindle poles. Curr. Opin. Cell Biol. 7:39–45.
- Geimer S, Clees J, Melkonian M, Lechtreck KF. 1998. A novel 95-kD protein is located in a linker between cytoplasmic microtubules and basal bodies in a green flagellate and forms striated filaments in vitro. J. Cell Biol. 140:1149–1158.
- Lechtreck KF, Frins S, Bilski J, Teltenkotter A, Weber K, Melkonian M. 1996. The cruciated microtubule-associated fibers of the green alga *Dunaliella bioculata* consist of a 31 kDa SF-assemblin. J. Cell Sci. 109(Pt 4):827–835.
- Lechtreck KF, Melkonian M. 1991. Striated microtubule-associated fibers: identification of assemblin, a novel 34-kD protein that forms paracrystals of 2-nm filaments in vitro. J. Cell Biol. 115:705–716.
- Harper JD, Thuet J, Lechtreck KF, Hardham AR. 2009. Proteins related to green algal striated fiber assemblin are present in stramenopiles and alveolates. Protoplasma 236:97–101.
- Lechtreck KF. 2003. Striated fiber assemblin in apicomplexan parasites. Mol. Biochem. Parasitol. 128:95–99.
- 88. Francia ME, Jordan CN, Patel JD, Sheiner L, Demerly JL, Fellows JD, de Leon JC, Morrissette NS, Dubremetz JF, Striepen B. 2012. Cell division in apicomplexan parasites is organized by a homolog of the striated rootlet fiber of algal flagella. PLoS Biol. 10:e1001444. doi:10.1371 /journal.pbio.1001444.
- Finn RD, Mistry J, Tate J, Coggill P, Heger A, Pollington JE, Gavin OL, Gunasekaran P, Ceric G, Forslund K, Holm L, Sonnhammer EL, Eddy SR, Bateman A. 2010. The Pfam protein families database. Nucleic Acids Res. 38:D211–D222.
- 90. Lupas A, Van Dyke M, Stock J. 1991. Predicting coiled coils from protein sequences. Science 252:1162–1164.