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Characterisation of TbSmee1 suggests endocytosis allows surface-bound cargo to enter the trypanosome flagellar pocket

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Summary: Characterisation of the essential trypanosome protein TbSmee1 suggests that endocytosis is required for flagellar pocket access of surface-bound cargo.

Abstract

All endo- and exocytosis in the African trypanosome *Trypanosoma brucei* occurs at a single subdomain of the plasma membrane. This subdomain, the flagellar pocket, is a small vase-

shaped invagination containing the root of the cell's single flagellum. Several cytoskeletonassociated multiprotein complexes are coiled around the neck of the flagellar pocket on its cytoplasmic face. One of these, the hook complex, was proposed to affect macromolecule entry into the flagellar pocket lumen. In previous work, knockdown of the hook complex component TbMORN1 resulted in larger cargo being unable to enter the flagellar pocket. In this study, the hook complex component TbSmee1 was characterised in bloodstream form *Trypanosoma brucei* and was found to be essential for cell viability. TbSmee1 knockdown resulted in flagellar pocket enlargement and impaired access to the flagellar pocket membrane by surface-bound cargo, similar to depletion of TbMORN1. Unexpectedly, inhibition of endocytosis by knockdown of clathrin phenocopied TbSmee1 knockdown, suggesting that endocytic activity itself is a prerequisite for the entry of surface-bound cargo into the flagellar pocket.

Introduction

The flagellated parasitic protist *Trypanosoma brucei* lives in the bloodstream of its mammalian hosts, in continuous exposure to the immune system. Endo- and exocytosis are used to remove bound antibodies from the cell surface and scavenge macromolecular nutrients from the surroundings (Borst and Fairlamb, 1998; Engstler et al., 2007; Webster et al., 1990). Remarkably, all endo- and exocytosis in *T. brucei* occurs at just a single subdomain of the plasma membrane (Engstler et al., 2004; Grunfelder et al., 2003). This subdomain is a small vase-shaped invagination, called the flagellar pocket, that houses the root of the cell's single flagellum (Halliday et al., 2021; Lacomble et al., 2009).

Molecules that enter the flagellar pocket, either in the fluid phase or when bound to the surface of the parasite (here "surface-bound"), are rapidly internalised by clathrin-mediated endocytosis and then routed to the endosomal/lysosomal system (Link et al., 2021; Overath and Engstler, 2004). Depletion of clathrin results in a gross enlargement of the flagellar pocket due to an imbalance between endocytosis and exocytosis, and causes cargo accumulation inside the flagellar pocket (Allen et al., 2003). Although endocytosis of fluid phase and surface-bound cargo is relatively well-characterised in *T. brucei*, the exact mechanism(s) by which cargo initially enters the flagellar pocket are less well-understood.

Coiled around the neck of the flagellar pocket on its cytoplasmic face are a number of poorly-characterised cytoskeleton-associated complexes, which might contribute to these processes (Esson et al., 2012; Halliday et al., 2019)(Fig. 1A). The flagellar pocket collar is a multiprotein complex shaped like a cuff bracelet, and demarcates the boundary between the

flagellar pocket and the flagellar pocket neck. It contains the protein TbBILBO1, and a number of its other components such as FPC4 and BILBO2 have recently been characterised (Albisetti et al., 2017; Bonhivers et al., 2008; Florimond et al., 2015; Isch et al., 2021). The centrin arm is a centrin-containing kinked rod which appears to be involved in the biogenesis of various cytoskeleton-associated structures (Selvapandiyan et al., 2007; Shi et al., 2008). The hook complex is a hook-shaped multiprotein complex that sits atop the flagellar pocket collar and alongside the centrin arm. Characterised components include the proteins TbMORN1, BOH1, BOH2, and Bhalin (Broster Reix et al., 2021; Morriswood et al., 2009; Pham et al., 2020; Pham et al., 2019).

Previous characterisation of TbMORN1 revealed that it is essential for the viability of bloodstream form *T. brucei* cells (Morriswood and Schmidt, 2015). Cells depleted of TbMORN1 have enlarged flagellar pockets, indicative of an endocytosis defect. In addition, although small endocytic reporters such as 10 kDa dextran (hydrodynamic radius ~1.86 nm) were still capable of entering the enlarged flagellar pocket, the access of larger macromolecules such as the lectin concanavalin A (ConA, hydrodynamic radius ~4.2 nm) or bovine serum albumin (BSA, hydrodynamic radius ~ 3.51 nm) conjugated to 5 nm gold particles was blocked (Ahmad et al., 2007; Armstrong et al., 2004; Morriswood and Schmidt, 2015). This phenotype was also observed upon depletion of the hook complex protein Bhalin (Broster Reix et al., 2021). On this basis, it was proposed that the hook complex might be regulating the passage of large macromolecules through the flagellar pocket neck, and thereby mediating cargo entry into the flagellar pocket (Morriswood and Schmidt, 2015).

In this study, the hook complex protein TbSmee1 was characterised in bloodstream form *T. brucei.* TbSmee1 (Tb927.10.8820) was initially identified in a proximity labelling screen using TbMORN1 as bait, and shown to localise to the shank part of the hook complex (Morriswood et al., 2013). It was subsequently characterised in procyclic form *T. brucei*, the life cycle stage found within the tsetse fly vector (Perry et al., 2018). In bloodstream form *T. brucei*, depletion of TbSmee1 phenocopied TbMORN1 and Bhalin, with ConA, BSA and surface-bound antibodies unable to access the enlarged flagellar pocket. Surprisingly, clathrin-depleted cells also exhibited this flagellar pocket access phenotype, suggesting that the observed effects following depletion of the hook complex proteins are due to impaired endocytosis, and therefore that endocytosis is required for the entry of surface-bound cargo into the flagellar pocket.

Results

TbSmee1 is a cytoskeleton-associated phosphoprotein.

Bioinformatic analysis predicted that TbSmee1 is composed of three structured domains separated by linker regions (Fig. 1B). A multiple sequence alignment of the TbSmee1 protein with orthologues from other trypanosome species indicated the presence of three blocks of highly-conserved sequence, corresponding to regions of predicted secondary structure (Fig. S1). Three-dimensional structural prediction by AlphaFold suggested a close association between the first two folded domains (https://alphafold.ebi.ac.uk/entry/Q38A92) (Jumper et al., 2021; Wheeler, 2021).

Two polyclonal antibodies (303, 304) were raised against a recombinant TbSmee1(1-400) truncation, and a third (508) was raised against two peptides (Fig. 1B, green lines). The specificity of the antibodies was confirmed using immunoblots of whole-cell lysates obtained from TbSmee1 RNAi cells and 3xTy1-TbSmee1 endogenous replacement cells. In situ tagging of the *SMEE1* gene in the 3xTy1-TbSmee1 cells was confirmed by PCR analysis of genomic DNA (Fig. S2A,B). All three antibodies detected a protein of ~ 85 kDa in the immunoblots (Fig. 1C, white arrows). This protein was strongly depleted in the RNAi (Tet +) samples (Fig. 1C). An extra band corresponding to 3xTy1-TbSmee1 was seen in lysates from the endogenous replacement (End. repl.) cells (Fig. 1C). This extra band was also detected using anti-Ty1 (BB2) antibodies, confirming its identity as Ty1-TbSmee1 (Fig 1C, lower panels).

To investigate TbSmee1 association with the cytoskeleton, cells were fractionated using non-ionic detergent (Fig. 1D). The fractions were analysed by immunoblotting, using PFR1,2 as markers for the cytoskeleton, and the ER chaperone BiP as a marker for the cytoplasm. TbSmee1 co-fractionated with PFR1,2, confirming that it was cytoskeleton-associated (Fig. 1E). Interestingly, the fuzzy appearance of the TbSmee1 band observed in immunoblots of whole-cell lysates was reduced or absent in the fractionation blots (compare Fig. 1C and Fig. 1E). TbSmee1 is heavily phosphorylated in vivo, and is a substrate and potential binding partner of the mitotic kinase TbPLK (Benz and Urbaniak, 2019; McAllaster et al., 2015; Nett et al., 2009; Urbaniak et al., 2013). The existence of different phosphoforms of a protein is known to cause a fuzzy appearance of bands in gels, so it seemed possible that exposure of TbSmee1 to endogenous phosphatases during the extraction step was causing dephosphorylation.

To investigate whether the band collapse could be attributed to dephosphorylation of TbSmee1, extracted cytoskeletons were incubated with exogenous phosphatase at various

timepoints prior to immunoblotting. Incubation with exogenous phosphatase resulted in a progressive collapse of the TbSmee1 band over a 20 min period (Fig. 1F). This band collapse was not seen when the samples were in the presence of phosphatase inhibitors either at RT or on ice, or kept on ice (Fig. 1F, last three lanes). The fuzzy appearance of TbSmee1 in immunoblots could therefore be attributed exclusively to phosphorylation.

TbSmee1 localises to the shank part of the hook complex.

The localisation of endogenous and tagged TbSmee1 protein was analysed in bloodstream form *T. brucei* cells using immunofluorescence microscopy (Fig. 1G). All three main stages of the cell division cycle (1K1N, 2K1N, 2K2N) were analysed. In 1K1N cells (i.e. those with a single kinetoplast, K, and nucleus, N), TbSmee1 localised to a single structure near the flagellum base, consistent with the position of the hook complex (Fig. 1G, arrow). In 2K1N and 2K2N cells there was a third subpopulation of TbSmee1 at varying distances along the cell body in addition to the replicated hook complex (Fig. 1G, arrowheads). The same TbSmee1 distributions were seen using all three anti-TbSmee1 antibodies; anti-Ty1 antibody labelling of 3xTy1-TbSmee1 cells also produced the same labelling pattern (Fig. S2C, D).

In colabelling experiments, TbSmee1 strongly overlapped with the shank part of the hook complex as labelled by TbMORN1 and TbLRRP1 (Fig. S3A, B). TbSmee1 also strongly overlapped with the hook complex protein Tb927.10.3010. This 133 kDa protein was one of the top hits in the BioID screen using TbMORN1 (Morriswood et al., 2013). Consistent with the convention started with TbSmee1, Tb927.10.3010 was named TbStarkey1 after another of Captain Hook's pirates. Two anti-peptide antibodies were generated against TbStarkey1, and their specificity was confirmed by immunoblotting and immunofluorescence imaging (Fig. S4A, B). Strong overlap was seen between TbSmee1 and TbStarkey1 on the shank part of the hook complex (Fig. S4C). As expected, TbSmee1 showed only a partial overlap with TbCentrin4, a marker of the centrin arm, and little overlap with TbBILBO1, a marker of the flagellar pocket collar (Fig. S3C, D).

To quantify the degree of colocalisation between TbSmee1 and the various hook complex and flagellar pocket collar marker proteins, correlation coefficients from the pairwise labelling experiments were calculated. Moderate correlation was seen between TbSmee1 and TbMORN1 and TbLRRP1, with lower values being measured for TbCentrin4 and TbBILBO1 (Fig. S3E).

TbSmee1 is transiently associated with the flagellum attachment zone (FAZ) tip.

The flagellum attachment zone (FAZ) is a cytoskeleton-associated apparatus that adheres the flagellum of *T. brucei* to the cell body (Sunter and Gull, 2016). During replication, a new FAZ is assembled and grows from its initiation point very close to the hook complex towards the anterior end of the cell (Sunter et al., 2015b; Zhou et al., 2015). It was previously shown that a tagged version of TbSmee1 is transiently localised to the tip of the new FAZ in replicating insect-stage *T. brucei* cells, in addition to its hook complex localisation (Perry et al., 2018).

Consistent with this, endogenous TbSmee1 showed a partial overlap with the posterior end of the FAZ (Fig. 2A, arrow). The extra TbSmee1 subpopulation in 2K1N and 2K2N cells overlapped with the tip of the new FAZ (Fig. 2A, arrowheads).

Using structured illumination microscopy, TbSmee1 was observed to be slightly in front of the tip of the new FAZ filament (Fig. 2B). This suggested that TbSmee1 might be present at the "groove". The groove, which can be detected using the monoclonal antibody DOT1, is a structure involved in remodelling of the microtubule cytoskeleton during cell replication in bloodstream form *T. brucei* (Hughes et al., 2013; Smithson et al., 2022). TbSmee1 at the FAZ tip appeared to be enveloped by the DOT1 labelling (Fig. 2C). In summary, in replicating cells TbSmee1 is present at the groove in addition to the hook complex, and travels in front of the newly-assembling FAZ filament.

The C-terminal part of TbSmee1 is required for targeting to the hook complex.

To determine what parts of TbSmee1 primary structure are responsible for targeting to the hook complex, a series of truncation constructs based on the predicted domain architecture of TbSmee1 were designed (Fig. 3A). Cell lines were generated that could inducibly express each of these truncations with an N-terminal Ty1 tag. The presence of the ectopic transgenes in the genomes of these cells was confirmed by PCR analysis of genomic DNA (Fig. S5A). To determine the localisation of each construct, and whether it could associate with the cytoskeleton, detergent-extracted cells were analysed using immunofluorescence microscopy. It should be noted that both endogenous *SMEE1* alleles were still present, meaning that targeting of the truncation constructs was being assayed in the presence of the endogenous Smee1 protein.

TbSmee1(161-766) correctly localised to the hook complex, indicating that Domain1 is not necessary for targeting (Fig. 3B). Of note, no dominant negative effects on cell growth were observed upon overexpression of the TbSmee1(161-766) construct. TbSmee1(265-766)

also localised correctly, indicating that the predicted linker region between Domain1 and Domain2 is not necessary for localisation (Fig. 3B). No other truncations were observed to localise to the hook complex. Expression of all TbSmee1 truncations was confirmed by immunoblotting with anti-TbSmee1 and anti-Ty1 antibodies (Figs 3C, S5A). Quantification of the immunoblots indicated that most were expressed at around the level of the endogenous protein (Fig. 3D).

