

Nek8445, a protein kinase required for microtubule regulation and cytokinesis in *Giardia lamblia*

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ABSTRACT *Giardia* has 198 Nek kinases whereas humans have only 11. *Giardia* has a complex microtubule cytoskeleton that includes eight flagella and several unique microtubule arrays that are utilized for parasite attachment and facilitation of rapid mitosis and cytokinesis. The need to regulate these structures may explain the parallel expansion of the number of Nek family kinases. Here we use live and fixed cell imaging to uncover the role of Nek8445 in regulating *Giardia* cell division. We demonstrate that Nek8445 localization is cell cycle regulated and this kinase has a role in regulating overall microtubule organization. Nek8445 depletion results in short flagella, aberrant ventral disk organization, loss of the funis, defective axoneme exit, and altered cell shape. The axoneme exit defect is specific to the caudal axonemes, which exit from the posterior of the cell, and this defect correlates with rounding of the cell posterior and loss of the funis. Our findings implicate a role for the funis in establishing *Giardia*'s cell shape and guiding axoneme docking. On a broader scale our results support the emerging view that Nek family kinases have a general role in regulating microtubule organization.

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INTRODUCTION

The protozoan parasite *Giardia lamblia* (synonymous with *G. intestinalis* and *G. duodenalis*) infects more than 200 million people each year (Lane and Lloyd, 2002; Lalle, 2010; Ansell *et al.*, 2015). The divergence of parasite protein kinases from their homologues in humans has been identified as an opportunity for the development of new therapeutic interventions (Rotella, 2012). The *Giardia* kinome (strain WB) contains 278 protein kinases, 80 of which constitute the core kinome from 49 different classes of kinases while the remainder are all Nek kinase homologues (Manning *et al.*, 2011). The highly expanded *Giardia* Nek family contains 198 kinases, compared with only 1 in budding yeast, 7 in *Arabidopsis*, 11 in humans, 13 in

Chlamydomonas, and 39 in *Tetrahymena* (Wloga *et al.*, 2006; Manning *et al.*, 2011; Takatani *et al.*, 2015).

Nek kinases are named for their homology to the *Aspergillus nidulans* NIMA (never in mitosis A) kinase. NIMA-related kinases (Nek or NRK) kinases have been shown to control mitotic entry, centrosome separation, and flagella length in other eukaryotes (Faragher and Fry, 2003; O'Connell *et al.*, 2003; Bradley and Quarmby, 2005; Wloga *et al.*, 2006; Chen and Gubbels, 2013), but may also have a more general role in regulating microtubule organization (Takatani *et al.*, 2015). The number of Nek kinases is typically expanded in the genomes of eukaryotes that build cilia or flagella (Parker *et al.*, 2007). Notably, *Giardia* possesses four pairs of flagella that undergo a complex developmental cycle (Nohynkova *et al.*, 2006; see Figure 1 for a diagram of *Giardia* structures). As pairs of flagella do not share paired basal bodies, maintaining the identity of each flagella is complex and the regulatory mechanism remains uncharacterized (Nohynkova *et al.*, 2006). The complexity of maintaining eight flagella identities could explain the development of a greatly expanded set of Nek kinases in *Giardia*.

Flagella positioning and identity are essential for *Giardia* cellular processes including pathogenicity. Unlike most other eukaryotes that dock their basal bodies just under the plasma membrane,

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Abbreviations used: BKI, bumped kinase inhibitor; DAPI, 4',6-diamidino-2-phenylindole; 3HA, triple-hemagglutinin tag; mNG, mNeonGreen; NEK or NRK, NIMA-related kinase; NIMA, never in mitosis A.

