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Polo-like kinase phosphorylation of the orphan kinesin KIN-G negatively regulates centrion arm biogenesis in *Trypanosoma brucei*

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eLife Assessment

This **important** study provides new insight into the regulation of cell organization and division in *Trypanosoma brucei* through the control of a kinesin motor protein by a polo-like kinase. The authors present **solid** evidence from rigorous biochemical and imaging analyses showing that phosphorylation modulates kinesin function and cellular organization. However, direct in vivo evidence that PLK phosphorylates kinesin-G is lacking.

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Abstract

The unicellular parasite *Trypanosoma brucei* assembles a motile flagellum that is required for locomotion, cell division plane placement, and cell-cell communication. Inheritance of the flagellum during the cell cycle relies on the faithful duplication/segregation of multiple flagellum-associated cytoskeletal structures, including a centrion-marked, bar-shaped structure termed centrion arm, which also determines the biogenesis site for Golgi. Biogenesis of the centrion arm requires the Polo-like kinase homolog TbPLK and the orphan kinesin KIN-G, but the mechanistic role of TbPLK in centrion arm biogenesis remains elusive. Here we report that TbPLK phosphorylates KIN-G, disrupts its microtubule-binding activity, and negatively regulates its function in the procyclic form of *T. brucei*. TbPLK phosphorylates KIN-G *in vitro* at multiple residues, some of which are phosphorylated *in vivo* in *T. brucei*, including the Thr301 residue within one of the microtubule-binding motifs of the kinesin motor domain. Phosphorylation of Thr301 by TbPLK inhibits the microtubule-binding activity of KIN-G *in vitro*, and expression of a Thr301 phospho-mimic mutant in *T. brucei* disrupts centrion arm integrity, thereby impairing Golgi biogenesis, flagellum attachment zone elongation, flagellum positioning, and cell division plane placement. Therefore, TbPLK negatively regulates KIN-G activity by phosphorylating Thr301, and dephosphorylation of Thr301 is required for KIN-G to fulfill its cellular function in promoting centrion arm biogenesis.

Introduction

The early branching parasitic protozoan *Trypanosoma brucei* causes sleeping sickness in humans and nagana in cattle in sub-Saharan Africa, and it possesses a single flagellum, which is required for cell motility, cell division, and cell-cell communication. The flagellum originates from the basal body, which has a typical structure of centrioles in eukaryotes, exits the cell body through the flagellar pocket, and extends toward the anterior tip of the cell by attaching to the cell membrane via the flagellum attachment zone (FAZ), a specialized cytoskeletal structure consisting of the

intra-cellular FAZ filament, the flagellum FAZ domain, and the junction between the flagellar membrane and the cell membrane (Sunter and Gull, 2016). At the proximal end of the flagellum and near the flagellar pocket region, there sit several flagellum-associated cytoskeletal structures, including the flagellar pocket collar (FPC) (Lacomble et al., 2009) and the hook complex (Esson et al., 2012). The horseshoe-shaped FPC, which is marked by TbBILBO1, wraps around the flagellum, whereas the hairpin-shaped hook complex, which consists of a fishhook-like structure marked by TbMORN1 and a bar-shaped structure named centrin arm and marked by TbCentrin2 (Esson et al., 2012; Morriswood, 2015), sits on the top of the FPC (Esson et al., 2012). Between the mature basal body and the pro-basal body, there originates a specialized set of four microtubules termed the microtubule quartet (MtQ), which passes through the hook complex and the FPC and extends, alongside the intracellular FAZ filament, to the anterior tip of the cell (Esson et al., 2012; Vaughan and Gull, 2016). The intracellular FAZ filament originates from a region within the hook complex structure (Esson et al., 2012) and elongates alongside the MtQ toward the anterior cell tip (Sunter et al., 2015; Zhou et al., 2015). In close proximity to the centrin arm, the Golgi apparatus is located (He et al., 2004), and it sits next to the endoplasmic reticulum exit site (ERES), thereby forming unique junctions to facilitate protein trafficking (Bangs, 2011).

During the S phase of the cell cycle in *T. brucei*, a new pair of mature basal body/pro-basal body is assembled, followed by the assembly of a new flagellum from the mature basal body and the subsequent assembly of the flagellum-associated cytoskeletal structures. The newly synthesized flagellum and its associated, newly synthesized cytoskeletal structures are segregated into the new-flagellum daughter (NFD) cell, whereas the old-flagellum daughter (OFD) cell inherits the existing flagellum and its associated cytoskeletal structures (Abeywickrema et al., 2019). Segregation or positioning of the newly assembled flagellum depends on the faithful duplication and segregation of its associated cytoskeletal structures, which further impacts the placement of the cell division plane/cleavage furrow and the faithful division of the daughter cells (Hu et al., 2015; Pham et al., 2020; Zhou et al., 2010; Zhou et al., 2011). During the cell cycle, the Golgi apparatus undergoes *de novo* duplication, assembling a new Golgi next to the old Golgi in a centrin arm-dependent manner (He et al., 2005), and the newly assembled Golgi is segregated into the NFD cell, likely by associating with the newly assembled centrin arm through the scaffold protein CAAP1 (Zhou et al., 2024).

The precise functions of the hook complex remain elusive, but its deficiency often leads to defects in the elongation of the new FAZ filament, the positioning of the new flagellum, the placement of cell division plane, and, consequently, cytokinesis (Pham et al., 2020; Pham et al., 2019; Zhou et al., 2010; Zhou et al., 2025; Zhou et al., 2024). Several regulatory proteins that control hook complex biogenesis/assembly have been identified, including the Polo-like kinase homolog TbPLK (de Graffenried et al., 2008), the kinetoplastid-specific protein phosphatase KPP1 (Zhou et al., 2018b), the E3 ubiquitin ligase complex CRL4^{WDR1} (Hu et al., 2017), the orphan kinesin KIN-G (Zhou et al., 2024), and the KIN-G-interacting protein WDR2 (Zhou et al., 2025). TbPLK phosphorylates the centrin arm protein TbCentrin2 to promote centrin arm duplication (de Graffenried et al., 2013), whereas its protein abundance is regulated by CRL4^{WDR1}, thereby impacting centrin arm biogenesis (Hu et al., 2017). KIN-G is a microtubule plus end-directed motor protein, which localizes to centrin arm and regulates centrin arm biogenesis (Zhou et al., 2024), and WDR2 promotes centrin arm biogenesis by recruiting KIN-G (Zhou et al., 2025). KIN-G is phosphorylated at multiple residues *in vivo* in trypanosome cells (Nett et al., 2009; Zhou et al., 2025), but the protein kinase responsive for KIN-G phosphorylation was not identified previously, and whether phosphorylation of KIN-G may regulate KIN-G function remains unknown.

Here we report that TbPLK phosphorylates KIN-G at multiple sites, including the Thr301 residue in one of the microtubule-binding motifs of the motor domain, and that phosphorylation of Thr301 inhibits the microtubule-binding activity of KIN-G. We further show that phosphorylation of Thr301 disrupts the cellular function of KIN-G by impairing centrin arm biogenesis, which further impacts FAZ elongation, flagellum positioning, cell division plane placement, and Golgi biogenesis. These results suggest that TbPLK negatively regulates KIN-G activity through phosphorylating

Thr301 and that dephosphorylation of Thr301 is required for KIN-G to exert its cellular function in the procyclic form of *T. brucei*. This finding underscores a novel control mechanism of KIN-G by the Polo-like kinase in regulating centrin arm biogenesis in *T. brucei*.

Results

TbPLK co-localizes with KIN-G at centrin arm and phosphorylates KIN-G *in vitro*

The findings that TbPLK and KIN-G are both required for the biogenesis of the centrin arm and its associated Golgi apparatus (de Graffenried et al., 2008; Zhou et al., 2024) and that KIN-G is phosphorylated *in vivo* in trypanosome cells (Nett et al., 2009; Zhou et al., 2025) prompted us to test whether TbPLK may phosphorylate KIN-G and thus may regulate KIN-G function. We first investigated when the two proteins may co-localize during the cell cycle by co-immunofluorescence microscopy analysis of cells expressing endogenously triple HA-tagged KIN-G. During G1 phase, TbPLK and KIN-G co-localized at the centrin arm, with KIN-G more enriched at the distal end of the centrin arm (Fig. 1A). During early S-phase when the centrin arm started to duplicate, KIN-G and TbPLK co-localized, almost entirely, to the elongating centrin arm (Fig. 1A, S-phase cell on the left). When cells gradually progressed to late S-phase, the centrin arm was further elongated and finally separated to two structures, KIN-G localized to the elongating centrin arm (Fig. 1A, S-phase cell in the middle) and the separated centrin arm structures, respectively (Fig. 1A, S-phase cell on the right). However, during these stages, TbPLK was enriched at the new FAZ tip and the flagella connector, with a small amount of TbPLK remaining on the old centrin arm, where it co-localized with KIN-G (Fig. 1A, S-phase cells in the middle and on the right). During G2 phase when the two centrin arms were farther separated, KIN-G remained on the two centrin arm structures, but TbPLK no longer resided on the centrin arm, but was enriched on the new FAZ tip and the flagella connector (Fig. 1A). Note that from G1 phase to G2 phase, TbPLK was additionally localized to the basal body (Fig. 1A).

We asked whether TbPLK may phosphorylate KIN-G, so we performed *in vitro* kinase assay using purified recombinant TbPLK and KIN-G proteins and ATP- γ -S for thio-phosphorylation and subsequent detection of thio-phosphorylated proteins by the anti-PNBM antibody (Allen et al., 2007). We detected auto-phosphorylation of TbPLK, indicating active TbPLK, and phosphorylation of KIN-G (Fig. 1B), suggesting that KIN-G is an *in vitro* substrate for TbPLK. To identify the *in vitro* TbPLK phosphosites on KIN-G, we performed *in vitro* kinase assay with purified recombinant TbPLK and KIN-G proteins and regular ATP, and then separated the proteins by SDS-PAGE, which showed that the phosphorylated KIN-G protein migrated slower than the non-phosphorylated KIN-G protein (Fig. 1C). The phosphorylated KIN-G protein band was excised and analyzed by mass spectrometry, which identified 29 *in vitro* TbPLK phosphosites (Fig. 1D), among which 10 sites have been previously identified as *in vivo* phosphosites (Nett et al., 2009; Zhou et al., 2025). Five out of these 10 phosphosites, Thr32, Thr72, Ser194, and Thr301, are located within the motor domain of KIN-G, with Thr301 located in the microtubule-binding motif #3 (Fig. 1D, E). The remaining five sites, Thr388, Thr523, Ser553, Ser556, and Thr617, are in the C-terminal coiled-coil motifs and in the loop between coiled-coil motifs #2 and #3 (Fig. 1D). The identification of ten *in vitro* TbPLK phosphosites that are also phosphorylated *in vivo* in trypanosome cells suggests that these sites could be potential *in vivo* TbPLK phosphosites.