Interestingly, TbSmee1(161-766) also localised to the FAZ tip in replicating cells (Fig. S5B). TbSmee1(265-766) did not localise to the FAZ tip, suggesting that the linker region between Domain1 and Domain2 (aa 161-264) might be required for targeting to this structure. In support of this hypothesis, TbSmee1(2-400) was found to localise to the FAZ tip despite not localising to the hook complex (Fig. S5C).

TbSmee1 is essential for the viability of BSF *T. brucei*, and its depletion causes gross enlargement of the flagellar pocket.

The effects of TbSmee1 depletion were analysed using tetracycline-inducible RNAi. Induction of RNAi resulted in a rapid cessation of population growth after around 24 h (Fig. 4A). Visual inspection of the stalled populations at 48 h and 72 h post-induction showed widespread lysis, confirming that TbSmee1 is essential for the viability of bloodstream form cells in vitro. A shorter timecourse with higher sampling frequency indicated that the growth defect began after around 20 h of RNAi, and was already clear at 24 h (Fig. 4A, inset).

TbSmee1 depletion was confirmed at the single-cell level using immunofluorescence microscopy (Fig. 4B). TbSmee1 signal was lost from both the hook complex and FAZ tip. The kinetics of TbSmee1 protein depletion on either side of the onset of the growth defect were assessed by immunoblotting. TbSmee1 protein levels were reduced to ~ 20-25% at 15 h and 18 h post-induction, with a further reduction to around 10-15% from 21 h onwards (Fig. 4C).

The effect of TbSmee1 knockdown on cell cycle progression was assessed by quantifying the numbers of 1K1N, 2K1N, 2K2N, and abnormal cell types in the same time window (Fig. S6A).

An increase in 2K1N and a decrease in 1K1N cells was visible from 21 h post-induction onwards, followed by an increase in 2K2N cells at the 24 h timepoint. This indicated that cell cycle progression was inhibited from around 21 h after induction of RNAi, correlating with the onset of the growth defect. To summarise, TbSmee1 protein was already significantly

depleted at 15 h, with subsequent effects on population growth and cell cycle progression visible from around 21 h onwards.

The effects of TbSmee1 depletion on a panel of hook complex proteins and components of the centrin arm and flagellar pocket collar were systematically evaluated (Fig. S7A-E). TbSmee1 depletion for 24 h did not result in observable effects on the expression levels or localisation of any of the candidates or on flagellum attachment. Conversely, depletion of TbMORN1 for just 16 h resulted in reductions in the levels of TbSmee1 and TbStarkey1 (Fig. S7B, F-H).

To determine the ultrastructural changes caused by TbSmee1 depletion, the cells were imaged by electron microscopy after high-pressure freezing, which gives better morphological preservation than chemically fixing the cells in the growth media. The most obvious morphological effect of TbSmee1 depletion was the gradual accumulation of cells with enlarged flagellar pockets (Fig. 4D). This presumably was the cause of the lethality phenotype, as depletion of TbMORN1 was also shown to result in progressive enlargement of the flagellar pocket until cells rounded up and lysed (Morriswood and Schmidt, 2015). Depletion of TbStarkey1 also frequently resulted in the generation of cells with enlarged flagellar pockets, despite causing no growth defect (Fig. S6B-D). This, together with the recently-published characterisation of Bhalin (Broster Reix et al., 2021), means that depletion of four separate hook complex proteins - TbMORN1, TbSmee1, TbStarkey1, Bhalin - results in flagellar pocket enlargement, though with varying magnitudes of effect.

Flagellar pocket enlargement is an early consequence of TbSmee1 depletion.

Although flagellar pocket enlargement appears to be a consistent phenotype resulting from hook complex protein depletion, it is not an uncommon phenotype in bloodstream form RNAi cells. Importantly, it can be either a direct or an indirect consequence of protein depletion (Ali et al., 2014; Allen et al., 2003; Hall et al., 2004; Hall et al., 2005; Price et al., 2007).

To determine whether flagellar pocket enlargement was an early/direct consequence of TbSmee1 depletion (i.e. occurring very soon after or even before the onset of the growth defect), flagellar pocket size was assayed using a fluorescent, fixable 10 kDa dextran reporter. Dextran is a polysaccharide that traffics in the fluid phase and is well-established as an endocytic marker. Control (- Tet) and TbSmee1-depleted (+ Tet) cells were incubated on ice in order to block endocytosis (Brickman et al., 1995). The cells were then incubated with the labelled dextran for 15 min to allow it to enter the flagellar pocket, and fixed afterwards (Fig. 5A). The magnitude of the dextran signal should therefore be proportional to flagellar

pocket volume. Visual analysis of the cells confirmed that dextran visibly labelled the flagellar pocket, with a much greater signal seen for cells with an enlarged flagellar pocket (Fig. 5B).

Flow cytometry was then used for high-throughput, unbiased, and quantitative analysis of the cells (Fig. 5C). At 15 h post-induction, no difference between TbSmee1-depleted cells and controls was observed. At 18 h and 21 h post-induction a slight "shoulder" on the +Tet traces became visible, indicating the emergence of cells with higher fluorescence values than in controls. By 24 h post-induction there was a clear hump visible in the traces, indicating a subpopulation with fluorescence values sometimes two orders of magnitude greater than those in control cells (Fig. 5C, arrow). Quantification of the flow cytometry data from multiple experiments showed that average fluorescence intensity of the whole + Tet population was noticeably higher than controls at 21 h and 24 h post-induction (Fig. 5D). Even at 18 h post-induction, i.e. before the onset of the growth defect, there was already a clear increase in the average fluorescence intensity in the + Tet population (Fig. 5D, blue arrow). This strongly suggests that flagellar pocket enlargement is an early and probably direct consequence of TbSmee1 depletion.

TbSmee1 depletion results in impaired flagellar pocket access of surface-bound cargo.

It was previously shown that the ability of large cargo to enter the flagellar pocket is affected after knockdown of the hook complex protein TbMORN1 (Morriswood and Schmidt, 2015). Specifically, the fluid phase marker 10 kDa dextran accumulates in the enlarged flagellar pocket of TbMORN1-depleted cells, while larger fluid phase cargo such as BSA-5 nm gold and large surface-bound cargo such as ConA (which binds to surface glycoproteins) do not access the flagellar pocket lumen.

To test whether TbSmee1 knockdown also impairs flagellar pocket access, TbSmee1 RNAi cells were incubated with both dextran and ConA to simultaneously monitor the uptake of fluid phase and surface-bound cargo. The cells were first incubated on ice to block endocytosis, and then with the reporters (also on ice) to allow ConA to bind and dextran to enter the flagellar pocket. The cells were then shifted to 37°C to reactivate endocytosis, and subsequently fixed and imaged (Fig. 6A).

In control (- Tet) cells, both dextran and ConA strongly overlapped in the part of the cell corresponding to the endosomal/lysosomal system (Fig. 6B, -Tet, arrow). To confirm that the internalised material was being trafficked to the lysosome, the cells were labelled with

antibodies specific for the lysosomal enzyme p67 (Kelley et al., 1999). The dextran reporter was not compatible with immunolabelling, but the ConA signal clearly overlapped with the lysosome marker p67 (Fig. S8, -Tet cells). In TbSmee1-depleted (+ Tet) cells, dextran filled the enlarged flagellar pocket while ConA was restricted to one or two small foci that appeared to be on the cell surface (Fig. 6B, +Tet, arrow). No overlap was seen between the dextran and ConA labels, suggesting that ConA was not able to enter the flagellar pocket. In addition, no overlap was observed between ConA and the lysosome marker p67 (Fig. S8, +Tet cells).

To quantify these observations, the cells were grouped into 4 categories: 1, complete overlap between the two labels; 2, partial overlap between the two labels; 3, no overlap between the two labels; 4, whole-cell labelling. Whole-cell labelling occurs when the cell has lost integrity, and labelling is found throughout the cytoplasm. Control (-Tet) cells all showed complete overlap between the dextran and ConA, while cells depleted of TbSmee1 for 24 h showed >25% of cells with partial or no overlap between the cargoes (Fig. 6C).

The experiments were repeated using fluorescently-labelled BSA (Fig. 6D). BSA traffics in the fluid phase and is a physiological cargo, unlike ConA (Coppens et al., 1987). As expected, in control cells there was strong overlap between dextran and BSA from the endosomal/lysosomal system (Fig. 6E, -Tet, arrow). In TbSmee1-depleted cells, there was once again no overlap between the reporters, indicating that BSA was unable to access the enlarged flagellar pocket. Surprisingly, despite BSA being fluid-phase cargo there was a punctate signal analogous to that seen with ConA, suggesting that it could at least partly bind to the cell surface (Fig. 6E, +Tet, arrow). Quantification of the data showed that the effect on BSA was more pronounced than that seen for ConA, with >55% of TbSmee1-depleted cells showing no or only partial overlap between the two reporters after 24 h of RNAi (Fig. 6F).

Endocytosis is required for flagellar pocket access of surface-bound cargo.

The inability of surface-bound cargo to enter the flagellar pocket of TbMORN1- and TbSmee1-depleted cells could either be due to a defect in endocytosis (indicated by the enlargement of the flagellar pocket), or a secondary effect.

To distinguish between these possibilities, the assays were repeated following clathrin depletion (Fig. 7). Clathrin is an essential endocytic coat protein and its loss results in a block in endocytosis. This results in the expected enlargement of the flagellar pocket and ultimately cell lysis, and in fact this phenotype was first described as a result of clathrin

depletion (Allen et al., 2003). In control cells, as expected, there was strong overlap between the dextran and ConA cargoes in the region of the endolysosomal system (Fig. 7B, -Tet, arrow). In clathrin-depleted cells however, there was again no overlap observed between the two reporters, and ConA did not appear able to access the enlarged flagellar pocket (Fig. 7B, +Tet, arrow). Quantification of the data showed that >50% of clathrin-depleted cells showed either partial or no overlap between the two reporters after 19 h of RNAi (Fig. 7C). The same effect was seen in assays using dextran and BSA, with little to no overlap between the reporters in clathrin-depleted cells (Fig. 7D-F). These results indicated that inhibition of endocytosis by itself resulted in the failure of either ConA or BSA to enter the enlarged flagellar pocket.

Endocytosis is required for flagellar pocket access of surface-bound antibodies.

One of the main functions of endocytosis in trypanosomes is to remove any antibodies bound to the surface glycoprotein coat. To determine whether the effects observed above using ConA and BSA reporters also apply to antibodies, antibody uptake assays were carried out. As the uptake of surface-bound antibodies occurs in a matter of seconds, a slightly modified assay protocol was used (Fig. 8A). The cells were first incubated on ice to block endocytosis, and then incubated with antiserum specific for the variant surface glycoprotein (VSG), which accounts for the overwhelming majority of the surface binding of the antibodies. The remaining cells were washed (at low temperature), after which a second sample was fixed (t=1). The remaining cells were shifted to 37°C to reactivate endocytosis and incubated for 2 minutes to allow the antibodies to enter the flagellar pocket and then be internalised by endocytosis.

The distribution of anti-VSG antibodies in control and TbSmee1-depleted cells was analysed using microscopy (Fig. 8B). In control (-Tet) cells, anti-VSG signals were observed at the cell surface (t=0), then at the location of the flagellar pocket (t=1) and finally at the endosomal/lysosomal system after allowing endocytosis (t=2) (arrows, upper panels). In TbSmee1-depleted cells (+Tet), no anti-VSG signals were observed in the enlarged flagellar pockets (t=1, t=2) (arrows, lower panels). There was considerable loss of anti-VSG signal between the t=0 and t=2 timepoints, and half of the TbSmee1-depleted cells displaying a phenotype had no observable signal at all, making the conclusions from these experiments somewhat tentative.

To quantify the results, the incidence of morphologically abnormal cells was counted, and then the various distributions of anti-VSG antibodies were classified into three categories

(Fig. 8C). Of the 35% of cells in the TbSmee1-depleted population that displayed a phenotype after 24 h of RNAi, 26% were classified as being completely rounded up ("BigEye") while 7% had the less-developed "enlarged pocket" state. The anti-VSG antibody distributions were grouped into three categories, respectively either a dot signal, another signal (usually a dot with other weaker labelling), and no signal. Although most of the morphologically-abnormal cells had no anti-VSG signal at all, TbSmee1-depleted cells with clearly internalised anti-VSG of the kind observed in control cells were never seen.

In summary, the inhibition of endocytosis caused by depletion of TbSmee1 prevents the internalisation of surface-bound anti-VSG antibodies, and appears to also prevent their entry into the enlarged flagellar pocket.

Discussion

In this study, the hook complex component TbSmee1 was characterised in bloodstream form *T. brucei.* It is only the third hook complex protein to be characterised in this life cycle stage after TbMORN1 and Bhalin, and like them is essential for the viability of the cells in vitro (Broster Reix et al., 2021; Morriswood and Schmidt, 2015). The localisation of TbSmee1 was found to be identical to that previously documented in the procyclic life cycle stage of *T. brucei* (Figs S3, 2) (Perry et al., 2018). It is constitutively present at the shank part of the hook complex, and additionally localises to the tip of the growing new FAZ in replicating cells. At the time of the preprint of this work being published, TbSmee1 was the first protein besides the DOT1 antigen and FLAM3 to be shown to localise to the groove structure associated with the tip of the growing new FAZ (Sunter et al., 2015a). A number of additional proteins present at the groove have since been identified (Smithson et al., 2022). It is unclear what role phosphorylation of TbSmee1 plays in its localisation, but given that it is both a substrate and putative direct binding partner of the mitotic kinase TbPLK, it is possible that its phosphorylation status may be related to its localisation to the groove structure.