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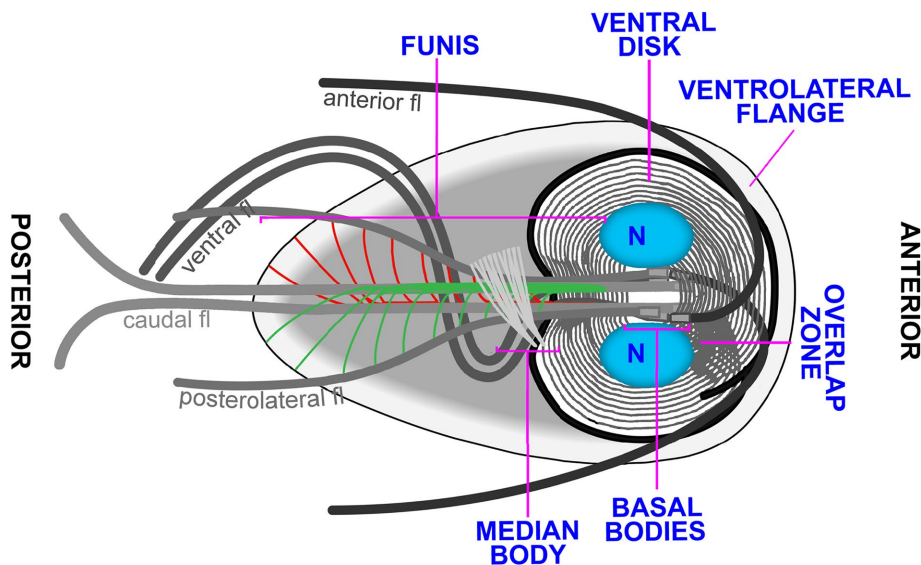


FIGURE 1: Diagram of *Giardia* emphasizing microtubule-based structures. The microtubule cytoskeleton includes the ventral disk, median body, funis, anterior, posterolateral, ventral, and caudal flagella sets.

Giardia positions its eight basal bodies between the two nuclei in the center of the cell. As a result, a significant portion of each axoneme runs through the cytoplasm of the cell before it exits and is sheathed by membrane (McInally and Dawson, 2016; McInally et al., 2019b). We use the term flagella to refer to the membrane-bound portion of axonemes, while the portion of axonemes running through the cell body is referred to as cytoplasmic axonemes. *Giardia* cytoplasmic axonemes serve as tracks for cargo trafficking and have also been implicated in force generation for cell division (Hardin et al., 2017). Each of *Giardia*'s flagella pairs beat with unique waveforms, further indicating the complexity of specifying flagella identity and function (Lenaghan et al., 2011). For example, the caudal axoneme pair, which runs the length of the cell body, is thought to flex the entire body of the cell to contribute to swimming via a flipper type mechanism (Lenaghan et al., 2011). The anterior and posterolateral flagella generate power strokes for motility, while the ventral flagella, with a unique fin-like appendage, beat in a sinusoidal waveform thought to generate fluid flows beneath and around *Giardia* trophozoites (Holberton, 1973; Lenaghan et al., 2011; Halpern et al., 2017).

In addition to four unique sets of flagella, *Giardia* has several novel microtubule structures including the ventral adhesive disk, median body, and funis (also known as the caudal complex). With the exception of the median body, each of these structures is associated with specific sets of basal bodies (McInally and Dawson, 2016). The ventral disk is a complex dome-shaped structure composed of approximately 100 microtubules in a one and a quarter turn helical sheet and is thought to mediate attachment through a suction cup-like mechanism (Erlandsen et al., 2004; Hansen et al., 2006; Brown et al., 2016; Nosala et al., 2018). The funis is a pair of microtubule sheets that originate at the basal bodies. One sheet runs above the cytoplasmic caudal axoneme pair while the other runs below the caudal cytoplasmic axonemes. Toward the posterior of the cell, the funis microtubules separate from the main sheet to form rib-like structures that lock the cytoplasmic axonemes of the caudal and posterolateral flagella together (Benchimol et al., 2004; Carvalho and Monteiro-Leal, 2004). The median body is a reservoir of micro-

tubules and associated binding proteins used to support *Giardia*'s rapid mitosis (Hardin et al., 2017; Horlock-Roberts et al., 2017). Remarkably, mitosis is completed in approximately 7 min and this includes duplicating axonemes and building two new ventral disks (Hardin et al., 2017). Currently, little is known about how the assembly and disassembly dynamics of these structures are regulated. As *Giardia* lacks an anaphase promoting complex, it has been suggested that kinases may play an enhanced role in regulating mitotic events (Gourguechon et al., 2013). Indeed, Nek kinases have been implicated in the regulated assembly and disassembly of microtubule-based structures during the cell cycle (Smith et al., 2012).