TbPLK phosphorylation of KIN-G disrupts KIN-G binding to microtubules

Because TbPLK co-localized with KIN-G during early cell cycle stages at the centrin arm and phosphorylates KIN-G in the microtubule-binding motif (Fig. 1), we tested the effect of TbPLK phosphorylation on the microtubule-binding activity and/or microtubule-gliding activity of KIN-G by *in vitro* microtubule-binding and -gliding assay (Zhou et al., 2024). Purified recombinant KIN-G protein was first settled onto glass coverslips in a flow chamber, and then rhodamine-labeled

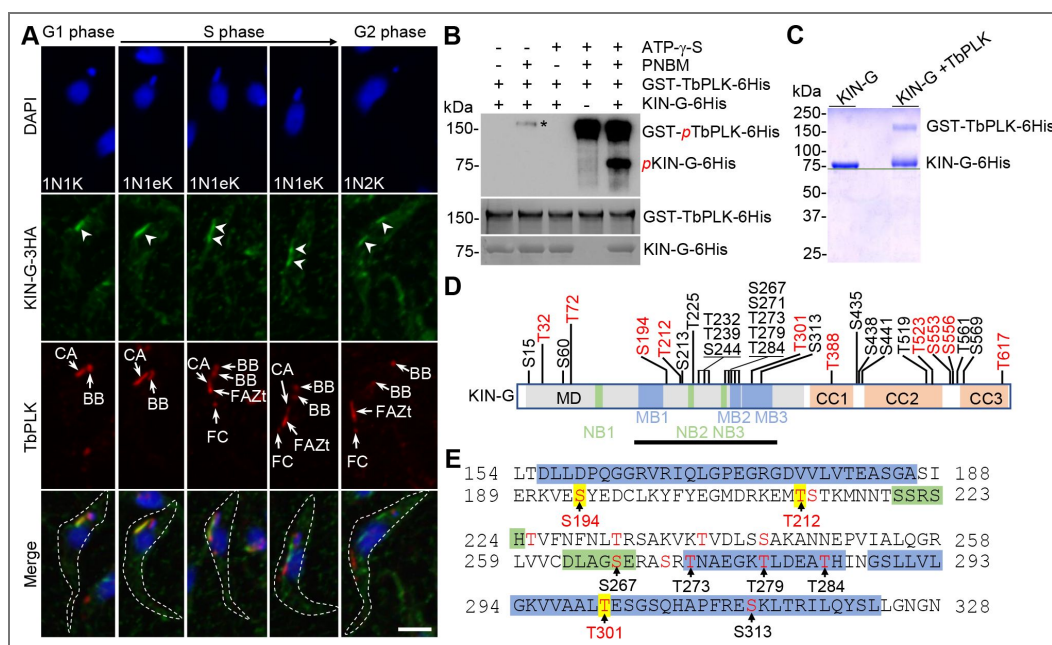


Figure 1. KIN-G is a substrate of TbPLK in *T. brucei*.

(A). Co-immunostaining of KIN-G-3HA and TbPLK during the cell cycle in the procyclic form. Open arrowheads indicate KIN-G signal at the centrin arm. TbPLK signal at different structures are indicated. BB: basal body; CA: centrin arm; FC: flagella connector; FAZt: FAZ tip. Scale bars: 5 μ m. (B). TbPLK phosphorylates KIN-G *in vitro*. The asterisk indicates a non-specific band. (C). *In vitro* phosphorylated KIN-G migrates slower than non-phosphorylated KIN-G on SDS-PAGE. (D). *In vitro* TbPLK phosphosites on KIN-G protein identified by mass spectrometry. Phosphosites highlighted in red indicate the *in vitro* and *in vivo* phosphosites. MD: motor domain; CC: coiled coil; MB: microtubule-binding motif; NB: nucleotide-binding motif. (E). Phosphosites within the KIN-G protein sequence spanning MB1 to MB3. Sequences highlighted in blue indicate the microtubule-binding motifs (MB1, MB2 and MB3), and sequences highlighted in green indicate the nucleotide-binding motifs (NB2 and NB3). Phosphosites highlighted in red in the yellow box indicate the *in vitro* and *in vivo* phosphosites.

microtubules were added into the flow chamber, where microtubules first bound to KIN-G and then moved toward their minus-ends (opposite to the plus end-directed moving direction of KIN-G) (Zhou et al., 2024 [DOI](#)). We detected strong binding of microtubules when only KIN-G was settled on the coverslips or together with the kinase-dead TbPLK mutant TbPLK^{K70R}, but we observed a significant reduction in the number of attached microtubules when TbPLK was co-settled with KIN-G (Fig. 2A, B [DOI](#)), suggesting that TbPLK activity inhibits the microtubule-binding activity of KIN-G. In the presence of TbPLK^{K70R}, the microtubule gliding speed of KIN-G was slightly, but insignificantly, reduced (Fig. 2C [DOI](#)), which suggests that the binding of TbPLK^{K70R} to KIN-G somewhat interfered with its motility on microtubules.

We next performed *in vitro* microtubule-binding and -gliding assay by adding TbPLK after incubating microtubules with settled KIN-G on the glass coverslip (Fig. 2D [DOI](#)). Without adding TbPLK, the number of bound microtubules decreased by an average of ~30% at 12 min, whereas when TbPLK was added into the chamber, the number of bound microtubules decreased by an average of ~83% at 12 min (Fig. 2D, E [DOI](#)), demonstrating that TbPLK-mediated phosphorylation caused the detachment of the microtubules that had attached to KIN-G before the addition of TbPLK. Together, these results suggest that the kinase activity of TbPLK inhibits the binding of KIN-G to microtubules.

Phosphorylation of Thr301 in KIN-G by TbPLK impairs the binding of KIN-G to microtubules

Because TbPLK phosphorylates the Thr301 residue within the microtubule-binding motif #3 of KIN-G (Fig. 1 [DOI](#)) and TbPLK disrupts the microtubule-binding activity of KIN-G *in vitro* (Fig. 2 [DOI](#)), we asked whether phosphorylation of Thr301 may disrupt the microtubule-binding activity *in vitro*. We mutated Thr301 to generate phospho-mimic and phospho-deficient mutants, KIN-G^{T301D} and KIN-G^{T301A}, respectively, purified the recombinant proteins, and used them in the *in vitro* microtubule-binding and -gliding assay. The results showed that KIN-G^{T301A} had similar microtubule-binding activity to wild-type KIN-G, but KIN-G^{T301D} lost microtubule-binding activity (Fig. 3A, B [DOI](#)), demonstrating that phosphorylation of Thr301 by TbPLK disrupted the microtubule-binding activity. Notably, the microtubule-gliding activity of KIN-G^{T301A} was moderately, but insignificantly, reduced (Fig. 3C [DOI](#)), suggesting that mutation of this residue somehow reduced the motility of KIN-G, although the reduction in motility does not affect the cellular function of KIN-G^{T301A} (see below).

We also mutated the Thr284 residue, an *in vitro* TbPLK phosphosite located in the microtubule-binding motif #2 (Fig. 1E [DOI](#)), to generate phospho-mimic and phospho-deficient mutants, KIN-G^{T284D} and KIN-G^{T284A}, respectively, and tested their microtubule-binding and -gliding activity. Neither KIN-G^{T284D} nor KIN-G^{T284A} showed any defects in microtubule-binding and -gliding activity (Fig. 3 [DOI](#)).

Phosphorylation of Thr301 in KIN-G by TbPLK causes defective cell proliferation

We ectopically expressed triple HA-tagged KIN-G^{T301D}, KIN-G^{T301A}, and KIN-G, which were recoded to resist RNAi, in the KIN-G RNAi cell line to investigate the potential effects on the cellular function of KIN-G in the procyclic form of *T. brucei*. Immunofluorescence microscopy showed that like wild-type KIN-G, both KIN-G mutants were localized to the centrin arm (Fig. 4A [DOI](#)), suggesting that phosphorylation of KIN-G does not affect its subcellular localization. Western blotting confirmed the knockdown of endogenously PTP-tagged KIN-G and the ectopic expression of triple HA-tagged wild-type and mutant KIN-G (Fig. 4B [DOI](#)). Expression of wild-type KIN-G and KIN-G^{T301A}, but not KIN-G^{T301D}, both rescued the growth defects of KIN-G RNAi cells (Fig. 4C [DOI](#)). We further quantitated the cell types of different nucleus and kinetoplast configurations in KIN-G RNAi cells and the three complementation cells. The KIN-G^{T301D} complementation cells, but not the KIN-G complementation cells and the KIN-G^{T301A}

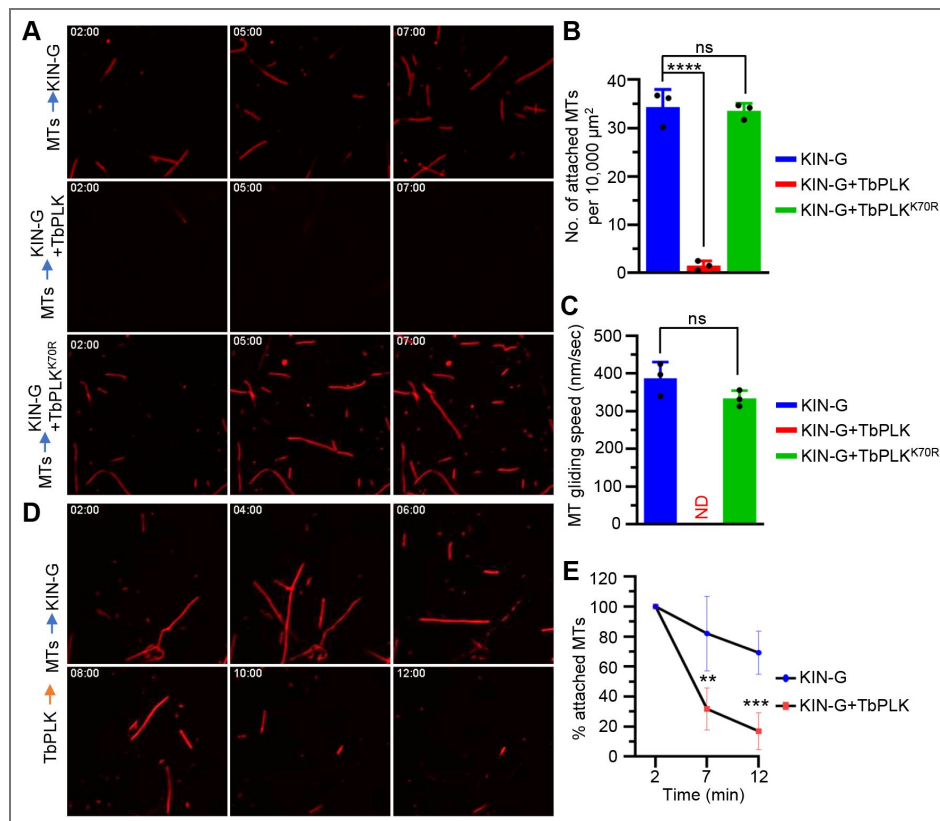


Figure 2. Phosphorylation of KIN-G by TbPLK disrupts the microtubule-binding activity of KIN-G.