Phenotypic characterisation of TbSmee1 showed that its depletion results in the enlargement of the flagellar pocket with concomitant effects on the entry of macromolecular cargo (Figs 4-6). There was extremely good agreement between the immunoblotting, growth curve, and cell cycle analysis data. At 18 h post-induction of RNAi, TbSmee1 protein levels were reaching their minimum but there was as yet no observable effect on cell cycle progression or population growth. From around 20-21 h post-induction, when TbSmee1 levels reached 10-15% of controls, cell cycle progression slowed, resulting in an increase in 2K1N and 2K2N cells. This was mirrored by a slowing of population growth.

analysis of a number of marker proteins for the hook complex, the centrin arm, and the flagellar pocket collar showed that none were strongly affected in terms of their protein levels or localisation by the depletion of TbSmee1. The lack of observable structural changes to the hook complex, flagellar pocket collar, and centrin arm following TbSmee1 depletion implied that the lethal phenotype might be due to a loss of a specific cellular function, rather than destabilisation of structural complexes.

In the previous characterisation of TbSmee1 in the procyclic form life cycle stage, depletion of TbSmee1 was found to cause structural changes to the hook complex and altered TbMORN1 distribution patterns (Perry et al., 2018). These analyses were however done after 6 days of RNAi induction, when TbSmee1 levels were already undetectable from 1 day onwards (a growth defect was observable from 3 days of RNAi onwards). It is very likely that similar changes would be observed in the bloodstream form stage at later RNAi timepoints, although the essential nature of TbSmee1 in bloodstream form cells might make this analysis difficult.

Flow cytometry analysis indicated that the flagellar pocket enlargement began around 18 h post-induction, prior to the effects on population growth and cell cycle progression. This therefore suggests that flagellar pocket enlargement is an early and likely direct consequence of TbSmee1 depletion, with the effects on cell cycle progression coming afterwards. One plausible hypothesis to account for these observations is that the spatial problems produced by flagellar pocket enlargement impair cell cycle progression. The relative lack of cell cycle checkpoints means that "monster" cells with re-duplicated organelles are then produced. The internal pressure caused by continuing enlargement of the flagellar pocket forces the cells into an increasingly spherical shape and eventually cell viability is lost.

The enlargement of the flagellar pocket was accompanied by a cargo access defect previously documented for TbMORN1 and Bhalin - small fluid-phase cargo such as 10 kDa dextran filled the enlarged flagellar pocket, while larger cargo such a ConA and BSA did not (Broster Reix et al., 2021; Morriswood and Schmidt, 2015). In TbSmee1-depleted cells, BSA was observed in one or more small foci next to the flagellar pocket. This was surprising, as BSA is known to traffic in the fluid phase and should not be binding to the cell surface. BSA was however also observed to associate with the flagellar pocket membrane in an electron microscopy-based study of cargo uptake (Gadelha et al., 2009). In a classic radiochemical study of cargo uptake in *T. brucei,* it was observed that "at low concentration, a small adsorbtive component may become prevailing" in assays using BSA (Coppens et al., 1987).

As the experiments conducted here and in the 2009 paper were using BSA well below physiological concentrations, it seems likely that these foci represent the small adsorbtive component that was indirectly observed in the biochemical assays. Therefore, in these assays, the ConA and BSA probes both report on the behaviour of surface-bound cargo, while the dextran reports on fluid-phase uptake.

As ConA is not a physiological cargo, the entry of anti-VSG antibodies into the flagellar pocket was also investigated. These assays turned out to be unexpectedly difficult, owing to much of the signal being lost during the wash steps. This may be related to the recently-reported high rate of VSG shedding (Garrison et al., 2021). Nonetheless, the t=0 and t=2 timepoints would appear to match expectations: in control cells, the anti-VSG antibodies are taken up by endocytosis, while in TbSmee1-depleted cells any remaining signal was found outside the area of the enlarged flagellar pocket and apparently still on the cell surface.

When the TbMORN1 phenotype was characterised, this was the first time that a protein had been shown to play a role in the entry of surface-bound cargo into the flagellar pocket. Subsequent characterisation of Bhalin and the work here on TbSmee1 shows that this cargo entry defect appears to be a consistent effect after enlargement of the flagellar pocket. That the same cargo entry defect is seen upon depletion of clathrin suggests that this is a previously-unobserved feature of all flagellar pocket enlargement ("BigEye") phenotypes. Therefore, active endocytosis appears to be required for the entry of surface-bound cargo into the flagellar pocket.

It should be noted that the results obtained in the clathrin RNAi experiments contradict previously-published observations, where ConA was claimed to accumulate inside the enlarged flagellar pocket of clathrin-depleted cells (Allen et al., 2003). Re-examination of previously-published data from clathrin RNAi cells suggests however that the effect has always been present, with ConA signal (and indeed other cargoes) absent from the flagellar pocket interior and predominantly concentrated in one or more foci just outside it (see for example Fig. S4 in (Allen et al., 2003), and Fig. S2 in (Zoltner et al., 2015)). This oversight is understandable, given that the focus in these clathrin papers was on whether or not cargo endocytosis was occurring, and not whether the cargo was able to access the flagellar pocket (Mark Field, personal communication). Importantly, it also means that similar observations have been made by (at least) two independent groups.

This raises two important questions. First, why does an inhibition of endocytosis apparently hinder the entry of surface-bound cargo into the flagellar pocket? Second, why does the

depletion of hook complex proteins - which are spatially removed from the actual sites of clathrin-coated vesicle formation - cause an endocytosis defect? The prevailing model for several years has been that hydrodynamic flow, driven by flagellar motility, is responsible for sorting surface-bound cargo to the posterior end of the cell where the flagellar pocket is located (Engstler et al., 2007). Once there, the surface-bound material enters the flagellar pocket through a narrow channel where the flagellar membrane and the flagellar pocket neck membrane are not as closely apposed (Gadelha et al., 2009). The results here suggest that the posterior sorting mechanism still functions when endocytosis is inhibited, but that the transit of surface-bound material through the channel and into the flagellar pocket might be impeded.

In this context, it is worth considering the somewhat confusing anti-VSG uptake results obtained at the t=1 timepoint in control cells, where the anti-VSG signal came from a single spot adjacent to the kinetoplast. As a temperature block was in place to prevent endocytosis, this would at first sight suggest that the anti-VSG antibodies can enter the flagellar pocket in the absence of endocytosis, and that the entire hypothesis proposed here is invalid. Given however that the flagellar pocket is less than 1 µm across, it is possible that there is insufficient resolution to distinguish the entrance to the flagellar pocket from the flagellar pocket is expanded due to the BigEye phenotype that these two localisations can be resolved at normal widefield resolution. Higher-resolution imaging of these cells is therefore an important target for follow-up work.

An earlier hypothesis for hook complex function was that it was somehow maintaining the integrity of the channel in the flagellar pocket neck. Nevertheless, the fact that the same phenotype is obtained following depletion of clathrin suggests that the hook complex is instead indirectly affecting endocytosis. This could potentially be by affecting either the localisation or activity of a number of lipid kinases, at least two of which are known to localise to the hook complex (Dean et al., 2017; Demmel et al., 2014). The activity of these enzymes could then be licensing the flagellar pocket membrane for endocytosis, for instance by generating the essential endocytic cofactor phosphatidylinositol-(4,5)-bisphosphate. Subsequent internalisation of membrane by endocytosis would then assist to pull in more membrane currently at the flagellar pocket entrance. How this would be integrated with the activity of the exocytic pathway is, however, unclear.

Thus, while hydrodynamic flow may be responsible for concentrating cargo at the entrance to the flagellar pocket, endocytic activity seems required for the entry of the cargo into the flagellar pocket itself, and endocytosis may assist or be responsible for pulling material in through the channel.

Materials and Methods

Recombinant protein expression and purification

The TbSmee1(1-400) open reading frame was amplified from Trypanosoma brucei brucei strain Lister 427 genomic DNA by PCR. The PCR product was ligated into the p3NH expression vector, which encodes an N-terminal His6 tag, using sequence and ligationindependent cloning (Li and Elledge, 2012). The plasmid was used to transform E. coli strain Rosetta II (DE3)pLysS by heat shock, and individual colonies were subsequently grown at 37°C in the presence of 100 µg/ml kanamycin to an OD600 ~ 0.8–1.0. Recombinant protein expression was induced by the addition of 50 µM IPTG, and the cells were then incubated overnight at 20°C with shaking. Cells were harvested by centrifugation (5000x g for 30 min). The pooled pellet from 6 L of cell culture was resuspended in 300 ml of lysis buffer (50 mM Hepes pH 7.0, 500 mM NaCl, 20 mM imidazole, 5% glycerol, 0.5% Triton-X, 1 mM TCEP, 200 mM PMSF and protease inhibitor cocktail). Pellet emulsions were first homogenised by mixing on ice using a T 10 basic Ultra-Turrax dispersing instrument (IKA). Final lysis was then achieved with sonication on ice, using 3 cycles of 3 min at 50% strength. Lysates were clarified by centrifugation (18,000× g, 45 min, 4°C). The lysates were added to a HiTrap Chelating HP 5 ml column (GE Healthcare) equilibrated with buffer A (20 mM Hepes pH 7, 300 mM NaCl, 40 mM imidazole, 2% glycerol, 1 mM TCEP), and eluted with a 100% step gradient of buffer B1 (20 mM Hepes pH 7, 300 mM NaCl, 400 mM imidazole, 2% glycerol, 1 mM TCEP). Selected peak fractions were examined by SDS-PAGE for protein content and purity. Fractions containing a dominant band at approximately 46 kDa were pooled and concentrated using Amicon Ultra centrifugal filter units with 10K pore size (MerckMillipore) according to the manufacturer's instructions. The His6 tag was removed by 3C protease during overnight dialysis in dialysis buffer (20 mM Tris-HCl pH 7, 300 mM NaCl, 2% glycerol, 1 mM DTT). Significant losses were incurred during this step. The TbSmee1(1-400) was applied to a previously equilibrated HiTrap Chelating HP 5 ml column charged with 50 mM CoCl₂ and coupled to a GSTrap HP 1 ml column (both GE Healthcare). Buffer A was used for equilibration. TbSmee1(1-1400) was mostly collected from the flow-through and a few initial collected fractions. Selected peak fractions were examined by 15% SDS-PAGE for protein content and purity. Fractions containing a dominant band at approximately 44 kDa were pooled and concentrated in Amicon Ultra centrifugal filter units (10K pore size) according to the manufacturer's instructions. Finally, TbSmee1(1-400) concentrates were applied to a previously equilibrated HiLoad 16/600 Superdex 200 pg column (GE

Healthcare) pre-equilibrated in dialysis buffer. Flow speed was adjusted to 1 ml/min and fractions of 1.5 ml were collected. Fractions corresponding to the targeted chromatographic peak were examined for protein content by 15% SDS-PAGE, pooled accordingly to their purity, concentrated, and stored at -80°C.

Antibody generation and affinity purification

Purified recombinant TbSmee1(1-400) was used for the generation of two polyclonal rabbit antisera (Eurogentec). Antisera (303, 304) were initially affinity-purified against the TbSmee1(1-400) antigen, but the neat antisera were later found to show high specificity and were also occasionally used. A third polyclonal antibody (508) was generated against two TbSmee1 peptides (Eurogentec) and affinity purified using the peptide antigens immobilised on a Sulfolink affinity column (ThermoFisher). Results shown were predominantly obtained using affinity-purified "303" and "304" anti-TbSmee1 antibodies; most immunoblotting data were generated using the "304" affinity-purified antibodies, as these had the lowest background in this application, while labelling was identical for "303" and "304" in immunofluorescence experiments. The results obtained with all three antibodies were consistent. The anti-Starkey1 anti-peptide rabbit polyclonal antibodies were generated and affinity purified in the same way.

Antibodies

The following primary antibodies have been described previously: rabbit anti-TbMORN1 (Morriswood et al., 2013), rabbit anti-TbBILBO1 (Esson et al., 2012), mouse anti-Ty1 ("BB2") (Bastin et al., 1996), mouse anti-TbLRRP1 (Zhou et al., 2010), mouse anti-TbCentrin4 ("6C5" (Ikeda and de Graffenried, 2012)), mouse anti-TbCentrin2 ("2B2H1") (de Graffenried et al., 2013), mouse anti-TbFAZ1 ("L3B2") (Kohl et al., 1999), mouse anti-PFR1,2 ("L13D6") (Kohl et al., 1999), mouse anti-FAZ filament ("DOT1")(Woods et al., 1989), rabbit anti-VSG(221) (Batram et al., 2014). The following antibodies came from commercial sources: goat anti-rabbit(IRDye800CW) (LI-COR), goat anti-mouse(IRDye680LT) (LI-COR), goat anti-rabbit and anti-mouse antibodies conjugated to AlexaFluor dyes (Molecular Probes).

Cell culture

Wildtype Lister 427 (monomorphic) BSF cells were cultured in HMI-9 medium (Hirumi and Hirumi, 1989) supplemented with 10% foetal bovine serum (FBS), 100 U/mI penicillin and 0.1 mg/ml streptomycin at 37°C and 5% CO_2 . The SM (single marker) cells (Wirtz et al., 1999) were cultured in the presence of G418 (2.5 µg/ml). Population density was monitored using a Z2 Coulter Counter (Beckman Coulter), and kept below $2x10^6$ cells/ml.