The *Giardia* Nek kinase homologues with the highest sequence identity for human Nek1 and Nek2 were previously studied (Smith et al., 2012). The Nek1 homologue *G/Nek1* localized to the ventral disk while the Nek2 homologue *G/Nek2* localized to the cytoplasmic axonemes of the anterior flagella. Both kinases reorganize during mitosis, and overexpression studies showed that *G/Nek1* has a role in ventral disk disassembly while *G/Nek2* overexpression interfered with excystation (hatching from the cyst form). Other *Giardia* Nek kinases have not yet been individually studied.

Here, we characterize the role of Nek kinase GL50803_8445 (Nek8445) in *Giardia* cell proliferation. Nek8445 has direct orthologues (>90% sequence identity) in other *Giardia* assemblages (strains), but exhibits only 30–40% sequence identity to other Nek paralogues in the expanded *Giardia* Nek kinome. This is consistent with a functional role in *Giardia* that is both essential and distinct from that of other Nek family kinases. We previously identified this kinase as a potential therapeutic target due to the presence of a small gatekeeper residue, which makes it susceptible to a specific class of kinase inhibitors with relatively low activity against human kinases (Hennessey et al., 2016). These “bumped kinase inhibitors” (BKIs) exploit the property that most mammalian protein kinases have a large gatekeeper residue that limits accessibility to a hydrophobic pocket in the active site. Various BKIs have been shown to selectively and effectively block kinase-associated processes in the life cycle of parasites such as *Toxoplasma gondii*, *Cryptosporidia parvum*, and *Plasmodium falciparum* (Keyloun et al., 2014). In *Giardia* there are at least eight small gatekeeper kinases (Hennessey et al., 2016). Although we identified inhibitors that target Nek8445, they also target additional small gatekeeper kinases, which currently limits their usefulness in uncovering the role of Nek8445.

We previously determined that Nek8445-depleted cells are severely impeded in growth, often abnormally multinucleated, and unable to maintain attachment (Hennessey et al., 2016). The initial study focused on defects at 48 h after morpholino knockdown, which corresponded with the endpoint of our growth assays. By this time cellular organization was severely disrupted and therefore the primary defect was not apparent. However, the cytokinesis and attachment defects are potentially explained by aberrant flagella function. Unlike other eukaryotes that use an actomyosin contractile ring to guide membrane delivery and drive cytokinesis, *Giardia* utilizes nascent cytoplasmic axonemes to coordinate