(A). Pre-incubation of KIN-G with TbPLK, but not TbPLK^{K70R}, disrupted KIN-G microtubule-binding activity. KIN-G, KIN-G and TbPLK mixture, and KIN-G and TbPLK^{K70R} mixture were first attached to coverslips and then microtubules (MTs) were added into the chamber. (B). Quantitation of KIN-G-bound microtubules in the presence or absence of TbPLK or in the presence of TbPLK^{K70R}. Error bars indicate S.D. from three independent experiments. ****: $p < 0.0001$; ns: no significance (one-way ANOVA). (C). Measurement of the microtubule-gliding speed of KIN-G in the presence or absence of TbPLK or in the presence of TbPLK^{K70R}. Error bars indicate S.D. from three independent experiments. ND: not done. ns: no significance (one-way ANOVA). (D). TbPLK disrupted KIN-G microtubule-binding activity. KIN-G was first attached to coverslips, MTs were added into the chamber, and finally TbPLK was added into the chamber. (E). Quantitation of KIN-G-bound microtubules following the incubation with or without TbPLK. Error bars indicate S.D. from three independent experiments. **: $p < 0.01$; ***: $p < 0.001$ (Student's *t*-test).

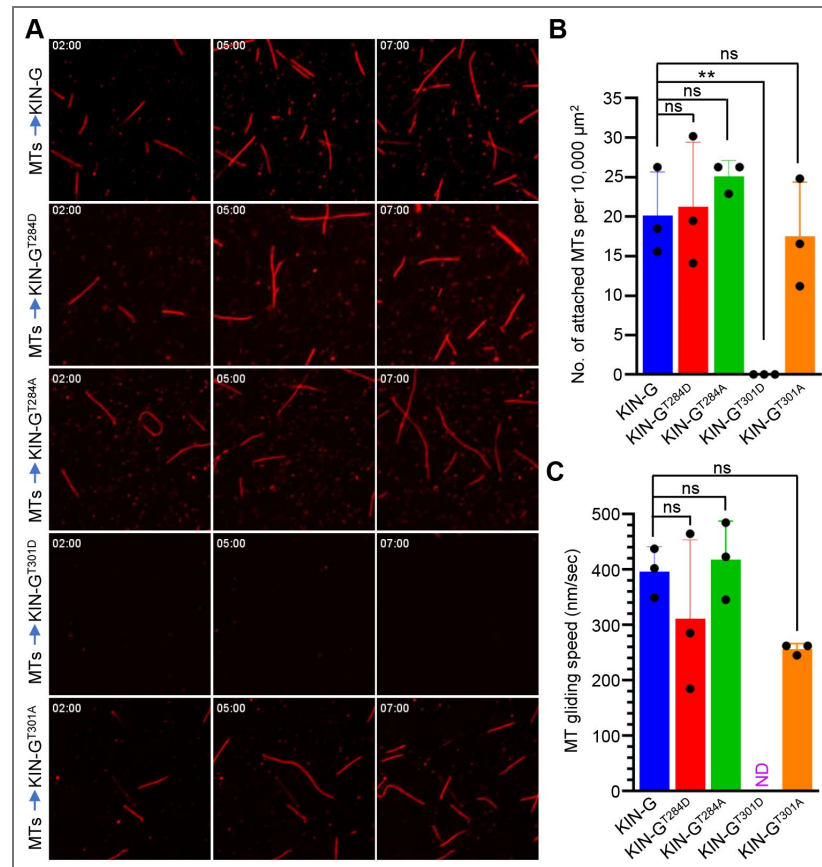


Figure 3. Phosphorylation of Thr301 on KIN-G by TbPLK disrupts the microtubule-binding activity of KIN-G.

(A). Microtubule-binding activity of KIN-G and its Thr284 and Thr301 mutants. (B). Quantitation of the bound microtubules of KIN-G and its mutants. Error bars indicate S.D. from three independent experiments. **: $p < 0.01$; ns: no significance (one-way ANOVA). (C). Measurement of the microtubule-gliding speed of KIN-G and its mutants. Error bars indicate S.D. from three independent experiments. ND: not done. ns: no significance (one-way ANOVA).

complementation cells, accumulated 2N1K (two nuclei and one kinetoplast) cells and xNyK ($x \geq 2$; $y \geq 1$) cells, similar to the phenotype caused by KIN-G RNAi (Fig. 4D [↗](#)). These results suggest that Thr301 phosphorylation disrupted the cellular function of KIN-G in trypanosomes.

Phosphorylation of Thr301 in KIN-G disrupts centrin arm and Golgi biogenesis

The potential effect of Thr301 phosphorylation on the function of KIN-G in centrin arm biogenesis was examined by immunofluorescence microscopy using anti-TbCentrin4 and the pan-centrin antibody 20H5. Measurement of the length of the new and old centrin arm structures showed that many KIN-G^{T301D} complementation cells contained a shorter new centrin arm, with the average length of the new centrin arm significantly reduced after tetracycline induction for 48 h (Fig. 5A, B [↗](#)). The results mimicked the deficiency in the biogenesis of the new centrin arm in KIN-G RNAi cells (Zhou et al., 2024 [↗](#)), suggesting that phosphorylation of Thr301 in KIN-G disrupted its function in promoting centrin arm biogenesis.

The potential effect of Thr301 phosphorylation on Golgi biogenesis was also investigated by immunofluorescence microscopy using the anti-TbGRASP antibody. We endogenously epitope-tagged the COPII coat protein subunit Sec13 with mCherry to serve as a marker for the ER exit site (ERES). In non-induced control, two major Golgi/ERES (strong TbGRASP/Sec13 fluorescence signal) were detected in bi-nucleated cells, and one or two smaller Golgi (weaker TbGRASP signal), each of which associated with a smaller ERES (weaker Sec13 signal), were also detected near the major Golgi/ERES in some bi-nucleated cells (Fig. 5C [↗](#), open arrows and solid arrowheads, respectively). In KIN-G^{T301D} complementation cells, the number of bi-nucleated cells containing two, three, or four Golgi/ERES was significantly reduced, whereas the number of bi-nucleated cells containing more than four Golgi/ERES was significantly increased (Fig. 5C, D [↗](#)). These results suggest that phosphorylation of Thr301 in KIN-G disrupted its function in promoting Golgi biogenesis.

Phosphorylation of Thr301 in KIN-G impairs FAZ elongation and flagellum positioning

We investigated the potential effect of Thr301 phosphorylation on the elongation of the new FAZ by immunofluorescence microscopy using the anti-CC2D antibody (Zhou et al., 2011 [↗](#)), which showed that the length of the new FAZ in the bi-nucleated KIN-G^{T301D} complementation cells was significantly shorter than that of the non-induced bi-nucleated cells (Fig. 6A, B [↗](#)), suggesting that Thr301 phosphorylation impairs FAZ elongation. Notably, in these bi-nucleated KIN-G^{T301D} complementation cells, the size of the NFD cell was also smaller, which is positively correlated to the length of the new FAZ (Fig. 6C [↗](#)), in agreement with the finding that the FAZ length determines the cell size in trypanosomes (Zhou et al., 2011 [↗](#)), acting as the “cellular ruler” of trypanosome cell morphology (Sunter and Gull, 2016 [↗](#)).

The formation of a shorter new FAZ in KIN-G^{T301D} complementation cells prompted us to examine the segregation of the flagellum-associated cytoskeletal structures. Co-immunofluorescence microscopy with YL1/2, which labels the mature basal body (Woods et al., 1989 [↗](#)), and anti-TbBILBO1 antibody, which labels the flagellar pocket collar (Bonhivers et al., 2008 [↗](#)), showed that in the bi-nucleated KIN-G^{T301D} complementation cells the duplicated basal bodies/flagellar pocket collars were not far segregated (Fig. 6D [↗](#)), with the average distance between the duplicated structures reduced significantly after tetracycline induction for 48 h (Fig. 6E, F [↗](#)). These results suggest that phosphorylation of Thr301 inhibits the segregation of flagellum-associated cytoskeletal structures. As the flagellum is nucleated from the mature basal body, these results suggest that Thr301 phosphorylation inhibits flagellum positioning.