Generation of transgenic cell lines

The Ty1-TbSmee1 endogenous replacement cell line was generated by transfection of 427 cells with a targeting fragment containing 285 bp of the TbSmee1 5' untranslated region (UTR), a blasticidin resistance gene, the intergenic region of alpha- and beta-tubulin, the sequence for a triple Ty1-tag and the first 399 bp of the TbSmee1 open reading frame (ORF) without the start codon. Clones were selected by growth in medium containing 5 µg/ml blasticidin. RNAi target sequences were chosen using RNAit (Redmond et al., 2003). TbSmee1 RNAi cells were generated by cloning the RNAi target sequence into the pGL2084 plasmid (Jones et al., 2014) and then transfecting 2T1 cells with the linearised plasmid (Alsford et al., 2005). Clones were selected by growth in medium containing phleomycin (2.5 µg/ml) and hygromycin (5 µg/ml). TbStarkey1 RNAi cells and TbCHC RNAi cells were generated by cloning the relevant RNAi target sequence into the p2T7_TAblue plasmid (Alibu et al., 2005) and then transfecting SM cells with the linearised plasmid. Clones were selected by growth in medium containing G418 (2.5 µg/ml) and hygromycin (5 µg/ml). Ty1-TbSmee1 truncation constructs were cloned into the pLew100_v5-Hyg plasmid using in vivo assembly (Watson and Garcia-Nafria, 2019). SM cells were transfected with the linearised plasmid. Clones were selected by growth in medium containing G418 (2.5 µg/ml) and hydromycin (5 µg/ml). For transfection >2.5 \times 10⁷ cells of the parental strain were washed and resuspended in 100 µl transfection buffer (90 mM Na₂PO₄, 5 mM KCl, 0.15 mM CaCl₂, 50 mM HEPES [4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid]-NaOH, pH 7.3) containing 10 µg DNA and transfected by electroporation using an AMAXA Nucleofector® Device (Lonza) with program "X-001 free choice". Transfected cells were incubated in 50 ml HMI-9 medium without selection overnight. The next day, drug selection was applied and clones were selected by limiting dilution. At least three separate clones of all cell lines were isolated to control for biological variability. Integration of targeting fragments at endogenous loci or the presence of Ty1-TbSmee1 truncation constructs in the genome were confirmed by PCR analysis of genomic DNA. Genomic DNA was isolated using a DNeasy Bloody & Tissue kit (QIAGEN) and relevant products were amplified by PCR. All cloned constructs used for cell line generation had their DNA sequence confirmed by sequencing.

Immunoblotting

For generation of dephosphorylated whole-cell lysates, cells were harvested by centrifugation (1000xg, 10 min) and the cell pellet was resuspended in 1 ml vPBS (PBS, 46 mM sucrose, 10 mM glucose) containing EDTA-free protease inhibitors (Roche). The washed cells were pelleted by centrifugation (750xg, 4 min). The cells were then resuspended in lysis buffer (0.5% IGEPAL, 0.1M PIPES-NaOH pH 6.9, 2 mM EGTA, 1 mM MgCl2, 0.1 mM EDTA, EDTA-free protease inhibitor cocktail) to a final concentration of

4x10⁵ cells/µl and incubated for 15 min at RT on an orbital mixer to allow dephosphorylation to occur. SDS-loading buffer was then added to a final concentration of 2x10⁵ cells/µl, the samples were further denatured by boiling (100°C, 10 min), and then stored at -20°C. SDS-PAGE was carried out using a Mini-Protean Tetra Cell (Bio-Rad), and protein transfer to nitrocellulose membranes using a Mini-Trans blot cell (Bio-Rad). Protein transfer and equal loading was confirmed using REVERT total protein stain (LI-COR) according to the manufacturer's instructions. Membranes were blocked using blocking buffer (PBS, 0.3% Tween 20, 10% milk)(30 min, RT, rocker). The membranes were then incubated in primary antibodies diluted in blocking buffer (1 h, RT, roller). After three washes in immunoblot buffer (PBS, 0.3% Tween 20) the membranes were visualised using an Odyssey CLx (LI-COR). Background subtraction, qualitative analysis, normalisation relative to total protein stain, and quantification were carried out using Image Studio Lite 5.2 and Empiria Studio 1.1 (LI-COR).

In vitro phosphatase assays

427 BSF cells were grown to approximately 1.5x10⁶ cells/ml and harvested by centrifugation (1,000xg, 10 min, 4 °C). The cells were washed in 1 ml ice-cold wash buffer (0.1M PIPES-NaOH pH 6.9, 2 mM EGTA, 1 mM MgCl2, 0.1 mM EDTA, phosphatase Inhibitor Cocktail 2) and pelleted by centrifugation (750xg, 3 min, 4 °C). For extraction of the cytoskeletons, the cell pellet was resuspended in 1 ml ice-cold extraction buffer (0.5% IGEPAL, 0.1M PIPES-NaOH pH 6.9, 2 mM EGTA, 1 mM MgCl2, 0.1 mM EDTA, EDTA-free protease inhibitor cocktail, phosphatase Inhibitor Cocktail 2) and incubated for 15 min on ice. The cell suspension was inverted every 5 min. The cytoskeletons were separated from the cytoplasm by centrifugation (750xg, 3 min, 4 °C). The cytoskeletons were washed with 0.5 ml ice-cold 1x NEBuffer for PMP supplemented with 1 mM MnCl₂ and pelleted by centrifugation (750xg, 3 min, 4 °C). After resuspension in 375 µl ice-cold 1x NEBuffer for PMP supplemented with 1 mM MnCl₂, all following samples were taken from this stock. An input sample (0 min) of 40 µl was taken and added to 20 µl of SDS-loading buffer. Two control samples were taken, 0.4 µl Phosphatase Inhibitor Cocktail 2 was added, and incubated on ice and at 26°C, respectively, for 20 min. 20 µl of SDS-loading buffer was then added. To assay for dephosphorylation, 130 µl were taken from the stock and 1 µl lambda phosphatase (400 U) was added. This sample and the remainder from the stock were incubated at 26 °C. After 1/2/3/5/10/20 min 20 µl samples were taken from each and were added to 10 µl SDS-loading buffer. All the samples were boiled at 104 °C for 10 min and stored at -20 °C. Samples were analysed by immunoblotting.

Fractionation

50 ml 427 BSF cells were grown to approximately $1.5x10^6$ cells/ml and then harvested by centrifugation (1000xg, 10 minutes, 4°C). The cell pellet was resuspended in 1 ml vPBS, transferred to a microfuge tube, and the cells again pelleted by centrifugation (750xg, 4 min, 4°C). The supernatant was discarded and the centrifugation step was repeated to remove remaining supernatant. The cell pellet was resuspended in 200 µl extraction buffer (0.5% IGEPAL, 0.1M PIPES-NaOH pH 6.9, 2 mM EGTA, 1 mM MgCl2, 0.1 mM EDTA, EDTA-free protease inhibitor cocktail) and incubated for 15 minutes at RT in an orbital mixer. A 5% input sample was taken (10 µl), put into a new microfuge tube and left on ice. The extracted cells were fractionated by centrifugation (3400xg, 2 minutes, 4°C) and the supernatant transferred to a fresh microfuge tube and the exact volume noted. The tube containing the extracted cells was centrifuged again at the same settings and this second residual supernatant discarded. The cytoskeleton pellet was then resuspended in 200 µl extraction buffer. 5% samples of supernatant and pellet fractions were taken and analysed by immunoblotting. Equal fractions of I, SN, P were loaded in each lane (I ~ $1.4x10^6$ cells).

Preparation of samples for immunofluorescence microscopy

Coverslips were washed in 70% ethanol and then incubated with 0.01% poly-L-lysine in a 24-well plate (>20 min, RT) and left to dry. 2x10⁶ cells were taken per coverslip and transferred to 15 ml Falcon tubes. The cells were pelleted by centrifugation (1000xg, 1 min per ml of liquid, RT) in a swing-bucket centrifuge. The supernatant was removed, and the cell pellet was gently resuspended in 1 ml ice-cold vPBS + Complete. The cells were again pelleted by centrifugation (1000xg, 2 min, RT) and subsequently resuspended in 1 ml icecold vPBS + Complete and directly added to the coverslips. The cells were attached to the coverslips by centrifugation (1000xg, 1 min, RT), and attachment was confirmed visually. The attached cells were then incubated in 1 ml ice-cold extraction buffer (0.5% IGEPAL 0.1M PIPES-NaOH pH 6.9, 2 mM EGTA, 1 mM MgCl2, 0.1 mM EDTA, EDTA-free protease inhibitor cocktail) (5 min, on ice). The extracted cells were washed two times with 1 ml vPBS + Complete and then fixed in 1 ml ice-cold 99.9% methanol (30 min, -20°C). The fixed cells were rehydrated using 1 ml PBS. The coverslips were blocked in 1 ml 3% BSA in PBS (30 min, RT), and sequentially incubated with clarified primary and secondary antibodies diluted in PBS (1 h, RT, humidified chamber for each) with three PBS washing steps (3 x 5 min, RT, rocker) after each incubation. After the final wash, glass slides were cleaned with 70% ethanol and a spot of DAPI-Fluoromount G (Southern Biotech) was placed on the surface. The coverslips were rinsed in ddH₂O, carefully dried, and then mounted. For preparation of samples for SIM imaging, different coverslips (high precision, No. 1.5 H) were used. Cells were harvested as above, washed, and fixed in 4% paraformaldehyde solution (10 min, ice

then 30 min, RT). The fixed cells were washed, attached to coverslips, permeabilised (0.25% Triton X-100 in PBS; 5 min, RT), and then labelled and mounted as described above.

Fluorescence microscopy

Images were acquired using a DMI6000B widefield microscope (Leica Microsystems, Germany) with a HCX PL APO CS objective (100x, NA = 1.4, Leica Microsystems, Germany) and Type F Immersion Oil (refractive index = 1.518, Leica Microsystems, Germany). The microscope was controlled using LAS-X software (Leica). Samples were illuminated with an EL6000 light source (Leica) containing a mercury short-arc reflector lamp (HXP-R120W/45C VIS, OSRAM, Germany). Excitation light was selected by using Y3 (545/25 nm), GFP (470/40 nm), and A4 (360/40 nm) bandpass filter cubes (Leica Microsystems, Germany). The power density, measured at the objective focal plane with a thermal sensor head (S175C, Thorlabs), was respectively 0.749 ± 0.086, 0.557 ± 0.069, 0.278 ± 0.076 W/cm² for the three filters. Emitted light was collected at ranges of 605/70 (Y3), 525/50 nm (GFP), and 470/40 nm (DAPI) respectively. The individual exposure times and camera gains were adjusted according to the different samples. RNAi samples (control and depleted) were imaged using identical settings. Differential interference contrast (DIC) was used to visualise cell morphology. 3D recording of each field of view was obtained using 40 Z-slices (step size = $0.21 \,\mu$ m). Fields of view were selected in the DIC channel in order to blind the user to the fluorescence signal and subjectively select for cells with optimum morphology. Images were captured using a DFC365 FX monochrome CCD camera (Leica, 6.45 µm pixel size). SIM images were acquired using an Elyra S.1 SIM microscope (Zeiss) and ZEN software (Zeiss).

Fluorescence microscopy image processing and analysis

Processing was carried out using FIJI (Schindelin et al., 2012) and a custom macro for the generation of maximum-intensity z-projections with single DIC z-slices overlaid. Overlaps between two proteins were confirmed in individual z-slices (thickness: 210 nm). The plugin ScientiFig was utilised to create the collage and adding the inserts (Aigouy and Mirouse, 2013). For correlation analysis between TbSmee1 and other flagellar pocket collar and/or hook complex associated proteins, 1K1N cells were selected using both DIC and DAPI channels. 2D sum slices projections were prepared for each stack of both green and red channels. The projections were clipped to 8-bit depth and a convoluted background subtraction was applied. All resultant individual 2D images, without channel overlay, were analysed pairwise to check for intensity-based correlation based on Zhang & Cordelières (Zhang and Cordelières, 2016). The Spearman's rank correlation results were further computed into Microsoft Excel sheets and analysed using R version 4.1.2 (R Core Team,

2022) in the environment RStudio 2021.09.1.372 (RStudio Team, 2020). The packages used for all descriptive analysis and plot generation were ggplot2 (Wickham, 2016), openxlsx (Schauberger and Walker, 2022), and psych (Revelle, 2023). All Fiji/ImageJ macros and R scripts for correlation analysis were written by Alyssa Borges and are available upon request.

Growth curves

RNAi cells were seeded at the required starting concentration in a volume of 22 ml and divided into two 10 ml aliquots in separate flasks. Tetracycline was added to a final concentration of 1 μ g/ml in one flask to induce RNAi, and refreshed every 24 h. The population density of the control and induced cells was measured every 24 h over a time course of 72h, or every hour over a time course of 8 h, using a Z2 Coulter Counter (Beckman Coulter). Depletion of the target protein was confirmed in every experiment by immunoblotting of whole-cell lysates.

Cell division cycle analysis

An aliquot of 10⁶ cells was taken, and the cells were fixed directly in media by addition of isothermal 25% glutaraldehyde to a final concentration of 2.5% (10 min, 37°C, gentle mixing). The cells were then pelleted by centrifugation (750xg, 10 min). The cell pellet was resuspended in 0.5 ml 2.5% glutaraldehyde in PBS, transferred to a microfuge tube, and incubated at RT (30 min, gentle mixing). The cells were pelleted again by centrifugation (750xg, 4 min), and the cell pellet resuspended in 500 µl PBS. The cells were then added to the coverslips inside the 24-well plate and attached by centrifugation (1000xg, 1 min, RT). The coverslips were mounted on poly-L-lysine-coated slides using DAPI-Fluoromount G (Southern Biotech). Imaging was as described for fluorescence microscopy above, using DAPI and DIC channels only. Cell division cycle stages (1K1N, 2K1N, 2K2N) were manually quantified from maximum intensity projections of the DAPI signal overlaid with single DIC z-slices. Depletion of the target protein at each timepoint was confirmed by immunoblotting of whole-cell lysates from the same experiments.