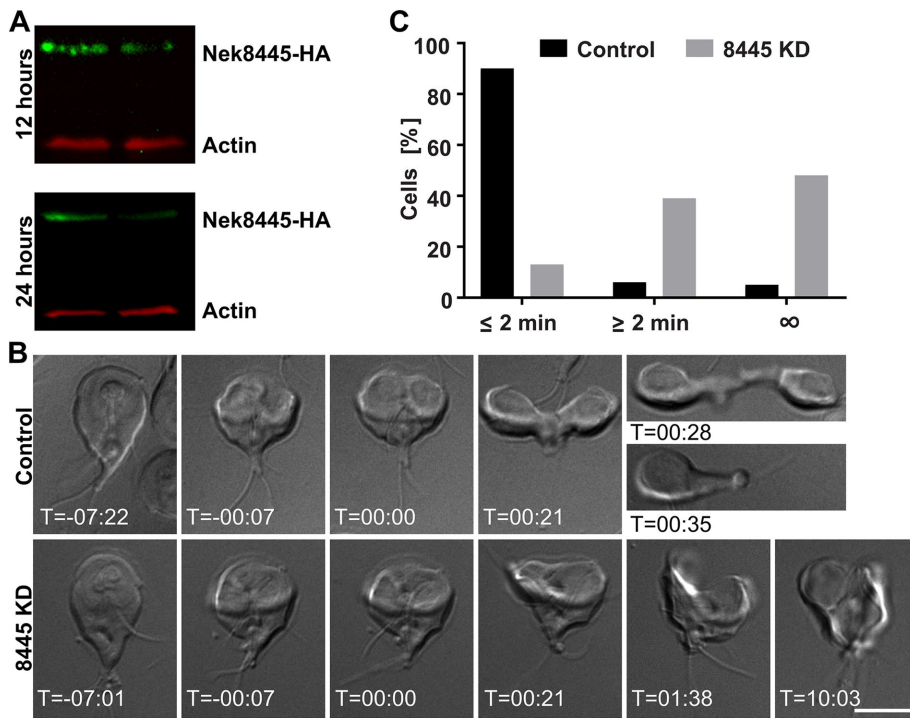


FIGURE 2: Nek8445 is required for cytokinesis. (A) Multiplex immunoblot showing Nek8445-HA depletion 12 and 24 h after Nek8445 antisense and control morpholino treatment. (B) Representative still images from a 4D time-lapse movie showing failed cytokinesis in Nek8445-depleted cells (see Supplemental Video S1). $T = 0$ set to the initiation of cytokinesis, which occurs ~ 7 min after the initiation of mitosis as indicated by anterior flagella reorganization. (C) Histogram of cytokinesis timing. Only 13% of Nek8445-depleted cells complete cytokinesis within 2 min vs. 90% for the control treatment ($n = 88$ control morpholino, $n = 56$ Nek8445 KD). Scale bar = 5 μm .

membrane trafficking into the furrow while the mature flagella propel daughter cells apart for abscission (Hardin *et al.*, 2017). The flagella may also have a role in parasite attachment (Holberton, 1973). It has been proposed that flows generated by the ventral flagella transit through gates in the ventral disk, which can close off to lock down attachment (Dawson and Nosala, 2017). Here we examine the role of Nek8445 in *Giardia*'s unique mode of cytokinesis.

RESULTS

Nek8445 function is required for cytokinesis

The initial observation that Nek8445 plays a role in cytokinesis was based on analysis of Nek8445-depleted cells fixed and analyzed 48 h after treatment with translation-blocking antisense morpholinos (Hennessey *et al.*, 2016). These cells were abnormally multinucleate (more than two nuclei) with a roughly corresponding increase in flagella number (Supplemental Figure S1). Cell shape and flagella organization were disrupted in the multinucleate cells. Therefore, it was not clear whether the cytokinesis defect was a primary consequence of Nek8445 knockdown or whether disrupted cell polarity led to failed cytokinesis. We recently developed methods for long-term four-dimensional (4D) differential interference contrast (DIC) time-lapse imaging to follow *Giardia* trophozoites through the cell cycle (Hardin *et al.*, 2017). We sought to film Nek8445-depleted cells undergoing cytokinesis to test for changes in the timing of events and gain insight into the underlying cause of failed cytokinesis. We planned to examine Nek8445-depleted cells from 20 to 26 h after morpholino treatment as in our previous study of *Giardia*

cytokinesis. However, the cells were poorly adherent and the majority of the attached cells had already failed cytokinesis or were abnormally organized (Supplemental Figure S2). Therefore, we decided to film cells from 12 to 18 h after knockdown. The majority of cells at this time point displayed typical organization as assessed by DIC imaging (Supplemental Figure S2), allowing us to follow cytokinesis in long-term 4D imaging experiments. Supporting significant depletion of Nek8445 at this time point, Western blotting indicated that antisense morpholino treatment reduced Nek8445 levels by 41% compared with cells treated with GeneTools LLC's standard control morpholino at 12 h post electroporation (~ 1.5 cell cycles after treatment; Figure 2A).