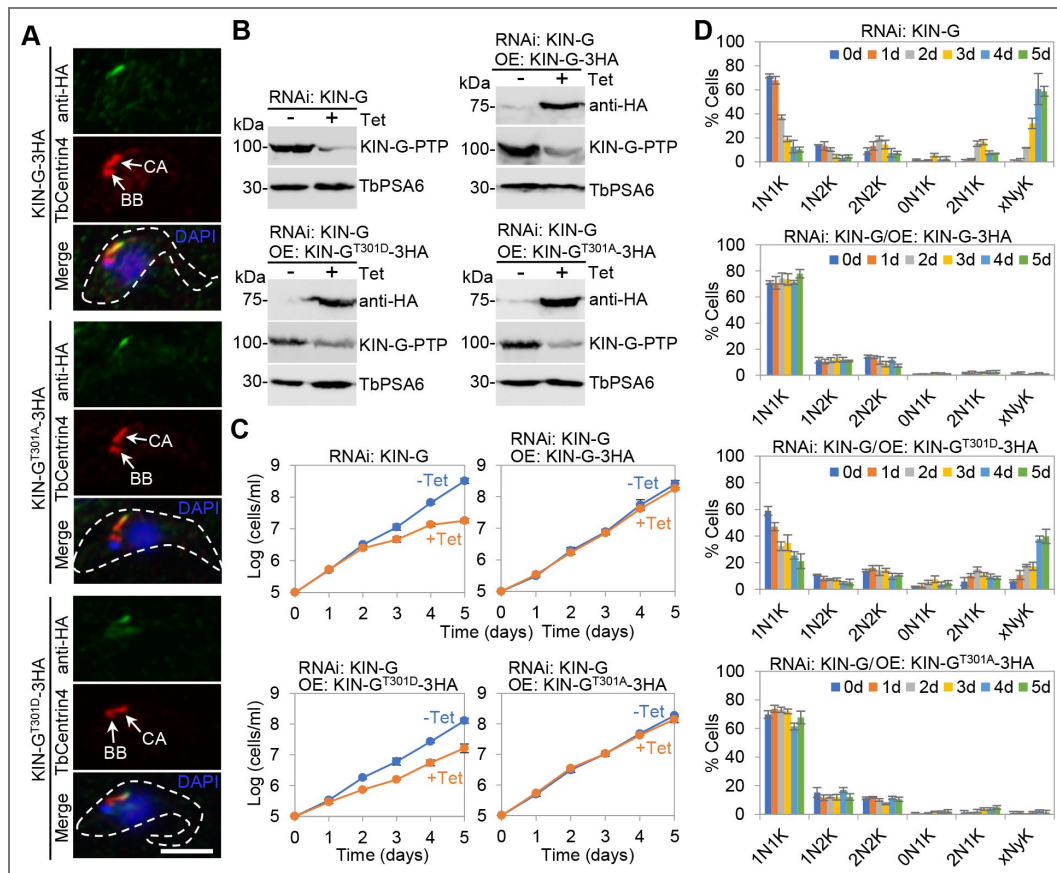


Figure 4. Expression of Thr301 phospho-mimic mutant of KIN-G disrupts cell proliferation.

(A). Subcellular localization of ectopically expressed KIN-G, KIN-G^{T301A}, KIN-G^{T301D}, co-stained with TbCentrin4. BB: basal body; CA: centrin arm. Scale bar: 5 μ m. (B). Western blotting to detect the levels of ectopically 3HA-tagged KIN-G and its mutants and the endogenously PTP-tagged KIN-G before and after tetracycline induction for 48 hours. TbPSA6 serves as a loading control. (C). Growth curves of KIN-G RNAi cell line and its complementation cell lines expressing KIN-G, KIN-G^{T301A}, or KIN-G^{T301D}. OE: overexpression. Error bars indicate S.D. from three independent experiments. (D). Quantitation of the numbers of nuclei (N) and kinetoplasts (K) of KIN-G RNAi cell line and its complementation cell lines. Error bars indicated S.D. from three independent experiments.

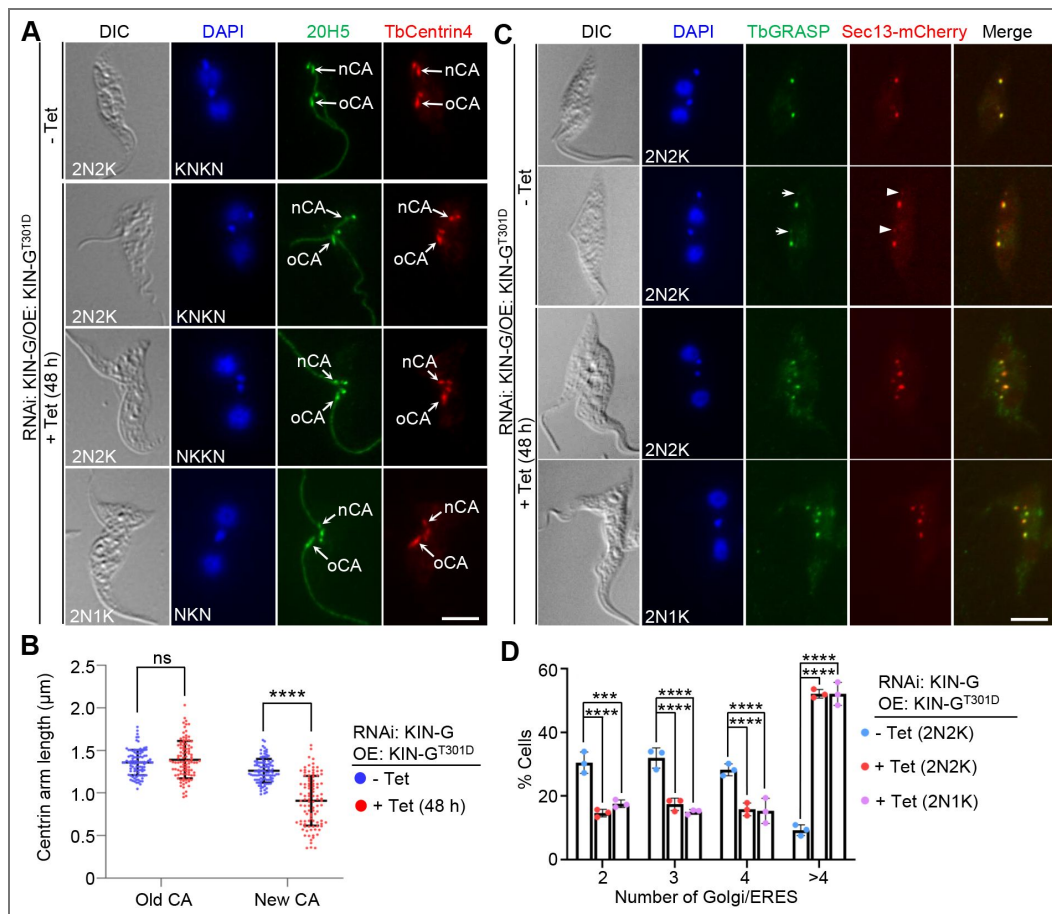


Figure 5. TbPLK phosphorylation of Thr301 on KIN-G disrupts centrin arm and Golgi biogenesis.

(A). Effect of the expression of KIN-G^{T301D} on centrin arm formation. Cells were co-immunostained with the pancentrin antibody 20H5 and the anti-TbCentrin4 antibody. Scale bar: 5 μm. (B). Measurement of centrin arm length in KIN-G RNAi cells expressing Thr301 phospho-mimic of KIN-G before and after tetracycline induction. Error bars indicate S.D. ns: no significance; ****: $p < 0.0001$ (Student's t -test). (C). Effect of the expression of KIN-G^{T301D} on the biogenesis of Golgi and ER exit site (ERES). Cells were co-immunostained with the anti-TbGRASP antibody to detect TbGRASP. ERES was labeled by mCherry-tagged Sec13. Scale bar: 5 μm. (D). Quantitation of cells with different numbers of Golgi/ERES in control and KIN-G RNAi cells. Error bars indicate S.D. from three independent experiments. ***: $p < 0.001$; ****: $p < 0.0001$ (one-way ANOVA).

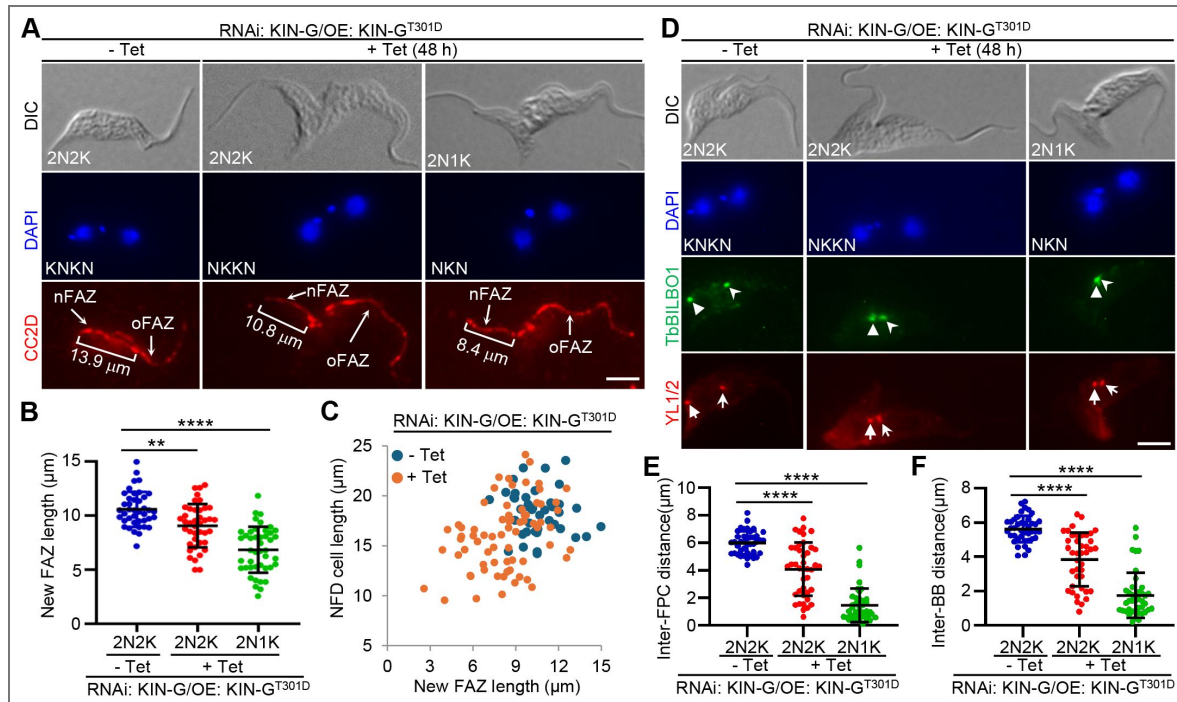


Figure 6. TbPLK phosphorylation of Thr301 on KIN-G impairs flagellar inheritance.