Preparation and imaging of electron microscopy samples

Induced and uninduced RNAi cells were grown for 24 h to a density of $1-2x10^6$ cells/ml in 50 ml medium and harvested by centrifugation (1000xg, 10 min, RT). The supernatant was removed to 4 ml and 4 ml FBS was added. The cells were pelleted again (1000xg, 10 min, RT) and the supernatant was removed to 200 µl. The cells were resuspended in the supernatant, and the suspension was then transferred to a PCR tube and pelleted by centrifugation (1,600xg, 10s, RT). The cells were then transferred into a carrier with a closed

lid to avoid air inclusions. High pressure freezing (HPF) was started immediately (Leica EM HPM100). After HPF the samples were transferred to an AFS (Leica EM AFS2) for freeze substitution and progressive lowering of temperature. Low temperature embedding and of raisin (DDSA, MNA, polymerisation Epon Epon812, 2,4,6 Tris(dimethylaminomethyl(phenol))) were then carried out. Ultra-thin cuts (60 nm) were carried out with an ultramicrotome (Leica EM UC7/FC7) and were placed on slotted grids. For contrasting they were incubated in 2% uranyl acetate for 8 min. Afterwards the grids were washed 3x in ddH₂O (boiled to remove CO₂) and incubated for 5 min on 50% Reynold's lead citrate in a petri dish with NaOH tablets. The grids were again washed 2x in ddH₂O. A 200 kV transmission electron microscope (Jeol, JEM-2100) with a TemCam F416 4k x 4k camera (Tietz Video and Imaging Processing Systems) and EMMenu 4.0.9.31 software (Tietz) were used. Uninduced control cells were viewed at a magnification of 12,000x, induced cells were viewed at a magnification of 8,000x.

Measurement of flagellar pocket enlargement

Induced and uninduced RNAi cells at a concentration of ~5×10⁶ cells/ml were harvested by centrifugation (1000xg, 4°C). The cells were resuspended in 45 µl ice-cold vPBS + protease inhibitors and incubated on ice to block endocytosis (10 min, ice). 5 µl labelled dextran (10 kDa, 50 mg/ml stock) was added and mixed by flicking. The mixture was then incubated to allow dextran to enter and fill the flagellar pocket (15 min, on ice, dark). At the end of the incubation, 1 ml ice-cold vPBS + protease inhibitors was added and the cells pelleted by centrifugation (750 xg, 2 min, 4°C). The cell pellet was resuspended in 0.5 ml vPBS + protease inhibitors, and the cells fixed by addition of 0.5 ml 8% pfa and 0.1% glutaraldehyde solution in vPBS + protease inhibitors (20 min, on ice then 60 min, RT). The fixed cells were pelleted by centrifugation (750 xg, 2 min) and washed twice with 1 ml vPBS + protease inhibitors. The dextran signal was immediately measured by flow cytometry using a FACSCalibur (BD Biosciences) running CellQuest ProTM Software (BD Biosciences). Later processing and analysis was carried out using FlowJo 10.8.1. Dot plots were created using Plots of Data (Postma and Goedhart, 2019). After flow cytometry, the remaining cells were attached to coverslips by centrifugation, mounted on clean glass slides using DAPI-Fluoromount G (Southern biotech), and imaged using fluorescence microscopy on the same day to confirm flagellar pocket labelling. Depletion of TbSmee1 was confirmed in every experiment by immunoblotting of whole-cell lysates.

Cargo (ConA, BSA) uptake assays

2x10⁶ cells per sample were harvested by centrifugation (1000 xg, 10 min, 4°C) and washed in 1 ml ice-cold vPBS. The washed cells were pelleted by centrifugation (750 xg, 2 min, 4°C),

and resuspended in 100 µl ice-cold vPBS. The cells were then incubated at low temperature (20 min, ice) to block endocytosis. During this incubation, the labelled cargoes were prepared. The 50 mg/ml dextran aliquot (10 kDa AF488 conjugate, Molecular Probes) was mixed using a sonicator bath (10 min, 37 Hz) and then vortexed. The 0.5 mg/ml ConA aliquot (TMR conjugate, Molecular Probes) was clarified by centrifugation (11,000 xg, 10 min, 4°C). Both cargoes were kept on ice and in the dark. At the end of the incubation, 10 µl of the 50 mg/ml dextran and 2 µl of the 0.5 mg/ml ConA were added to the chilled cells, mixed by flicking, and incubated at low temperature (15 min, on ice, in the dark). T=0 samples were quenched and fixed at this point; t=30 samples had an additional incubation to allow endocytosis of cargo (30 min, 37°C, in the dark) before guenching and fixing. To quench samples, 1 ml ice-cold vPBS was added, and the cells were pelleted by centrifugation (750 xg, 2 min, 4°C). The pelleted cells were resuspended in 50 µl vPBS by flicking, and then fixed by addition of 0.5 ml ice-cold fix solution (4% paraformaldehyde solution, 0.1% glutaraldehyde in vPBS) (20 min, on ice, in the dark, then 30 min, RT, in the dark). The fixed cells were pelleted by centrifugation (750 xg, 2 min, RT), washed in 2 ml vPBS, resuspended in 1 ml vPBS, and then attached to poly-L-lysine-coated coverslips by centrifugation. Coverslips were mounted on glass slides using DAPI-Fluoromount G (Southern Biotech) and imaged immediately. Assays using BSA (AlexaFluor555 conjugate, Molecular Probes) were done in the same way. The 20 mg/ml BSA aliquot was clarified by centrifugation (750 xg, 1 min, RT)), and 3 µl was added to the cells/dextran mixture. Depletion of TbSmee1 was confirmed by immunoblotting of whole-cell lysates in every experiment.

Anti-VSG uptake assay

2x10⁶ cells per sample were harvested by centrifugation (1000 xg, 1 min per ml of liquid + 1, 4 °C) and washed in 1 ml ice-cold vPBS. The washed cells were pelleted by centrifugation (1000 xg, 3 min, 4 °C), resuspended in 50 μ l ice-cold vPBS and incubated at low temperature to block endocytosis (20 min, ice). During incubation, the primary and secondary antibodies were clarified by centrifugation (10,000 xg, 10 min, 4 °C). The primary antibody (anti-VSG antiserum) was diluted 1:50 in ice-cold vPBS. The secondary antibody was diluted to required concentration (1:3000) in 3% BSA in PBS. Both were kept on ice in the dark until used. At the end of the incubation, 50 μ l of diluted anti-VSG was added to the cells and quickly mixed by flicking, followed by an incubation to allow binding (15 min, ice). To fix cells for t=0, 0.5 ml ice-cold vPBS and 0.5 ml ice-cold 8% pfa in vPBS were sequentially added at the end of the incubation and kept on ice. For timepoints t=1 & t=2, 1 ml ice-cold vPBS was added. The cells were washed by centrifugation (1000 xg, 2 min, 4 °C) and resuspended in 1 ml ice-cold vPBS. To fix t=1, 350 μ l of ice-cold 16% pfa was

added and the cells were kept on ice. The cells for t=2 were incubated (2 min, 37°C) to allow redistribution, flagellar pocket entry, and endocytosis of the anti-VSG. 350 μ l of ice-cold 16% pfa was then added to fix the cells. All three timepoints were incubated on ice (20 min) after addition of pfa, followed by a subsequent incubation at RT (30 min). The fixed cells were pelleted by centrifugation (1000 xg, 2 min, RT), resuspended in 1 ml RT vPBS, and attached to poly-L-lysine-coated coverslips by centrifugation (1000 xg, 1 min, RT). The attached cells were then permeabilised in 1 ml 0.25% Tx100 in PBS (5 min, RT). The permeabilised cells were washed two times in 1 ml PBS and the coverslips with cells were placed on a drop of clarified and diluted secondary antibody (70 μ l) in a humidified chamber and incubated (1h, RT, dark). The coverslips were washed 3 times in 1 ml PBS (5 min, RT, rocker, dark). After rinsing the cells in ddH₂O, they were mounted onto a drop of DAPI-Fluoromount G (Southern Biotech) on a coverslip that was cleaned with 70% EtOH.

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Competing interests

No competing interests declared.

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Data availability

All relevant data can be found within the article and its supplementary information. Raw data files are available upon request.

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Figures

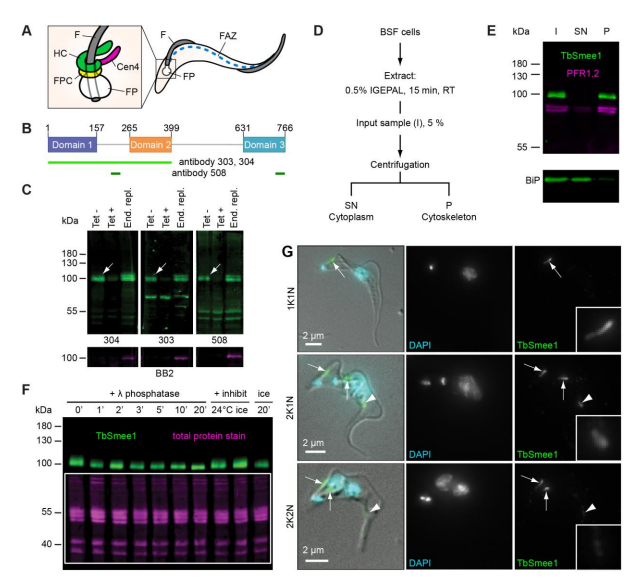


Fig. 1. TbSmee1 is a cytoskeleton-associated phosphoprotein. (A) Schematic representation of a trypanosome cell with flagellar pocket (FP), flagellum (F), and flagellum attachment zone (FAZ) indicated. Inset shows cytoskeleton-associated structures in the flagellar pocket neck area: hook complex (HC); flagellar pocket collar (FPC); centrin arm (Cen4). (B) Schematic representation of TbSmee1 (766 amino acids), predicted domains shown in purple, orange, and teal. The regions used for the generation of the three anti-TbSmee1 antibodies (303, 304, 508) are shown in light/dark green bars below the schematic. (C) Validation of the specificity of the three anti-TbSmee1 antibodies. Immunoblots against whole cell lysates from uninduced TbSmee1 RNAi cells (Tet -), induced TbSmee1 RNAi cells (Tet +), and Ty1-TbSmee1 endogenous replacement cells (End. repl.) using the three anti-TbSmee1 antibodies (304, 303, 508). All three antibodies detected an ~ 85 kDa protein in the Tet- samples (upper panels, white arrows). Exemplary results from

multiple (n>3) independent experiments are shown. (D) Fractionation scheme. Bloodstream form (BSF) cells were extracted with the non-ionic detergent IGEPAL and separated into cytoplasm and cytoskeleton fractions by centrifugation. (E) TbSmee1 is associated with the cytoskeleton. Immunoblot of whole-cell input (I), cytoplasmic supernatant (SN) and cytoskeletal pellet (P) fractions with anti-TbSmee1 antibodies. Equal fractions were loaded in each lane. Exemplary results from multiple (n>3) experiments are shown. (F) TbSmee1 is phosphorylated in vivo. Trypanosome cell lysates were incubated with λ phosphatase for the indicated times (0-20 min), followed by TbSmee1 detection by immunoblotting. Equal loading of samples was confirmed using total protein stain (magenta). Exemplary results from multiple (n>3) independent experiments are shown. (G) TbSmee1 localisation through the cell cycle. Bloodstream form trypanosomes were extracted using non-ionic detergent, fixed, and labelled with anti-TbSmee1 antibodies. DNA was stained using DAPI. In 1K1N cells, TbSmee1 localised to a bar-shaped structure (arrow) close to the kinetoplast. In 2K1N cells the structure had replicated (arrows). TbSmee1 was additionally present on a third structure farther along the cell body (arrowhead). Exemplary images from multiple (n>3) independent experiments are shown.

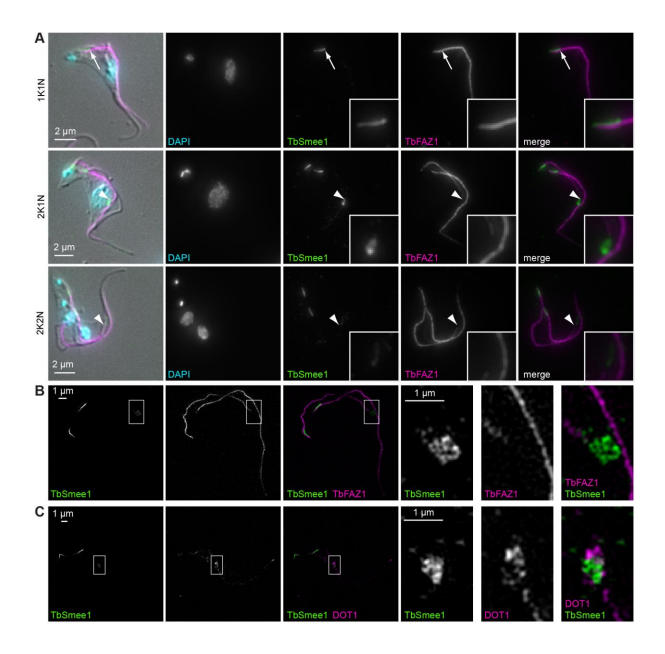


Fig. 2. TbSmee1 is associated with the tip of the elongating new FAZ in replicating cells. (A) Bloodstream form trypanosomes were extracted with non-ionic detergent, fixed, and labelled with the indicated antibodies. DNA was stained with DAPI. Maximum intensity projections are shown, with a single DIC z-slice overlaid. In 1K1N cells, TbSmee1 partially overlapped (arrow) with the posterior end of the FAZ. In 2K1N and 2K2N cells, the additional TbSmee1 structure (arrowhead) was present at the tip of the elongating new FAZ. Exemplary images from multiple (n>3) independent experiments are shown. (B) SIM image of the same preparations. The TbSmee1 structure (inset) lay just ahead of the elongating FAZ tip. (C) SIM image of a detergent-extracted cell colabelled for TbSmee1 and DOT1.