Upon the initiation of cytokinesis, the cleavage furrow often failed to progress (Figure 2B and Supplemental Video S1). Under our assay conditions, 90% of control morpholino-treated cells completed cytokinesis within 2 min ($n = 88$), similar to our previous results (Hardin *et al.*, 2017). In contrast, only 13% of Nek8445-depleted cells ($n = 56$) completed cytokinesis within 2 min. Additionally, 48% of Nek8445-depleted cells never completed cytokinesis compared with just 4.5% noncompletion observed for cells treated with the control morpholino (Figure 2C). Although Nek kinases are implicated in centrosome duplication and regulation of mitosis, we did not

observe any mitotic defects as indicated by typical nuclear segregation in Nek8445-depleted cells (Supplemental Video S1). Remarkably, the cell division defect resulting from Nek8445 knockdown is stronger than the phenotype of any cytoskeleton or membrane trafficking component previously studied (Hardin *et al.*, 2017), suggesting a central regulatory role for Nek8445 in coordinating cytokinesis.

Nek8445 localization is cell cycle regulated

We previously reported that Nek8445-3HA localized to the plasma membrane and nuclear envelope in interphase cells (Hennessey *et al.*, 2016). Many cell cycle-regulated proteins, including Nek kinases, change their localization during mitosis (Mahjoub *et al.*, 2004; Smith *et al.*, 2012; Chen and Gubbels, 2013). To follow Nek8445 localization throughout the cell cycle in live cells, the C-terminus of Nek8445 was tagged with the green fluorescent protein, mNeonGreen (Shaner *et al.*, 2013), generating Nek8445-mNeonGreen (Nek8445-mNG). This construct was integrated into the genome to ensure endogenous expression levels (Gourguechon and Cande, 2011). At the initiation of mitosis, Nek8445-mNG translocated from the cell cortex to become concentrated around the two nuclei and along the cytoplasmic axonemes (Figure 3A and Supplemental Video S2). Late in mitosis, Nek8445-mNG also concentrate at what appear to be the tips of the cytoplasmic axonemes as the axonemes were being rearranged (Supplemental Video S2). During cytokinesis, Nek8445-mNG remained in association with the nuclear envelopes as well as portions of the anterior, caudal, and posterolateral cytoplasmic axonemes.