(A) Effect of the expression of KIN-G^{T301D} on the elongation of the flagellum attachment zone (FAZ) filament. Cells were immunostained with the anti-CC2D antibody. Scale bar: 5 μ m. **(B)** Measurement of the length of the new and the old FAZ filaments in control and KIN-G RNAi cells expressing KIN-G^{T301D}. Error bars indicate S.D. **: $p < 0.01$; ****: $p < 0.0001$ (one-way ANOVA). **(C)** Measurement of the cell body length of the new-flagellum daughter (NFD) cell and its correlation with the length of the new FAZ for control cells and KIN-G RNAi cells expressing KIN-G^{T301D}. **(D)** Effect of the expression of KIN-G^{T301D} on the segregation of flagellar pocket collar (FPC) and basal body (BB). Cells were co-immunostained with anti-TbBILBO1 antibody and YL1/2 antibody. Scale bar: 5 μ m. **(E, F)** Measurement of the inter-FPC distance **(E)** and inter-BB distance **(F)** in control cells and KIN-G RNAi cells expressing Thr301 phospho-mimic mutant of KIN-G. Error bars indicate S.D. ****: $p < 0.0001$ (one-way ANOVA).

Phosphorylation of Thr301 in KIN-G causes mis-placement of the cell division plane

Knockdown of KIN-G by RNAi caused cytokinesis defects (Zhou et al., 2024 [DOI](#)), likely due to the mis-placement of the cell division plane, but this phenotype was not explored previously. Therefore, we investigated the potential defect in cell division plane placement in KIN-G RNAi cells by analyzing the localization of the orphan kinesin protein KLIF, which marks the cell division plane during cytokinesis in trypanosomes (Zhou et al., 2018a [DOI](#)). In the non-induced control cells, the leading edge of the KLIF-labeled cell division plane was always placed at the mid-portion of the ventral side of the NFD cell (Fig. 7A [DOI](#), yellow arrow). In contrast, in the dividing KIN-G RNAi cells, the leading edge of the KLIF-labeled cell division plane was placed near the posterior cell tip (Fig. 7A [DOI](#), yellow arrows). We further investigated the effect of KIN-G RNAi on the positioning of the nascent posterior of the OFD cell, which is formed near the mid-portion of the ventral edge of the NFD cell through microtubule bundling and cytoskeleton remodeling (Wheeler et al., 2013 [DOI](#)). We used the epitope-tagged GB4 protein (Fig. 7B [DOI](#)), a microtubule plus end-localized protein (Rindisbacher et al., 1993 [DOI](#)), as a marker for the cell posterior, and performed immunofluorescence microscopy. In control cells, the GB4-labeled nascent posterior was located at the mid-portion of the ventral side of the NFD cell, whereas in KIN-G RNAi cells, the GB4-labeled nascent posterior was placed next to the GB4-labeled existing posterior (Fig. 7B [DOI](#)), which was confirmed by measurement of the inter-posterior distance (Fig. 7C [DOI](#)).

The KIN-G^{T301D} complementation cells also appear to have cytokinesis defects (Fig. 4C, D [DOI](#)). The number of bi-nucleated cells undergoing cytokinesis or with a visible cleavage furrow was significantly increased upon tetracycline induction (Fig. 7D, E [DOI](#)), suggesting that the KIN-G^{T301D} complementation cells initiated cytokinesis, but failed to complete cytokinesis. Notably, those cells with a visible cleavage furrow appeared to contain a mis-placed cell division plane in the KIN-G^{T301D} complementation cells (Fig. 7D [DOI](#), yellow arrows). The mis-placement of the cell division plane was more prominent in the bi-nucleated cells with the NKKN configuration and the NKN configuration, in which the duplicated kinetoplasts were only partly separated or not separated, respectively, than in the bi-nucleated cells with the KNKN configuration (Fig. 7F [DOI](#)). Therefore, phosphorylation of Thr301 in KIN-G disrupted cell division plane placement, mimicking the defect caused by KIN-G RNAi.

Discussion

In this report, we have identified the orphan kinesin KIN-G as a substrate of TbPLK, one of the two Polo-like kinase homologs in *T. brucei* (Kurasawa et al., 2020 [DOI](#)), and discovered a negative role of TbPLK phosphorylation in regulating the biochemical activity and cellular function of KIN-G in the procyclic form of *T. brucei*. TbPLK is known to play multiple roles in basal body rotation, centrin arm assembly, FAZ elongation, cytokinesis, and Golgi biogenesis (de Graffenried et al., 2013 [DOI](#); de Graffenried et al., 2008 [DOI](#); Hammarton et al., 2007 [DOI](#); Ikeda and de Graffenried, 2012 [DOI](#); Kumar and Wang, 2006 [DOI](#); Li et al., 2010 [DOI](#); Lozano-Nunez et al., 2013 [DOI](#)), but the mechanistic roles of TbPLK in regulating these cellular processes have not been fully elucidated, mainly because most of its substrates have not been identified. TbPLK phosphorylates the basal body-localized protein SPBB1 (Hu et al., 2015 [DOI](#)), the centrin arm protein TbCentrin2 (de Graffenried et al., 2013 [DOI](#); de Graffenried et al., 2008 [DOI](#)), and the cytokinesis regulators CIF1 and CIF2 (Kurasawa et al., 2022 [DOI](#)), which exemplifies its role in regulating the rotation/segregation of basal body, the biogenesis of centrin arm and Golgi, and the initiation of cytokinesis, respectively. Phosphorylation of TbCentrin2 by TbPLK at Ser54 does not appear to be essential, as expression of the Ser54 phospho-deficient mutant TbCentrin2^{S54A} caused weak growth defects, whereas expression of the Ser54 phospho-mimic mutant TbCentrin2^{S54D} caused severe growth defects, suggesting that the dephosphorylation of TbCentrin2 at Ser54 is required for TbCentrin2 to execute its function in centrin arm assembly (de Graffenried et al., 2013 [DOI](#)). However, it remains unclear how phosphorylation of Ser54 by TbPLK contributes to the regulation of TbCentrin2 function and what protein phosphatase is responsible for TbCentrin2 dephosphorylation. KPP1 is potentially

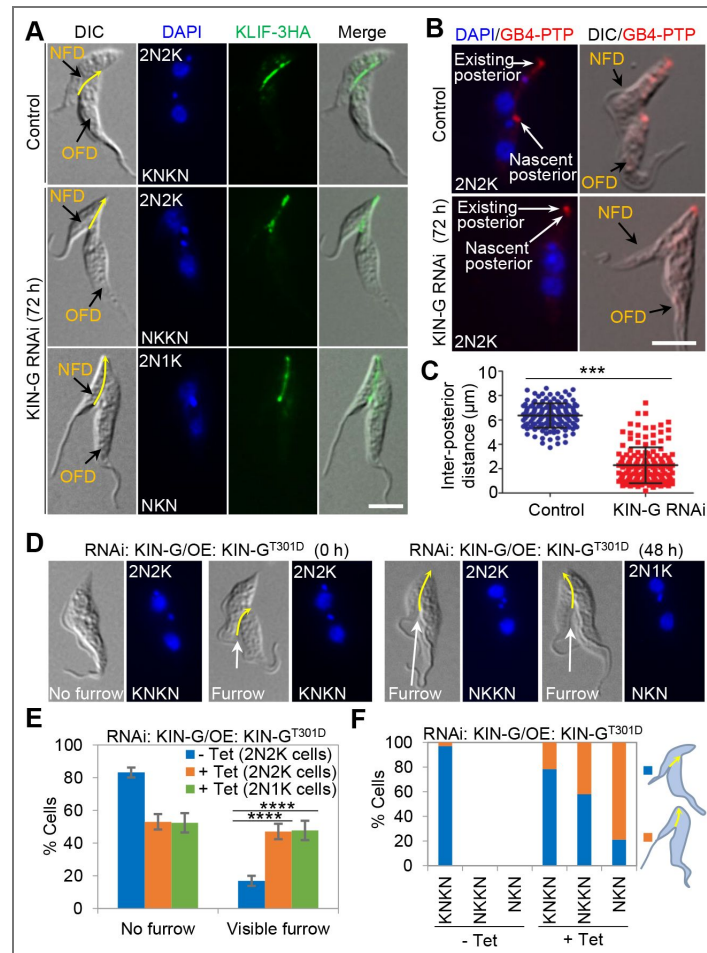


Figure 7. Phosphorylation of KIN-G at Thr301 by TbPLK disrupts cell division plane placement.

(A). Immunofluorescence microscopy to detect the cell division plane with endogenous triple HA-tagged KLIF in dividing cells from non-induced control and KIN-G RNAi cells. Yellow arrows indicate the KLIF-marked cell division plane. NFD: new-flagellum daughter; OFD: old-flagellum daughter. Scale bar: 5 μm. (B). Immunofluorescence microscopy to detect the NFD posterior and the OFD nascent posterior with PTP-tagged GB4 protein. Scale bar: 5 μm. (C). Measurement of the inter-posterior distance of bi-nucleated cells from non-induced control and KIN-G RNAi cells. Error bars indicate S.D. *** $p < 0.001$ (Student's *t*-test). (D). Non-induced and tetracycline-induced KIN-G RNAi cells expressing KIN-G^{T301D}. Shown are a non-dividing cell without a visible cleavage furrow and three dividing cells with a visible cleavage furrow. Yellow arrows indicate the cell division plane. Scale bar: 5 μm. (E). Quantitation of bi-nucleated cells with or without a visible cleavage furrow from non-induced and tetracycline-induced KIN-G RNAi cells expressing KIN-G^{T301D}. Error bars indicate S.D. from three independent experiments. **** $p < 0.0001$ (one-way ANOVA). (F). Percentage of dividing bi-nucleated cells with a normally placed cell division plane or an abnormally placed cell division plane from non-induced and tetracycline-induced KIN-G RNAi cells expressing KIN-G^{T301D}.

responsible for TbCentrin2 dephosphorylation (Zhou et al., 2018b), but this has not been experimentally verified. The previous work (de Graffenried et al., 2013) and our current work suggest that regulation of centrin arm assembly in procyclic trypanosomes involves the phosphorylation of multiple centrin arm-localized proteins, such as TbCentrin2 and KIN-G, by TbPLK.