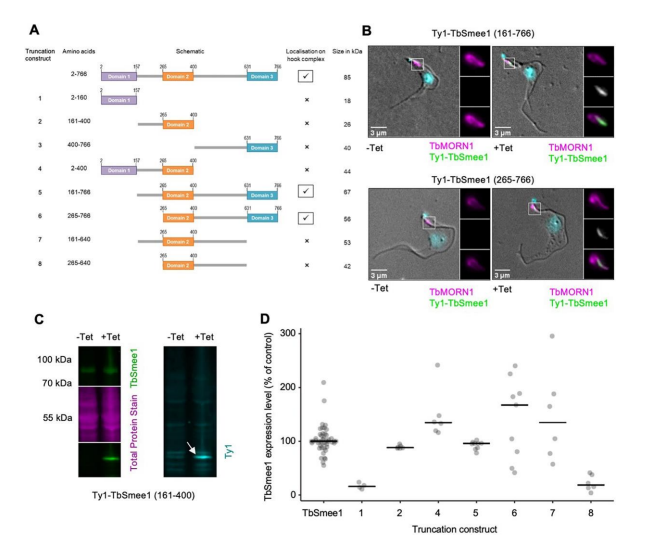


Fig. 3. TbSmee1 domains 2 and 3 are required for targeting to the hook complex. (A) Schematics of the 8 TbSmee1 truncation constructs tested. (B) TbSmee1 domains 2 and 3 are required for localisation to the hook complex. Stably-transfected cells inducibly expressing the indicated TbSmee1 truncations were used. Ty1-TbSmee1 was detected using anti-Ty1 antibodies. In the absence of induction (-Tet), no signal was seen. After induction of expression (+Tet), the Ty1-TbSmee1 truncations overlapped with the shank part of TbMORN1 (insets). Images shown are maximum intensity projections of the fluorescence channels overlaid with a single DIC z-slice. Exemplary images are shown. (C) Confirmation of Ty1-TbSmee1 truncations (+Tet) were immunoblotted using anti-TbSmee1 (left panel) and anti-Ty1 (right panel) antibodies. Total protein stain (magenta) was used as a loading control. In the exemplary blot shown, the Ty1-TbSmee1 was detected at ~ 44 kDa, as expected. (D) Quantification of immunoblotting data. Anti-TbSmee1 signals in the immunoblots were normalised relative to total protein staining. TbSmee1 levels in uninduced

control cells (TbSmee1) were expressed relative to the mean value of all control samples for each clone in each experiment. Ty1-TbSmee1 levels were expressed relative to the levels of endogenous TbSmee1 for each clone in each experiment. The data shown were obtained from two independent experiments with each Ty1-TbSmee1 truncation; each experiment used three separate clones.

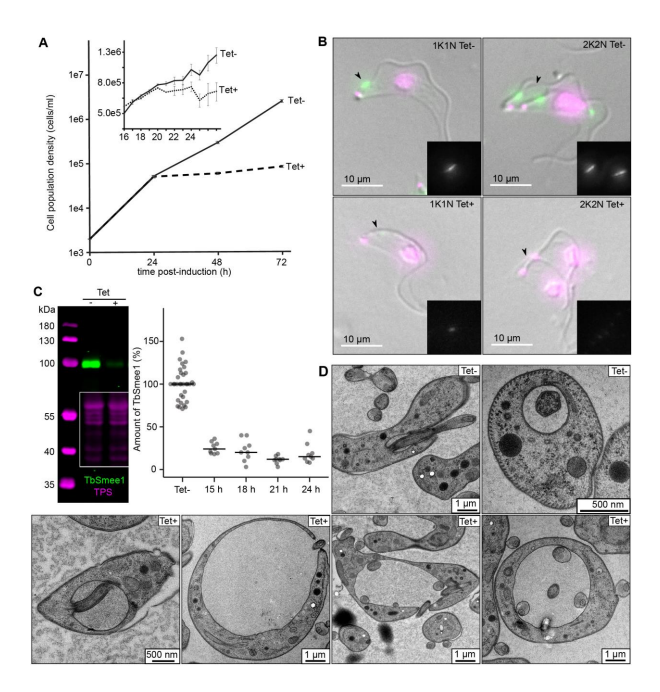


Fig. 4. TbSmee1 is essential for the viability of bloodstream form *T. brucei.* (A) Depletion of TbSmee1 causes a strong growth defect. The population density of uninduced control (Tet-) and TbSmee1-depleted (Tet+) cells were measured after induction of RNAi over a 72 h timecourse. Inset shows data from experiments with readings taken at hourly intervals. (B) Confirmation of TbSmee1 depletion at the single-cell level. Uninduced control (Tet-) and TbSmee1-depleted (Tet+) RNAi cells were extracted, fixed, and labelled with anti-TbSmee1 antibodies (green). DNA was labelled with DAPI (magenta). Maximum intensity projections of the fluorescence channels are shown overlaid with a single DIC z-slice. Insets show the TbSmee1 signal. Exemplary 1K1N and 2K2N cells from each condition are shown. (C) Confirmation and quantification of TbSmee1 depletion. Whole-cell lysates from control

(Tet-) and TbSmee1-depleted (Tet+) RNAi cells were immunoblotted using antibodies against TbSmee1 (green). Total protein stain (TPS, magenta) was used for signal normalisation, using the same approach as described in Figure 3D. An exemplary immunoblot is shown. TbSmee1 depletion was quantified at the indicated timepoints post-induction. (D) Depletion of TbSmee1 results in an enlargement of the flagellar pocket. Electron microscopy images of control (Tet-) and TbSmee1-depleted (Tet+) RNAi cells are shown. The cells were fixed 24 h post-induction. All data were obtained from multiple ($n \ge 3$) independent experiments, each using three separate clones.

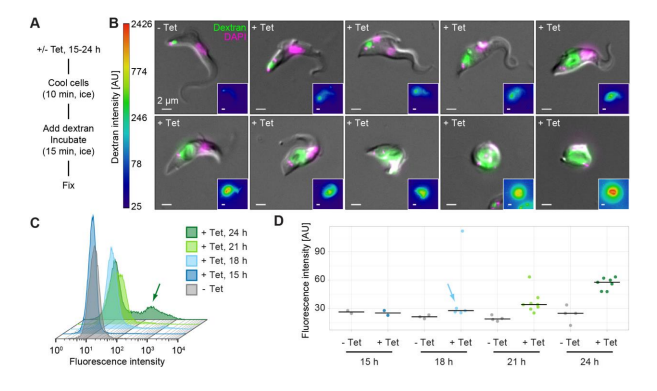


Fig. 5. Flagellar pocket enlargement is an early consequence of TbSmee1 depletion. (A) Summary of dextran uptake protocol. (B) The intensity of the dextran signal reports on flagellar pocket volume. Cells were incubated with fluorophore-conjugated dextran (green) prior to being fixed and imaged using fluorescence microscopy. DNA was labelled with DAPI (magenta). Exemplary cells from control (- Tet) and TbSmee1-depleted (+ Tet) conditions are shown. The + Tet cells (18, 21, or 24 h of induction) exhibited various degrees of flagellar pocket enlargement and progressive morphological aberration. Insets show the dextran signal with a log-scale LUT. (C) Flow cytometry analysis of control (- Tet) and TbSmee1-depleted (+ Tet) cells incubated with fluorescent dextran at various timepoints after induction of RNAi. At later timepoints there is a clear emergence of a subpopulation of cells with much greater fluorescence intensity (arrow). Exemplary traces from a single experiment are shown. (D) Quantification of flow cytometry data. The geometric mean of the fluorescence intensity in control (- Tet) and TbSmee1-depleted (+ Tet) cells was measured at various timepoints after induction of RNAi. Bars indicate median values; a visible shift was visible as early as 18 h post-induction (arrow). All data were obtained from multiple (n≥2) independent experiments; each experiment used three separate clones.

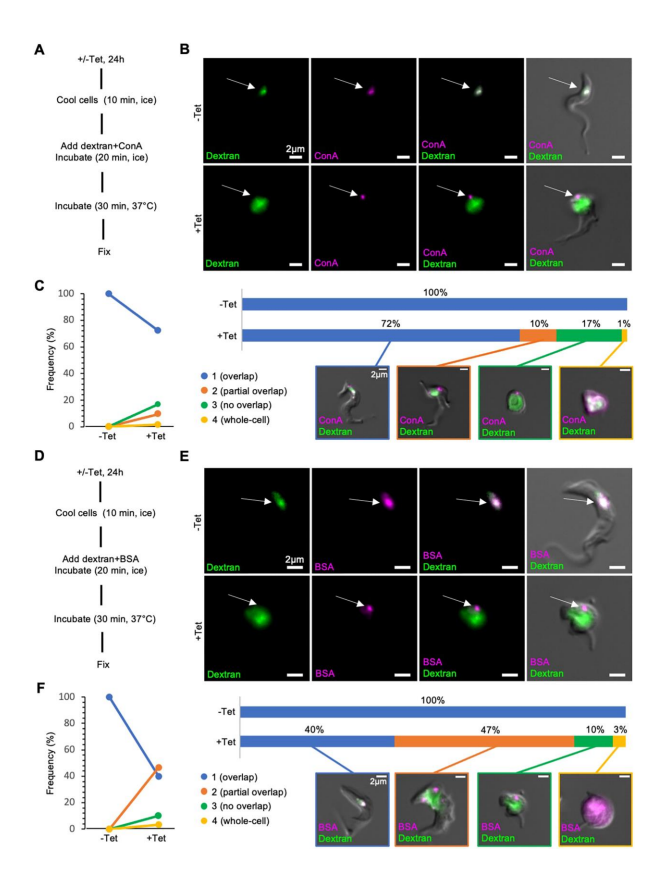


Fig. 6. Surface-bound reporters cannot enter the flagellar pocket of TbSmee1-depleted cells. (A) Summary of paired dextran+ConA uptake protocol. (B) ConA is unable to enter the enlarged flagellar pocket of TbSmee1-depleted cells. In control (-Tet) cells, the dextran and

ConA signals strongly overlapped (arrow, upper panels). In TbSmee1-depleted cells (+Tet), little to no overlap between the two reporters was observed (arrow, lower panels). Maximum intensity projections of the fluorescence channels are shown overlaid with a single DIC z-slice. (C) Quantification of dextran+ConA uptake experiments. Cells from control (-Tet) and TbSmee1-depleted (+Tet) samples were manually classified into four categories (1-4) based on the degree of overlap between the ConA and dextran reporters. Results are shown as a slope chart (left) and stacked bar chart (right); exemplary cells for each category are shown. Percentages represent total cell counts (442 cells) that were obtained from multiple independent experiments (n>3); each experiment included three separate clones. (D) Summary of paired dextran+BSA uptake protocol. (E) BSA is unable to enter the enlarged flagellar pocket of TbSmee1-depleted cells. (F) Quantification of dextran+BSA uptake experiments. Percentages represent total cell counts (103 cells) that were obtained from multiple independent experiments (n>3); each experiment included three separate clones.

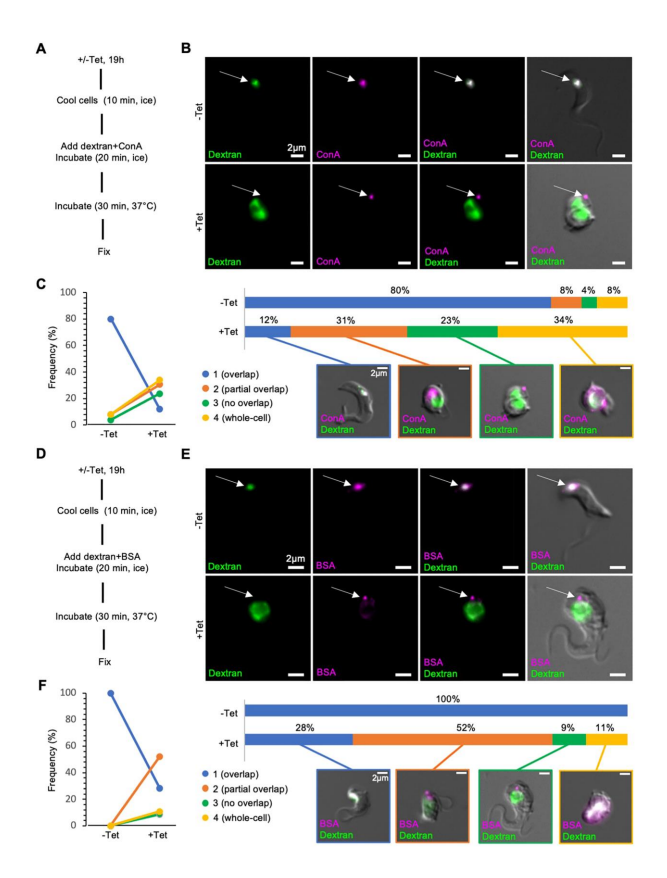


Fig. 7. Blockade of surface reporters is due to an inhibition of endocytosis. (A) Summary of paired dextran+ConA uptake protocol. (B) ConA is unable to enter the enlarged flagellar pocket of clathrin-depleted cells. Maximum intensity projections of the fluorescence channels are shown overlaid with a single DIC z-slice. (C) Quantification of dextran+ConA

uptake experiments, using the same approach as in Figure 6. Percentages represent total cell counts (84 cells) that were obtained from multiple independent experiments (n>3); each experiment included three separate clones. (D) Summary of paired dextran+BSA uptake protocol. (E) BSA is unable to enter the enlarged flagellar pocket of clathrin-depleted cells. (F) Quantification of dextran+BSA uptake experiments. Percentages represent total cell counts (58 cells) that were obtained from multiple independent experiments (n>3); each experiment included three separate clones.