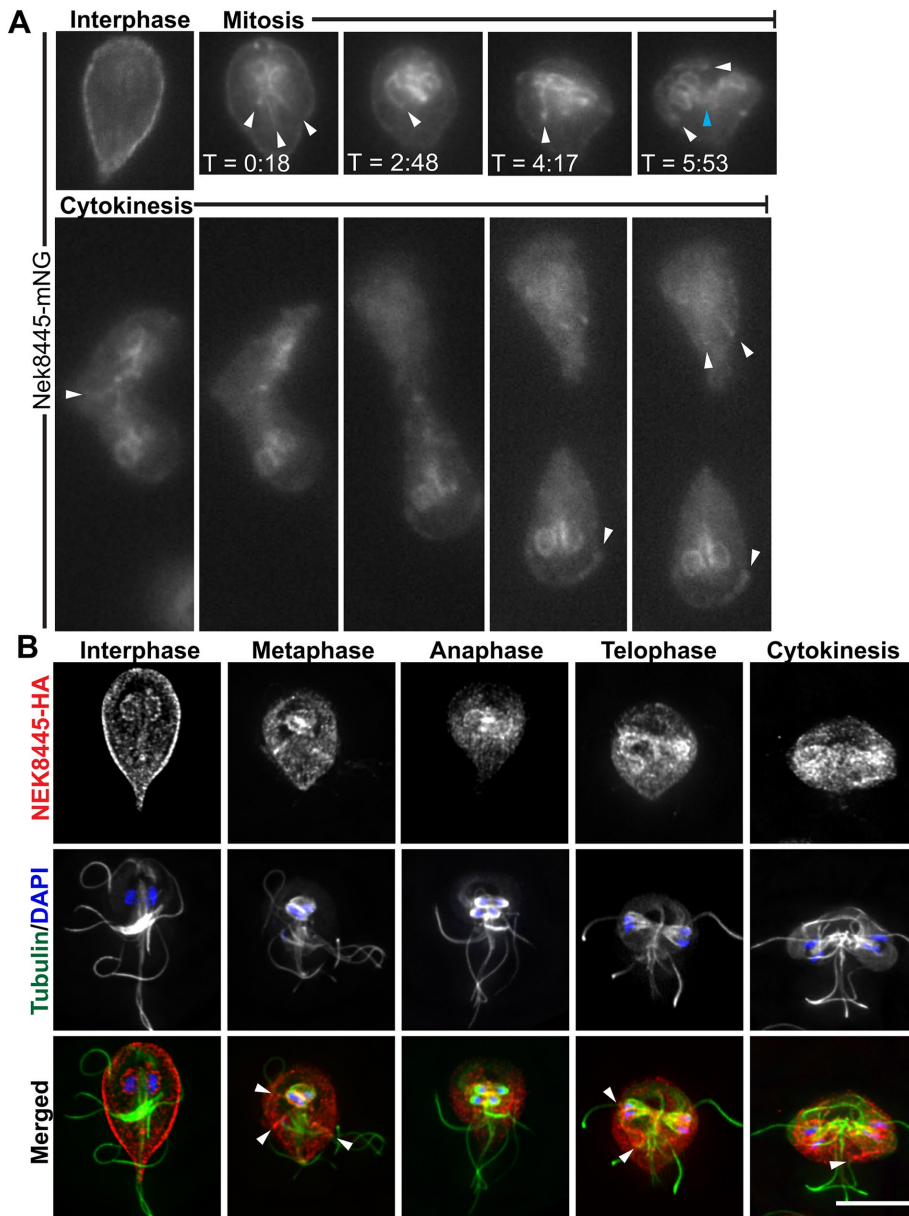


FIGURE 3: Nek8445 relocates during mitosis to associate with axoneme microtubules. (A) Still images from a time-lapse movie of Nek8445-mNG showing Nek8445 translocating from the cell cortex to the nuclear envelope, and axoneme microtubules (see Supplemental Video S2). White arrowheads indicate Nek8445 in association with cytoplasmic axonemes; cyan arrowhead marks nascent axonemes pointing into the developing furrow. (B) Immunofluorescence analysis of Nek8445-HA (red), microtubules (6-11B-1 antibody; green), and nuclei (DAPI; blue) throughout the cell cycle. Scale bar = 5 μ m.

To confirm association of Nek8445 with the cytoplasmic axonemes of flagella during mitosis and cytokinesis, we colocalized Nek8445-3HA and microtubules in fixed cells (Figure 3B). Indeed, Nek8445-3HA colocalized with the cytoplasmic axonemes of flagella including some association with the nascent axonemes that act as tracks for delivering membrane to the cleavage furrow (Hardin et al., 2017). Overall the two chimeric proteins have similar localization; however, Nek8445-3HA clearly associates with the nuclear envelope of interphase cells while Nek8445-mNG only weakly associated with the nuclei until the initiation of mitosis.

our measurements because they were too convoluted to trace. Similarly, we did not attempt to measure the posterolateral or ventral axonemes as they were often difficult to identify, due to being mispositioned, in the Nek8445 knockdown cells.