One of the intriguing questions is how TbPLK-phosphorylation of KIN-G negatively regulates the function of KIN-G. KIN-G is a microtubule plus end-directed motor protein, and its motor activity is required for cellular function (Zhou et al., 2024), which suggests that KIN-G needs to travel along microtubules to execute its function. The best candidate microtubules that KIN-G may associate with are the specialized set of four microtubules (MtQ), which have their plus ends directed toward the distal tip of the cell in *T. brucei* (Gull, 1999; Robinson et al., 1995). Presumably, KIN-G may be first recruited to the centrin arm through binding to WDR2 (Zhou et al., 2025) and then it associates with and travels along the MtQ to transport cargos in the centrin arm region to facilitate centrin arm assembly. In this regard, phosphorylated KIN-G is unable to bind to MtQ due to the loss of microtubule-binding activity (Figs. 2 and 3) and, hence, is unable to transport its cargos to their destinations. The lack of rescue of KIN-G RNAi by the expression of the phospho-mimic mutant KIN-G^{T301D} and the complete rescue of KIN-G RNAi by the expression of the phospho-deficient mutant KIN-G^{T301A} (Fig. 4) support this notion, and these findings further suggest that dephosphorylation of Thr301, but not phosphorylation of Thr301, is required for KIN-G function. The regulation of KIN-G function by TbPLK phosphorylation appears to be similar to the regulation of TbCentrin2 by TbPLK phosphorylation (de Graffenried et al., 2013). The best candidate protein phosphatase for KIN-G dephosphorylation is the centrin arm-localized KPP1 (Zhou et al., 2018b), which also attenuates TbPLK kinase activity by dephosphorylating TbPLK at Thr125 (An et al., 2021).

We have provided evidence to demonstrate that phosphorylation of Thr301 in KIN-G disrupted its cellular function, causing defects in centrin arm assembly, FAZ elongation, flagellum positioning, cell division plane placement, cytokinesis, and Golgi biogenesis (Figs. 5-7), which mimics the defects caused by KIN-G RNAi (Zhou et al., 2024). There is an intriguing question of why KIN-G appears to regulate so many cellular processes, while it only localizes to the centrin arm throughout the cell cycle (Zhou et al., 2024). It should be noted that previous work showed that the deficiency in centrin arm integrity impairs FAZ elongation, flagellum positioning, cell division plane placement, cytokinesis, and Golgi biogenesis (de Graffenried et al., 2013; de Graffenried et al., 2008; He et al., 2005; Zhou et al., 2010; Zhou et al., 2025; Zhou et al., 2024). Since the proximal end of the FAZ is embedded within the hook complex between the centrin arm and the shank part of the hook complex (Morriswood et al., 2013) and assembly of the FAZ occurs from its proximal end (Sunter et al., 2015; Zhou et al., 2015), the primary role of the centrin arm likely is to act as a hub to coordinate FAZ assembly. In this regard, all of the other defects described above, except Golgi biogenesis, are the secondary effects of the defective FAZ assembly. The impairment of Golgi biogenesis by the deficiency in centrin arm integrity led to the interpretation that centrin arm determines the site where Golgi is assembled (He et al., 2005), although the underlying mechanism has not been established. Presumably, there is a physical connection between the centrin arm and the Golgi apparatus, which may be mediated by the scaffold protein CAAP1, because CAAP1 resides in between the centrin arm and the Golgi apparatus and depletion of CAAP1 disconnects the two structures and disrupts Golgi biogenesis (Zhou et al., 2024). All in all, it appears that deficiency in centrin arm integrity disrupts FAZ elongation and Golgi biogenesis, the former of which further causes flagellum mis-positioning, cell division plane mis-placement, and unequal cytokinesis. Therefore, the primary function of KIN-G is to regulate the formation of the centrin arm. Phosphorylation of Thr301 disrupts the microtubule-binding activity of KIN-G and, hence, the plus end-directed motility along the MtQ, suggesting that the regulation of centrin arm assembly by KIN-G requires its motor activity, in agreement with the results obtained with the expression of the motility-dead mutant KIN-G^{G266A} in KIN-G RNAi cells, which had almost identical phenotypes as the KIN-G^{T301D} complementation cells (Zhou et al., 2024).

The defect in FAZ assembly due to defective centrin arm assembly, observed in KIN-G RNAi cells (Zhou et al., 2024 [DOI](#)) and the KIN-G^{T301D} complementation cells (Fig. 7 [DOI](#)), is believed to be the cause of the mis-positioning of the newly synthesized flagellum and the mis-placement of the cell division plane. Previous work has demonstrated that positioning of the newly synthesized flagellum relies on the faithful duplication/segregation of its associated cytoskeletal structures, including the FAZ (Ikeda and de Graffenried, 2012 [DOI](#); Zhou et al., 2011 [DOI](#)). Elongation of the new FAZ appears to promote basal body migration (Zhou et al., 2011 [DOI](#)), thereby facilitating flagellum segregation. The cell division plane in procyclic trypanosomes is placed along the longitudinal cell axis from the anterior tip of the NFD cell to the nascent posterior of the OFD cell (Wheeler et al., 2013 [DOI](#)), and its placement appears to depend on the length of the new flagellum (Kohl et al., 2003 [DOI](#)) and the length of the new FAZ (Zhou et al., 2011 [DOI](#)). However, the mis-placement of the cell division plane in CC2D RNAi cells containing a short-length new FAZ but a normal-length new flagellum (Zhou et al., 2011 [DOI](#)) argues that it is the length of the new FAZ that determines the cell division plane. This notion is supported by the observed positive correlation between the length of the new FAZ and the length of the NFD cell in the control cells, CC2D RNAi cells (Zhou et al., 2011 [DOI](#)), FAZ10 RNAi cells (Moreira et al., 2017 [DOI](#)), KIN-G RNAi cells (Zhou et al., 2024 [DOI](#)), and the KIN-G^{T301D} complementation cells (Fig. 6C [DOI](#)). The mis-positioning of the new flagellum and the mis-placement of the cell division plane collectively contributed to the defects in cytokinesis and cell proliferation.

The complete rescue of KIN-G RNAi phenotypes by expression of the KIN-G^{T301A} mutant (Fig. 4 [DOI](#)) raised an interesting question about the role of TbPLK phosphorylation of KIN-G in trypanosome cells. TbPLK phosphorylates KIN-G throughout the entire protein, and Thr301 is one of the ten *in vitro* and *in vivo* phosphosites that was characterized in this work. Four other *in vitro* and *in vivo* phosphosites in the motor domain of KIN-G are located outside of the microtubule-binding motifs and the nucleotide-binding motifs (Fig. 1D [DOI](#)); whether the phosphorylation/dephosphorylation of these sites may affect microtubule binding or nucleotide binding of KIN-G is unknown and remains to be explored. Five *in vitro* and *in vivo* phosphosites are in the C-terminus of KIN-G, with three sites located in the three coiled-coil motifs and two sites located in the loop region between coiled-coil motifs #2 and #3 (Fig. 1D [DOI](#)). The potential role of phosphorylation of these sites by TbPLK is also unknown and needs further exploration. However, because the C-terminal tail in kinesin proteins is known to bind cargo or cargo adaptor proteins (Hirokawa et al., 1998 [DOI](#)), the phosphorylation of the five residues in the C-terminus of KIN-G may impact the binding of KIN-G to its cargo or cargo adaptor proteins. Future experiments will be directed to exploring the potential biochemical and physiological roles of these phosphosites, which may uncover the other role(s) of TbPLK-mediated phosphorylation of KIN-G.

In summary, we have identified the orphan kinesin KIN-G as a substrate of TbPLK and demonstrated that phosphorylation of Thr301 by TbPLK disrupted KIN-G microtubule-binding activity. We further showed that expression of the phospho-mimic mutant KIN-G^{T301D} in trypanosome cells impaired the cellular function of KIN-G. Therefore, dephosphorylation of Thr301 in KIN-G is necessary for KIN-G to exert its function in regulating centrin arm assembly, thereby facilitating Golgi biogenesis and FAZ elongation, which further promotes flagellum positioning and cell division plane placement.

Materials and Methods

Trypanosome cell culture, RNAi, and complementation

The procyclic form of *T. brucei* strain Lister427 was cultured in the SDM-79 medium at 27°C. The KIN-G RNAi cell line, which was generated previously (Zhou et al., 2024 [DOI](#)), was cultured in SDM-79 medium supplemented with 10% heat-inactivated fetal bovine serum (Sigma-Aldrich), 15 µg/ml G418, 50 µg/ml hygromycin, and 2.5 µg/ml phleomycin at 27°C. Cells were diluted with fresh medium whenever the cell density reached 5×10⁶/ml. RNAi was induced with 1 µg/ml tetracycline, and cell growth was monitored daily with a hemacytometer.

The coding sequence of KIN-G was recoded to make it RNAi resistant, and the recoded KIN-G gene was cloned into pLew100-3HA-PAC vector. The pLew100-KIN-G-3HA-PAC plasmid was then used for site-directed mutagenesis to mutate the TbPLK-phosphorylated residues to aspartate or alanine, and then the resulting plasmids were used to transfect the KIN-G RNAi cell line. Transfectants were selected with 1 µg/ml puromycin in addition to 15 µg/ml G418, and 50 µg/ml hygromycin, and 2.5 µg/ml phleomycin, and then cloned by limiting dilution in a 96-well plate containing SDM-79 medium supplemented with appropriate antibiotics and 20% fetal bovine serum. Expression of recoded KIN-G and its phospho-mimic and phospho-deficient mutants, and knockdown of endogenous KIN-G were induced with 1 µg/ml tetracycline. Cell growth was monitored by daily counting of the cells with a hemacytometer.

In situ epitope tagging of proteins

Epitope tagging of KIN-G, KLIF, GB4, and Sec13 from their respective endogenous locus was carried out using the PCR-based method (Shen et al., 2001 [DOI](#)). KIN-G was tagged with a C-terminal triple HA epitope or PTP epitope, KLIF was tagged with a C-terminal triple HA epitope, GB4 was tagged with a C-terminal PTP epitope, and Sec13 was tagged with an N-terminal mCherry tag. Transfectants were selected with appropriate antibiotics, and clonal cell lines were obtained by limiting dilution in a 96-well plate as described above.