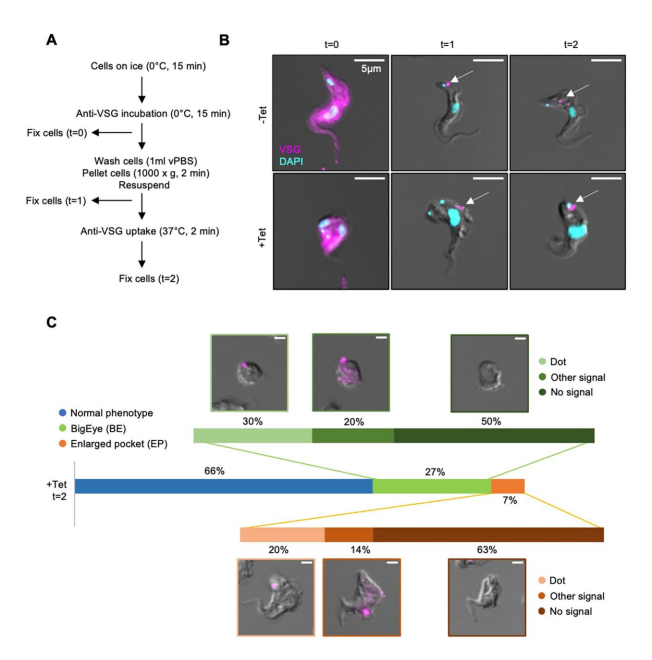


Fig. 8. Anti-VSG does not enter the flagellar pocket of TbSmee1-depleted cells. (A) Summary of anti-VSG uptake protocol. (B) Anti-VSG is unable to enter the enlarged flagellar pocket of TbSmee1-depleted cells. In control (-Tet) cells, anti-VSG signals were observed at the cell surface (t=0), then at the location of the flagellar pocket (t=1) and finally at the endosomal/lysosomal system after allowing endocytosis (t=2) (arrows, upper panels). In TbSmee1-depleted cells (+Tet), no anti-VSG signals were observed in the enlarged flagellar pocket or in the endosomal/lysosomal system (t=1, t=2) (arrows, lower panels). Maximum intensity projections of the fluorescence channels are shown overlaid with a single DIC *z*-slice. (C) Quantification of anti-VSG uptake experiments. Cells from TbSmee1-depleted samples (+Tet, t=2) were manually classified into three categories (Normal phenotype, BigEye, and Enlarged pocket). The anti-VSG signal could not be observed in enlarged

flagellar pockets in BigEye cells or cells with slightly enlarged flagellar pockets. Signals in these cells were classified into three categories (Dot, Other signal or No signal). Results are shown as a stacked bar chart; exemplary cells for each category are shown. Percentages represent total cell counts (408 cells) that were obtained from multiple independent experiments (n>3); each experiment included three separate clones.

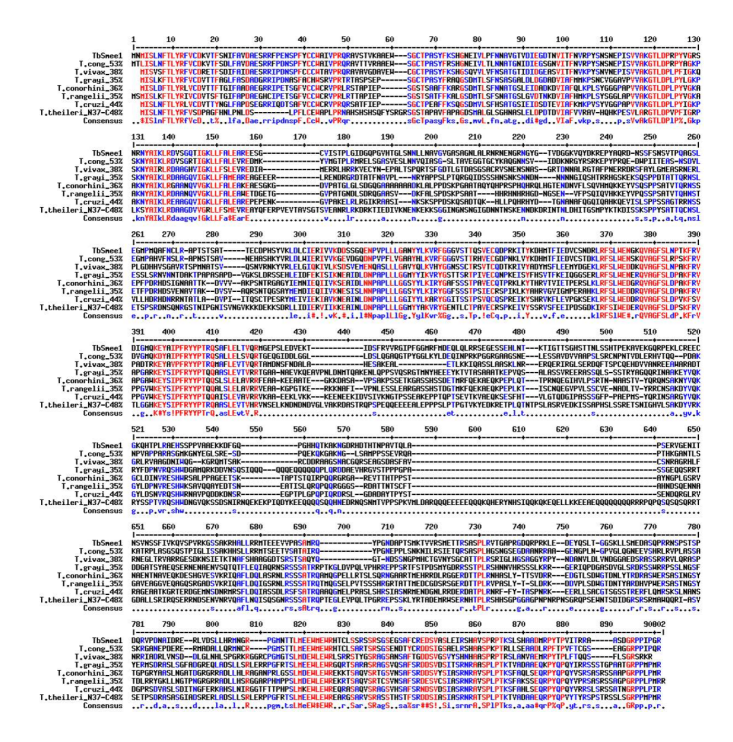


Fig. S1. The TbSmee1 primary structure contains three blocks of highly-conserved sequence. Multiple sequence alignment of Trypanosoma brucei Smee1 (TbSmee1) and homologous proteins from other trypanosome species, generated using MultAlin (Corpet, 1988) with default parameters. Amino acid numbers are indicated in black numerals above the alignment. Moderately (50-90%) conserved residues are highlighted in blue. Highly-conserved (>90%) or completely conserved residues are highlighted in red. The consensus sequence is shown below the alignment. Abbreviations: TbSmee1, Trypanosoma brucei Smee1; T.cong, Trypanosoma congolense; T.vivax, Trypanosoma vivax; T.grayi, Trypanosoma grayi. T.conorhini, Trypanosoma conorhini; T.rangelii, Trypanosoma rangelii; T.cruzi, Trypanosoma cruzi; T.theileri, Trypanosoma theileri. The % sequence identity of each homologue to TbSmee1 is indicated after the name.

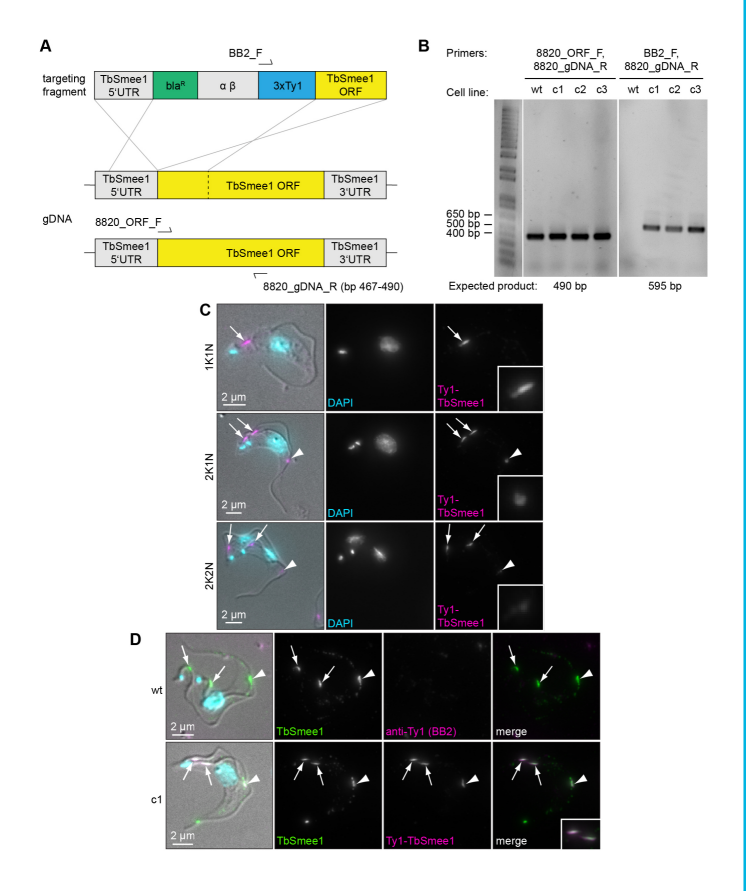


Fig. S2. Generation of Ty1-Tb5mee1 endogenous replacement cell line. (A) Schematic showing the in situ tagging procedure and annealing sites for PCR primers. The two endogenous alleles of TbSmee1 in the gDNA are shown in yellow, flanked by 5'UTR and 3'UTR sequences. T. brucei cells were transfected with a targeting fragment with homology arms for recombination with the 3' end of the 5'UTR and 5' end of the ORF. The targeting fragment contained a blasticidin resistance gene (blaR), the intergenic region from the alpha/beta tubulin locus (a) and a 3xTy1 epitope tag preceded by an ATG start codon. Homologous recombination removed the endogenous ATG start codon of the TbSmee1 ORF. (B) Confirmation of targeting fragment integration at the endogenous TbSmee1 locus by PCR analysis of genomic DNA. Genomic DNA from wild-type (wt) and candidate Ty1-TbSmee1 clones (c1, c2, c3) was analysed by PCR. Left panel: positive control using 8820 0RF F and 8820 gDNA R primers; 490 bp product expected. Right panel: integration test using BB2 F and 8820 gDNA R primers. A product is only expected if the 3xTy1 sequence has integrated upstream of the TbSmee1 ORF (see primer annealing sites in panel A). Two independent experiments were carried out, each using all three separate clones. (C) Ty1-TbSmee1 displays the same localisations through the cell cycle as endogenous TbSmee1. Detergent-extracted cells were fixed with methanol and labelled with anti-Ty1 antibodies; DNA was stained using DAPI. Exemplary cells from the three main cell cycle states (1K1N, 2K 1N, 2K2N) are shown. Maximum-intensity z-projections of the fluorescence channels are shown, together with a single DIC z-slice overlay. The Ty1-TbSmee1 signal is shown in magenta in the overlay and highlighted with arrows. Arrowheads indicate the Ty1-TbSmee1 present at the tip of the new FAZ. Insets show an enlarged view of the TbSmee1 signal from the hook complex or FAZ tip. Multiple (n>3) independent experiments were carried out using three separate clones. (D) The anti-Ty1 signal is specific for Ty1-TbSmee1. Wild-type (wt) and Ty1-TbSmee1 cells were extracted with detergent, fixed with methanol, and labelled with anti-TbSmee1 and anti-Ty1 antibodies. Hook complex (arrows) and FAZ tip (arrowheads) localisations are indicated. No anti-Ty1 signal was seen in wild-type cells; strong overlap between the anti-Ty1 and anti-TbSmee1 signals was seen in the Ty1-TbSmee1 cells. Maximum-intensity zprojections of the fluorescence channels are shown, together with a single DIC z-slice overlay.

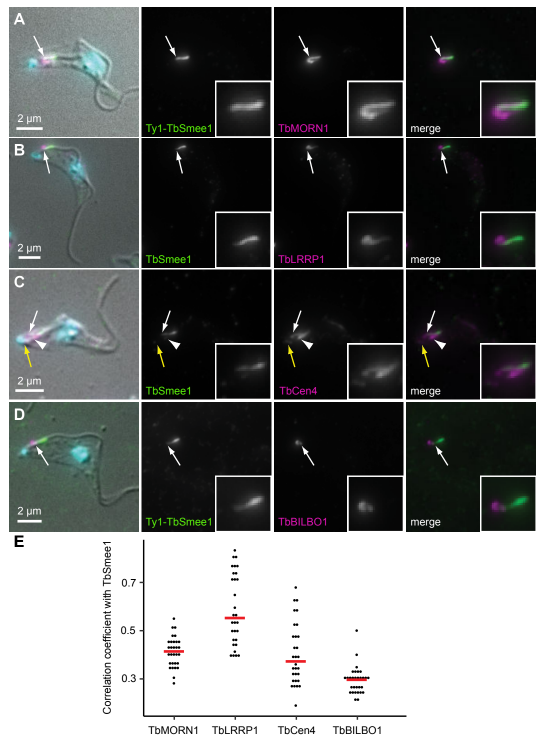


Fig. S3. TbSmee1 is localised to the shank part of the hook complex. Bloodstream form trypanosomes were extracted with non-ionic detergent, fixed, and labelled with the indicated antibodies. Either wild-type or Ty1-TbSmee1 cells were used. Insets show an enlarged view of the hook complex region. (A) TbSmee1 overlaps with the shank part of the hook complex protein TbMORN1 (arrow). (B) TbSmee1 overlaps with the shank part of the hook complex protein TbLRRP1 (arrow). (C) TbSmee1 partially overlaps with TbCen4. TbCen4 is present at the basal and probasal bodies (yellow arrow), centrin arm (arrowhead), and a small additional projection (white arrow). (D) TbSmee1 does not overlap with the flagellar pocket collar protein TbBILBO1 (arrow). (E) Summary of measured correlation coefficients for each of the colabelling experiments; red bars show median values. TbSmee11 showed a moderate correlation with TbMORN1 (0.41) and TbLRRP1 (0.55) and a weak correlation with TbCen4 (0.37) TbBILBO1 (0.3). Each dot represents a single cell in the 1K1N stage (N = 30). All fluores-cence images are maximum intensity z-projections, and an overlay with a single DIC section is shown. Overlap was manually confirmed in single z-slices. Results were obtained from multiple (n>3) independent experiments; exemplary images are shown.