We also noted that the rounded cells lacked median bodies (Figure 4A). Upon quantitation, we found that 46.8% ($n = 158$) of all Nek8445-depleted cells lacked median bodies versus just under 15% ($n = 302$) of the control morpholino-treated cells. Note that this microtubule reservoir is consumed during mitosis and then grows in size as cells prepare for the next round of division (Hardin et al., 2017; Horlock-Roberts et al., 2017). Additionally, we identified

We previously established that Rab11-positive vesicles traffic along cytoplasmic axonemes and into the cleavage furrow to support furrow progression (Hardin et al., 2017). The localization of Nek8445 to the cytoplasmic axonemes prompted us to ask whether Nek8445 and Rab11 might have a functional relationship. Colocalization of HA-Rab11 with Nek8445-mNG revealed that while both proteins translocate from the plasma membrane to the intracytoplasmic axonemes during mitosis and cytokinesis, they were rarely seen to be on cytoplasmic axonemes at the same time (Supplemental Figure S3). Therefore, it is unlikely that Nek8445 is directly regulating membrane trafficking associated with furrowing. Instead, Nek8445 may have a role in regulating axoneme/flagella function, which is required for force generation during cytokinesis (House et al., 2011; Hardin et al., 2017).

Nek8445-depleted cells have short flagella and lack median bodies

The microtubule cytoskeleton plays a major role in establishing *Giardia*'s unique shape and in generating forces for driving cells apart during cytokinesis (Mariane et al., 2005; Tumova et al., 2007; Hardin et al., 2017). Defects in microtubule regulation could account for defects in cell division (Mariane et al., 2005). Therefore, we examined tubulin staining in Nek8445-depleted cells (Figure 4A). We observed that ~17% ($n > 300$) of the Nek8445-depleted cells had noticeably short flagella and either round or oval-shaped cell bodies (Figure 4, A and B).

We measured the entire length of anterior and caudal axonemes from randomly selected control and Nek8445 translation-blocking morpholino-treated cells. The anterior axonemes were modestly shorter (control morpholino: $17.5 \pm 0.34 \mu\text{m}$, $n = 76$; Nek8445 KD $15.5 \pm 0.48 \mu\text{m}$, $n = 60$) while the caudal axonemes were 30% shorter (control morpholino: $17.7 \pm 0.27 \mu\text{m}$, $n = 80$; Nek8445 KD: $12.3 \pm 0.31 \mu\text{m}$, $n = 64$). The values reported here likely underreport the shortening defect as the knockdown cells with the most severe flagella shortening had tangled axonemes, which we omitted from