Immunofluorescence microscopy

To prepare intact trypanosome cells for immunofluorescence microscopy, cells were washed once with PBS, settled onto glass coverslips, fixed with methanol at −20°C for 30 min, and rehydrated with PBS at room temperature. To prepare trypanosome cytoskeletons for immunofluorescence microscopy, cells were washed once with PBS, adhered onto glass coverslips, treated with 1% Nonidet-P40 in PEME buffer (100 mM PIPES, pH6.9, 2 mM EGTA, 1 mM MgSO₄, and 0.1 mM EDTA) for 2 seconds at room temperature, and then fixed with cold methanol at −20°C for 30 min. Intact cells or cytoskeletons on the coverslips were incubated with the blocking buffer (3% BSA in PBS) at room temperature for 1 h, and then incubated with the primary antibody at room temperature for 1 h. The following primary antibodies were used: anti-HA monoclonal antibody (1:400 dilution, Sigma-Aldrich), anti-Protein A polyclonal antibody (1:400 dilution, Sigma-Aldrich), anti-CC2D polyclonal antibody (1:1,000 dilution) (Zhou et al., 2011 [DOI](#)), 20H5 monoclonal antibody (1:400 dilution) (He et al., 2005 [DOI](#)), YL 1/2 monoclonal antibody (1: 1000 dilution) (Woods et al., 1989 [DOI](#)), anti-TbCentrin4/LdCentrin1 polyclonal antibody (1:1000 dilution) (Selvapandiyan et al., 2007 [DOI](#)), anti-TbBILBO1 antibody (1: 400 dilution) (Bonhivers et al., 2008 [DOI](#)), and anti-TbGRASP polyclonal antibody (1:400 dilution) (He et al., 2004 [DOI](#)). Cells or cytoskeletons on the glass coverslips were washed three times (5 min each) with PBS and incubated with secondary antibodies at room temperature for 1 h. The following secondary antibodies were used: FITC-conjugated anti-mouse IgG, FITC-conjugated anti-rabbit IgG, Cy3-conjugated anti-rabbit IgG, and Cy3-conjugated anti-rat IgG (all from MilliporeSigma). Cells or cytoskeletons were washed three times (5 min each) with PBS, mounted with DAPI-containing VectaShield mounting medium (Vector Labs), and imaged with the Olympus IX71 fluorescence microscope. Images were acquired and processed using the Slidebook 5 software.

Expression and purification of recombinant proteins

To express and purify recombinant proteins for *in vitro* microtubule-binding and gliding assays, KIN-G was cloned into the pET26 vector, and the resulting plasmid was used to generate the phospho-deficient and phospho-mimic mutants by site-directed mutagenesis. The full-length TbPLK was cloned into the pET41a vector, and the resulting plasmid was used to generate the kinase-dead mutant TbPLK^{K70R} by site-directed mutagenesis. These plasmids were each used to transform the *E. coli* BL21 strain. Expression of recombinant proteins was induced with 0.1 mM isopropyl β-d-thio-galactopyranoside at room temperature for KIN-G-6His and GST-TbPLK-6His and at 15°C overnight for GST-TbPLK^{K70R}-6His. Bacterial cells expressing KIN-G-6His and its mutants, GST-TbPLK-6His, and GST-TbPLK^{K70R}-6His were lysed by sonication in bacteria lysis

buffer (50 mM NaH_2PO_4 , 300 mM NaCl, 10 mM imidazole, pH 8.0), and cell lysate was cleared by centrifugation at $20,000 \times g$ for 10 min at 4°C . Cleared cell lysate was incubated with the Chelating Sepharose Fast Flow (GE Healthcare) beads charged with nickel ion at 4°C for 30 min. Beads were washed five times with 1 ml wash buffer (50 mM NaH_2PO_4 , 1 M NaCl, 120 mM imidazole, pH 8.0) for each wash. Recombinant proteins were eluted with elution buffer (50 mM NaH_2PO_4 , 300 mM NaCl, 250 mM imidazole, pH 8.0), and eluted proteins were concentrated and buffer-exchanged to the kinase assay buffer (10 mM HEPES, pH 7.5, 10 mM MgCl_2 , 50 mM NaCl, 1 mM EGTA, 1 mM DTT) with Amicon Ultra Centrifugal Filters 10K (MilliporeSigma).

***In vitro* kinase assay using the thiophosphorylation method**

In vitro kinase assay was performed by using a semisynthetic epitope to detect thiophosphorylated kinase substrates (Allen et al., 2007 [DOI](#)). Purified recombinant proteins were incubated in the kinase assay buffer (10 mM HEPES pH7.5, 10 mM MgCl_2 , 50 mM NaCl, and 0.4 mM DTT) supplemented with 1 mM ATP- γ -S at room temperature for 30 min. 50 mM p-nitrobenzyl mesylate (PNBM), which alkylates thiophosphates to form thiophosphate ester epitopes that can be recognized by the anti-thiophosphate ester antibody, was added to the kinase reaction and was incubated at room temperature for 60 min. Thiophosphorylated proteins were separated by SDS-PAGE, transferred onto a PVDF membrane, and then immunoblotted with the anti-thiophosphate ester (anti-ThioP) monoclonal antibody (1:5,000 dilution, ThermoFisher).

Identification of TbPLK-phosphosites in KIN-G by mass spectrometry

Purified recombinant KIN-G-6His was mixed with purified recombinant GST-TbPLK-6His in the kinase assay buffer (see above) containing 0.2 mM ATP at room temperature for 30 min. Thiophosphorylated proteins were separated by SDS-PAGE and stained with coomassie blue dye. The gel slice containing the phosphorylated KIN-G-6His band was excised, and incubated with 160 ng trypsin at 37°C for 4 h, according to published procedures (Rosenfeld et al., 1992 [DOI](#)). Peptides were extracted with 50 ml of 50% acetonitrile and 5% formic acid, dried using SpeedVac, resuspended in 2% acetonitrile and 0.1% formic acid, and injected onto Thermo LTQ Orbitrap XL (ThermoFisher Scientific), according to published procedures (Lee and Li, 2021 [DOI](#)). Peptide samples were analyzed on an LTQ Orbitrap XL interfaced with an Eksigent nano-LC 2D plus ChipLC system (Eksigent Technologies). Samples were loaded onto a ChromXP C18-CL trap column (200 mm i.d. x 0.5 mm length) at a flow rate of 3 nL/min. Reverse-phase C18 chromatographic separation of peptides was carried on a ChromXP C18-CL column (75 mm i.d x 10 cm length) at 300 nL/min. The LTQ Orbitrap was operated in a data-dependent mode to simultaneously measure full-scan MS spectra in the Orbitrap and the five most intense ions in the LTQ by CID, respectively. In each cycle, MS1 was acquired at a target value of $1\text{E}6$ with a resolution of 100,000 (m/z 400) followed by top five MS2 scan at a target value of $3\text{E}4$. The mass spectrometric setting was: spray voltage at 1.6 KV, and charge state screening and rejection of singly charged ion enabled. Ion selection thresholds were set at 8,000 for MS2, 35% normalized collision energy, activation Q was set at 0.25, and dynamic exclusion was employed for 30 sec. Raw data files were processed and searched against the *T. brucei* proteome database using the Mascot and Sequest HT (version 13) search engines. The search conditions were set as follows: peptide tolerance of 10 p.p.m. and MS/MS tolerance of 0.8 Da, with two missed cleavages permitted and the enzyme set as trypsin.

Tubulin polymerization and microtubule binding and gliding assay

In vitro tubulin polymerization was performed according to our published procedure (Hu et al., 2024 [DOI](#)). Non-labeled porcine brain tubulin (Cytoskeleton, Inc., Cat#: T240-A80, 80 μg) was mixed with rhodamine-labeled porcine brain tubulin (Cytoskeleton, Inc., Cat#: TL590M, 20 μg) in BRB80-DTT buffer (80 mM Potassium-PIPES, pH 6.8, 1 mM MgCl_2 , 1 mM EGTA, 1 mM DTT) supplemented with 1 mM Guanylyl-(α,β)-methylene-diphosphonate (GMP-CPP). The mixture was incubated at 4°C for 5 min and then clarified by centrifugation at $353,000 \times g$ at 4°C for 5 min in a TLA120.1 rotor in an ultracentrifuge (Beckman Coulter TL-100) to remove aggregates. The supernatant was snap-

frozen in liquid nitrogen, aliquoted, and stored at -80°C . To generate microtubule seeds, an aliquot of the supernatant was diluted (1:4 v/v) with the BRB80-DTT buffer to a final concentration of 0.5 mg/ml, and incubated at 37°C for 30 min. Microtubule seeds were centrifugated at $353,000 \times g$ at 27°C for 5 min, and the pellet was resuspended in the BRB80-DTT buffer.

To polymerize microtubules from the microtubule seeds, 20 μg of rhodamine-labeled tubulin was mixed with the microtubule seeds in the presence of 1 mM GTP. Microtubules were then successively assembled by incubating at 37°C in the BRB80-DTT-GTP buffer (BRB80-DTT buffer, 1.0 mM GTP) with 0.1 μM Taxol for 20 min, 1 μM Taxol for 10 min, and 10 μM Taxol for 20 min. Assembled microtubules were diluted with 10 μM Taxol in the BRB80-DTT-GTP buffer.

Microtubule-binding and -gliding assay was conducted in a chamber assembled on a glass slide with two double-sided tapes, which were spaced 8 mm apart, and covered with a 22 mm $\times$ 30 mm cover glass. Twenty microliters of 3 μM recombinant KIN-G-6His or its mutants, which were suspended in the BRB80-DTT-ATP-Taxol buffer containing 800 ng/ml BSA, was loaded into the chamber and incubated for 3 min to allow the attachment of KIN-G-6His to the cover glass. The chamber was washed twice with 20 μl of the BRB80-DTT-ATP-Taxol buffer, and 20 μl of 120 nM rhodamine-labeled microtubules in the BRB80-DTT-ATP-Taxol buffer containing 2.25 mg/ml BSA was then loaded into the chamber. After incubation for 2 min, the chamber was observed for microtubule binding and gliding under the Nikon A1 confocal microscope, and images were captured every 5 seconds. Movie files of the AVI format were generated by NIS Elements (Nikon).

To test the effect of TbPLK phosphorylation on KIN-G microtubule binding and gliding, two experiments were performed. In experiment #1, 4 μM recombinant KIN-G-6His was pre-incubated with 2 μM GST-TbPLK-6His or GST-TbPLK^{K70R}-6His in the presence of 1 mM ATP at 37°C for 7 min, and the protein mixture was clarified by centrifugation at $21,000 \times g$ for 30 sec. The supernatant was used for microtubule-binding and -gliding assay as described above. In experiment #2, 4 μM recombinant KIN-G-6His was settled onto glass coverslip in the chamber. Microtubules were then added into the chamber, and incubated for 5 min. Subsequently, 0.2 μM GST-TbPLK-6His or the BRB80-DTT-ATP-Taxol buffer was added into the chamber, and image acquisition was started after 2 min of incubation in the Nikon A1 confocal microscope.