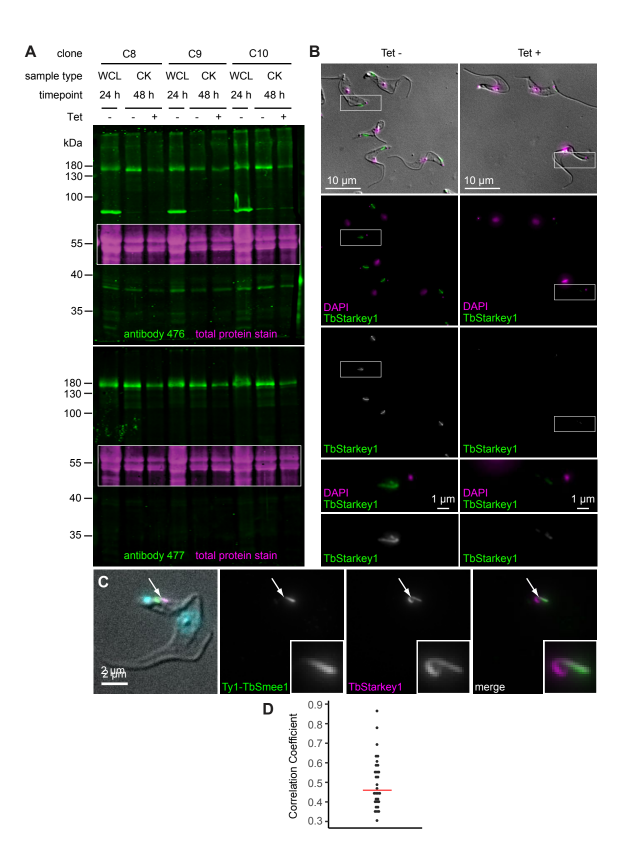


Fig. S4. Validation of anti-TbStarkey1 antibodies. (A) Validation of anti-TbStarkey1 antibody specificity by immunoblotting. Three separate TbStarkey1 RNAi clones (C8, C9, C10) were analysed by immunoblotting. Both whole-cell lysates (WCL) and detergentextracted cytoskeleton (CK) samples were obtained at the indicated timepoints (24 h, 48 h) from control (Tet-) and induced (Tet+) cells. The samples were immunoblotted using two separate anti-TbStarkey1 affinity-purified antibodies (antibody 476, 477). Both antibodies recognised a >130 kDa protein whose abundance was depleted after 48 h of RNAi. The 476 antibody additionally showed a significant cross-reaction with a protein of <100 kDa in WCL but not CK samples. A section of the total protein stain of each membrane is shown as an inset (magenta). (B) Validation of anti-TbStarkey1 antibody specificity by immunofluorescence microscopy. Control (Tet-) and TbStarkey1-depleted (Tet+) RNAi cells were extracted with non-ionic detergent, fixed with methanol, and labelled with anti-TbStarkey1 antibodies (green); DNA was stained using DAPI (magenta). TbStarkey1 localised to the hook complex, and signal was lost upon depletion. The boxed areas are shown enlarged in the bottom panels. Identical results were obtained using both anti-TbStarkey1 antibodies; exemplary images using the 477 antibodies are shown. (C) TbSmee1 overlaps with the shank part of the hook complex protein TbStarkey1 (arrow). All fluorescence images are maximum intensity projections, and an overlay with a single DIC section is shown. Overlap was manually confirmed in single z-slices. Results were obtained from multiple (n>3) independent experiments; exemplary images are shown. (D) Summary of measured correlation coefficients. Each dot represents a single cell; red lines show median values.

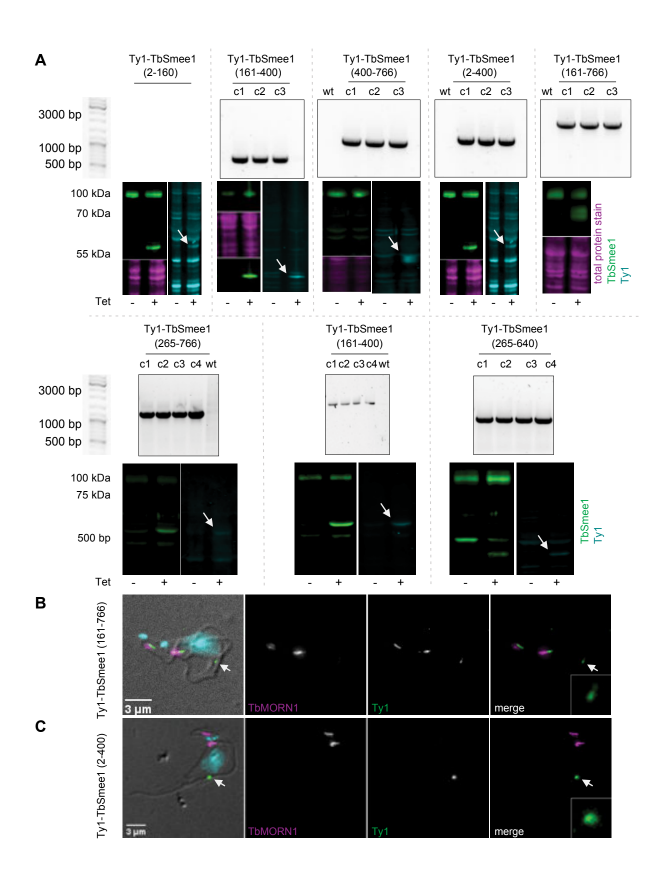


Fig. S5. Validation of TbSmee1 truncations. (A) Validation of Ty1-TbSmee1 truncation cell lines by PCR analysis of genomic DNA and immunoblotting. The results for all 8 TbSmee1 truncations are shown, separated by dotted lines. Upper panels: PCR analysis of genomic DNA to confirm the presence of the truncation construct. PCR was used to amplify DNA from clones (c1, c2, c3) and wild-type control (wt) genomic DNA. Primers annealing to the sequence encoding Ty1 epitope and within the truncation were used. Lower panels: confirmation of Ty1-TbSmee1 truncation construct expression by immunoblotting. Wholecell lysates from uninduced (-Tet) and induced (+Tet) cells were analysed by immunoblotting with anti-TbSmee1 (green) and anti-Ty1 (cyan) antibodies. Both antibodies detected proteins corresponding to the predicted size of the Ty1-TbSmee1 truncations; arrows indicate the target protein in the anti-Ty1 blots. A portion of the total protein staining of the membranes is shown in magenta. (B) Ty1-TbSmee1(161-766) localises to both the hook complex and the FAZ tip. Cells expressing the Ty1-TbSmee1(161-766) construct were extracted with non-ionic detergent, fixed with methanol, and labelled with anti-TbMORN1 and anti-Ty1 antibodies. DNA was stained with DAPI (cyan). Ty1-TbSmee1(161-766) was observed at both the hook complex and the FAZ tip (arrow). (C) Ty1-TbSmee1(2-400) localises to the FAZ tip but not the hook complex. Cells expressing the Ty1-TbSmee1(2-400) construct were extracted, fixed, and labelled as above. Ty1-TbSmee1(2-400) was observed exclusively at the FAZ tip (arrow). Images in panels B and C are maximum intensity z-projections with a single DIC z-slice overlay. Multiple (n>2)independent experiments using three separate clones for each construct were carried out; exemplary cells are shown.

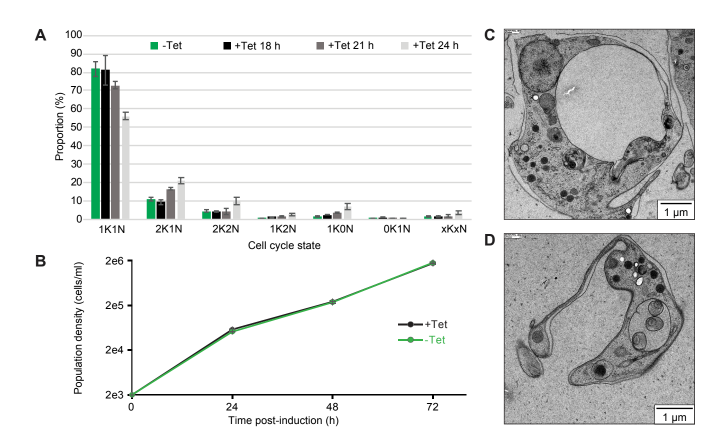


Fig. S6. Additional results for TbSmee1 and TbStarkey1 RNAi experiments. (A) TbSmee1 depletion causes changes to cell cycle state distribution. Control (-Tet) and TbSmee1-depleted (+Tet) RNAi cells from 18 h, 21 h, 24 h post-induction were fixed using glutaraldehyde; DNA was stained using DAPI. The various cell cycle states (1K1N, etc) were manually quantified from images taken of the fixed cells. Data were obtained from three independent experiments, each using three separate clones; at least 240 cells were quantified for each timepoint. (B) Depletion of TbStarkey1 has no effect on population cell growth. Control (-Tet) and TbStarkey1-depleted (+Tet) cells were followed over a 72 h timecourse, and population density (cells/ml) was measured every 24 h. Data were obtained from three independent experiments, each using 3 separate clones. (C, D) Depletion of TbStarkey1 causes morphological abnormality. TbStarkey1-depleted cells were prepared for electron microscopy using high-pressure freezing and imaged. Cells with enlarged flagellar pockets could readily be observed.

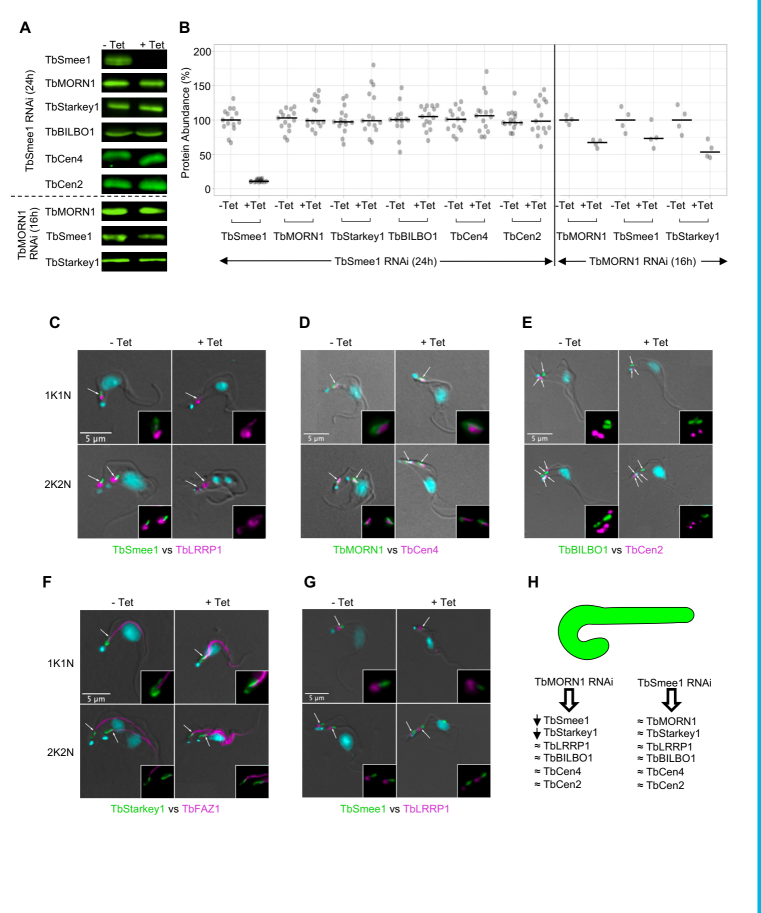
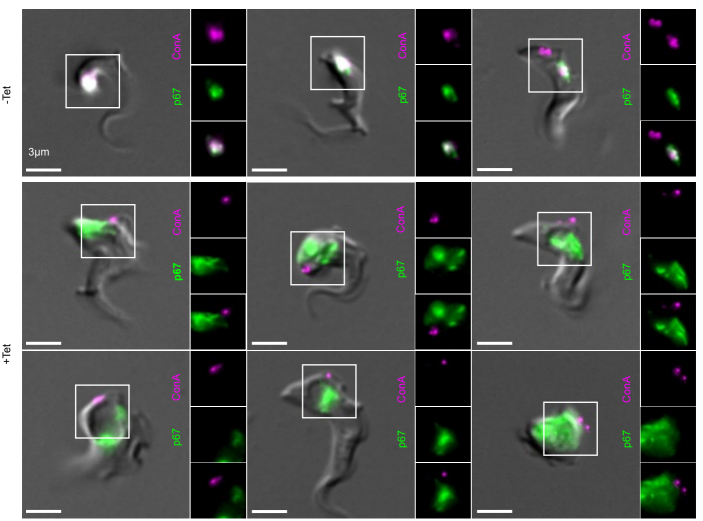


Fig. S7. Depletion of TbSmee1 affects other cytoskeleton-associated proteins. TbSmee1 RNAi cells were induced for 24 h and analysed by immunoblotting of whole-cell lysates and immunofluorescence imaging of detergent-extracted cells. (A) Exemplary immunoblots showing the effect of 24 h TbSmee1 depletion on marker proteins for the hook complex, flagellar pocket collar, and centrin arm. For comparison, the effect of 16 h TbMORN1 depletion was also analysed. (B) Quantification of immunoblot data. TbSmee1 depletion did not affect any of the proteins analysed. Total protein staining was used for signal normalisation. Normalised TbSmee1 signals in +Tet cells were expressed relative to the -Tet signal for each sample. TbSmee1 -Tet signals were expressed relative to the mean of all TbSmee1 -Tet values in the dataset. Immunoblots from TbMORN1 RNAi experiments were quantified in the same way; TbMORN1 depletion resulted in a loss of both TbSmee1 and TbStarkey1 signal. The data shown were obtained from multiple (n>2) independent experiments, each using three (TbSmee1 RNAi) or two (TbMORN1 RNAi) separate clones. (C-F) TbSmee1 depletion does not affect the localisation of the marker proteins. Control (-Tet) and TbSmee1-depleted (+Tet) cells were extracted with detergent, fixed with methanol, and labelled with the indicated antibodies; DNA was stained using DAPI. The position of the hook complexes in exemplary 1K1N and 2K2N cells are shown with arrows; panel E also indicates basal bodies. Data were obtained from 2 independent experiments for each labelling combination, each using 3 separate clones. (G) TbSmee1 and TbLRRP1 in TbMORN1-depleted cells.(H) Summary of the observed effects (arrow = depletion, squiggles = no change) on the abundance of marker proteins caused by either TbMORN1 or TbSmee1 depletion, based on immunoblotting and immunofluorescence data.



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Fig. S8. Depletion of TbSmee1 prevents trafficking of ConA to the lysosome. Control (-Tet) and TbSmee1-depleted cells (+Tet; 24 h timepoint) were incubated on ice with ConA (magenta). The cells were then shifted to 37 °C for 30 min to allow internalisation. The cells were then fixed and labelled with antibodies specific for the lysosome marker p67 (green). Control (-Tet) cells showed strong overlap between the two labels, indicating that ConA had been trafficked to the lysosome. +Tet cells showed no overlap between the two labels. Maximum intensity projections of the fluorescence channels are shown overlaid with a single DIC z-slice. Overlap in -Tet cells was confirmed in single z-slices. Single channels from the boxed area in each image are shown as insets. Data obtained from multiple (n>2) independent experiments each using three separate clones; exemplary cells are shown.