Microtubules bound to KIN-G or its mutants were quantified in the last image of the time-lapse imaging process or at the indicated time point of the time-lapse imaging process. Nonspecifically coverslip-attached microtubules, which showed no gliding during the time-lapse imaging process, were not counted as KIN-G-bound microtubules and, thus, were excluded. Microtubule-gliding speed was calculated using the microtubules that exhibited continuous movement for at least 100 sec (20 frames) in time-lapse imaging.

Data availability

All data are included in the manuscript and available.

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Peer reviews

Reviewer #1 (Public review):

Summary:

This manuscript identifies the orphan kinesin KIN-G as a substrate of Polo-like kinase (TbPLK) in *Trypanosoma brucei* and demonstrates that phosphorylation of Thr301 inhibits KIN-G microtubule binding and disrupts its cellular function. Using a combination of in vitro

kinase assays, phosphosite mapping, microtubule binding and gliding assays, and in vivo complementation with phosphomimetic and phosphodeficient mutants, the authors link TbPLK-mediated regulation of KIN-G to defects in centrin arm integrity, FAZ elongation, Golgi organization, flagellum positioning, and division plane placement. The study provides a mechanistic advance in understanding how TbPLK regulates centrin arm biogenesis and integrates KIN-G into the growing regulatory network controlling hook complex and FAZ assembly. Overall, the work is technically strong, internally consistent, and builds logically on previous studies from this group and others.

Strengths:

A major strength of the manuscript is the clear mechanistic link between phosphorylation of Thr301 and loss of microtubule binding activity. The use of phosphomimetic (T301D) and phosphodeficient (T301A) mutants in an RNAi-rescue framework provides a clean and convincing demonstration of functional relevance in vivo. The integration of biochemical assays with detailed cell biological phenotyping (centrin arm length, FAZ elongation, basal body segregation, and cytokinesis markers) is particularly effective and makes the central conclusion robust. The observed phenotypic cascade from centrin arm defects to FAZ and division plane abnormalities is also well aligned with existing models of trypanosome morphogenesis.

Weaknesses:

My (more or less main) concern relates to the interpretation of the Golgi phenotype. The conclusion that phosphorylation of KIN-G "impairs Golgi biogenesis" is currently based on fluorescence microscopy using TbGRASP and Sec13 markers and on quantification of the number and distribution of Golgi/ERES puncta in binucleated cells. While these data convincingly demonstrate altered Golgi/ERES number and spatial organization, they do not distinguish between true defects in Golgi biogenesis or duplication and alternative possibilities such as fragmentation, vesiculation, or mislocalization of Golgi membranes. Given the central role of Golgi-centrin arm organization in the proposed model, ultrastructural analysis (for example, by EM or electron tomography) would greatly strengthen this aspect of the study by providing direct evidence for structural alterations of the Golgi and its association with the centrin arm and ERES. Such data would elevate this part of the manuscript from a descriptive fluorescence phenotype to a true structural cell biological insight. I appreciate that this experiment goes beyond the current dataset, but it would substantially enhance the mechanistic depth of the Golgi-related conclusions and strengthen the causal chain linking centrin arm defects to Golgi abnormalities. However, I have to confess, the inclusion of such data would make this reviewer particularly enthusiastic about the work. If this is not feasible, I would recommend tempering the wording of "Golgi biogenesis" to a more conservative description, such as altered Golgi organization or duplication, and explicitly acknowledging the limitations of fluorescence-based analysis for this conclusion.

An additional conceptual point concerns the dual role of TbPLK in centrin arm regulation. TbPLK is known to promote centrin arm biogenesis through phosphorylation of TbCentrin2, yet in this study, TbPLK phosphorylation of KIN-G negatively regulates centrin arm assembly. This dual positive and negative regulatory role is intriguing but could be discussed more explicitly. The manuscript would benefit from a clearer conceptual framework addressing how phosphorylation of KIN-G might serve as a temporal or spatial switch to restrain KIN-G activity at specific stages of centrin arm assembly.

Finally, a schematic model summarizing the proposed regulatory pathway from TbPLK phosphorylation of KIN-G to centrin arm assembly, FAZ elongation, division plane placement, and Golgi organization would aid the reader.

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Reviewer #2 (Public review):

Summary:

The authors identify KIN-G as an in vitro substrate for phosphorylation by TbPLK and show that several of the in vitro P-ated sites, including T310, overlap with P-ation sites seen in live cells. The authors further show that PLK-mediated P-ation inhibits KIN-G binding to microtubules in vitro, as does a KIN-G-T301D mutant, and that expression of a KIN-G-T301D Phospho-mimic in *T. brucei* phenocopies KIN-G RNAi knockdowns, producing defects in cell division, morphogenesis of the centrion arm, FAZ and other cellular structures, as well as a misplaced cytokinesis furrow.

Understanding cytoskeletal rearrangements that drive cell division in *T. brucei* is an important and unresolved problem, so the work addresses important questions that are of great interest. PLK and KIN-G have previously been shown to be important for cell division and morphogenesis of cytoskeletal structures that drive cell division in *T. brucei*. The current work advances our understanding by suggesting a potential mechanism by which PLK and KIN-G might participate, namely through PLK-dependent P-ation to control KIN-G MT binding activity.

Strengths:

The authors use a rigorous combination of biochemistry, phosphoproteomics, cell biology, and mutant analysis to support their conclusion that PLK-mediated P-ation of KIN-G negatively regulates KIN-G microtubule binding, and this may explain the observation that a KIN-G T301 phosphomimic mutant blocks cell division and perturbs biogenesis of cytoskeletal structures that drive cell division and morphogenesis. Combining rigorous and informative in vitro studies with mutant analysis in live cells is a great strength. The work is solid and important, though a few pieces are needed to fully connect the in vitro findings with the in vivo observations, as detailed below.

Weaknesses:

Overall, I find this work to be solid and to provide an important addition to our understanding of mechanisms controlling cell division in *T. brucei*. The biochemistry, in particular, is rigorous and convincingly demonstrates PLK can P-ate KIN-G, altering its MT-binding ability. Analysis of phospho-mutants of KIN-G in live *T. brucei* supports the conclusion that P-ation of KIN-G at T301 negatively affects KIN-G function in vivo. I think, however, that the results fall short of supporting the title, because, although the data convincingly show that PLK can phosphorylate KIN-G at T301 in vitro, and that T301 is P-ated in vivo, they do not formally demonstrate (nor even test) whether PLK is the kinase responsible for this phosphorylation in vivo (experiments to address this seem quite feasible). I also do not see where the authors try to reconcile the absence of phenotype for KIN-G-T301A with the implied importance of KIN-G phosphorylation by PLK in cell division, which calls into question the need for P-ation of KIN-G-T301 in cell division. Suggestions for addressing these concerns are provided below.

My two main questions are:

(1) What is the biological relevance of KIN-G P-ation at T301?

a) The authors report no defect for the KIN-G-T301A mutant, so what then is the need for T301 P-ation, if the cell gets along fine without it? One step toward addressing this would be to ask what fraction of KIN-G shows P-ation at T301. Although published studies indicate P-ation at T301, it isn't known what percentage of KIN-G in the cell is P-ated. One might

anticipate, for example, that T301-P is a small minority of the population in asynchronous cultures and that T301 P-ation increases at specific cell cycle stages.

b) Published work links PLK to cell division, FAZ elongation, etc... The current work suggests that one role of PLK is to P-ate KIN-G at T301. In contrast, however, the current work also indicates that P-ation of KIN-G at T301 is unnecessary for normal cell division, FAZ elongation, etc....

c) Some experiments or at least commentary on points a and b above would strengthen the paper.

(2) Is PLK the kinase that P-ates Kin-G T301 in vivo?

a) The authors show PLK P-ates T301 (and other residues) in vitro, and that T-301 is P-ated in vivo. To bring the analysis full circle, it would be informative to examine KIN-G P-ation in a PLK mutant or upon inhibition of PLK with published inhibitors. This seems to be a very doable experiment with the tools available.

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Reviewer #3 (Public review):

Summary:

Here, the authors investigate the role of the *Trypanosoma brucei* polo-like kinase TbPLK in the function of flagellum-associated cellular structures in trypanosomes. They set out to test the hypothesis that a key substrate of TbPLK is the kinesin protein KIN-G, and that TbPLK phosphorylation of KIN-G regulates its functions in cells.

Strengths:

Using in vitro biochemistry with purified proteins, the authors convincingly demonstrate that TbPLK phosphorylates KIN-G at 29 sites. Moreover, they convincingly show that phosphorylation at one site, T301, impairs the binding of purified KIN-G to purified microtubules. Using immunofluorescence-based imaging approaches, they also show that TbPLK colocalizes with KIN-G at centrin arms during the early S-phase of the cell cycle. Centrin arms are structures that are located near the basal body and flagellum and are important for new flagellum biogenesis, Golgi positioning, and cell division. To evaluate the function of KIN-G phosphorylation in cells, they depleted KIN-G by RNAi, simultaneously expressed phospho-mimetic (T301D) and phospho-ablative mutant proteins, and used immunofluorescence to examine the impact on flagellum-associated cellular structures. They show that expression of the phospho-mimetic mutant KIN-G-T301D causes the following defects: reduced cell proliferation, disruption of centrin arm and Golgi biogenesis, impairment of FAZ elongation and flagellum positioning, and misplacement of the cell division plane. The data convincingly support the conclusion that KIN-G phosphorylation on T301 plays an important role in regulating the cellular functions of this kinesin motor protein.

Weaknesses:

Some of the broader conclusions are not directly supported by the data. For example, the title states "Polo-like kinase phosphorylation of the orphan kinesin KIN-G negatively regulates centrin arm biogenesis in *Trypanosoma brucei*," but the data do not directly address the specific role of TbPLK in phosphorylating KIN-G in cells. Moreover, some of the more specific conclusions in the paper, for example, that "phosphorylation of KIN-G" causes various cellular defects, are a bit of an overstatement. The supporting data rely on the expression of a phospho-mimetic mutant of KIN-G. Presumably, phosphorylation in cells is a normal part of

KIN-G regulation, and it is not just phosphorylation, but rather hyperphosphorylation that is being mimicked by the mutant. Some rewording of the specific conclusions is warranted, and the broader conclusion would be better supported with additional experimental evidence.

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