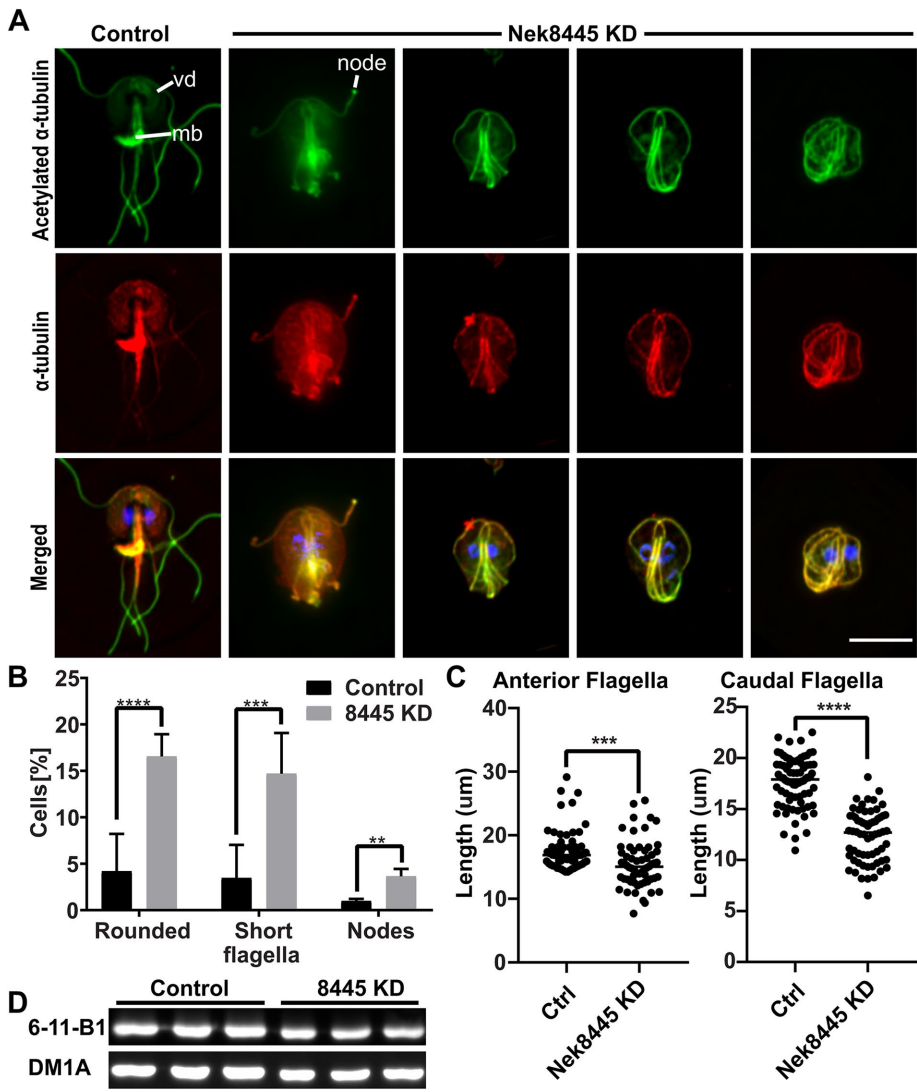


FIGURE 4: Nek8445 regulates microtubule organization. (A) Immunofluorescence staining of acetylated and total α -tubulin. Acetylated tubulin is stained with the anti-acetylated α -tubulin antibody 6-11B-1 (green) and total tubulin is stained with the anti- α -tubulin antibody DM1A (red). Note changes in overall tubulin staining after Nek8445 depletion including reduced ventral disk (vd) staining, the lack of a median body (mb) in rounded cells, and nodes of tubulin staining at the ends of flagella. Scale bar = 5 μm . (B) Quantification of mutant phenotypes observed after Nek8445 depletion ($n > 300$ cells). (C) Measurement of anterior and caudal axoneme length from control morpholino and Nek8445 knockdown cells ($n = 76$ measurements from control cells and $n = 64$ measurements for 8445 knockdown cells). (D) Immunoblotting for total tubulin (DM1A) vs. acetylated tubulin (6-11B-1 antibody) in Nek8445 and control morpholino-treated cells. Lysates were prepared from equal numbers of cells for each replicate. There is no significant difference in the ratio of acetylated to total tubulin although overall tubulin levels are decreased by ~25% in Nek8445-depleted cells.

abnormal accumulation of tubulin in nodes at the tips of flagella in the Nek8445 knockdown cell (Figure 4, A and B). Short flagella and accumulation of axoneme components at the tips of flagella is consistent with a defect in axoneme/flagella length homeostasis (Qin et al., 2004; Absalon et al., 2008; Meng and Pan, 2016).

Microtubule staining also appears altered in Nek8445 knockdown cells. Specifically, we noticed that staining of the ventral disk was often reduced while the axonemes of Nek8445-depleted cells appeared more clearly delineated with less background staining. Our initial immunofluorescence localization was performed with the

anti-acetylated α -tubulin antibody 6-11B-1, which is known to recognize K40 acetylation and stain all known microtubule-based structures in *Giardia* (Nohynkova et al., 2006; Halpern et al., 2017). The altered staining indicates that either acetylation levels or microtubule array organization were perturbed by Nek8445 depletion. Microtubule acetylation is thought to soften microtubules so that they are more flexible and able to resist mechanical stress (reviewed in Janke and Montagnac, 2017). Notably, human Nek3 has been shown to regulate the levels of acetylated tubulin (Chang et al., 2009). Therefore, we questioned whether Nek8445 has a role in regulating microtubule acetylation as this could potentially account for the observed defects in axoneme length and ventral disk staining.

To differentiate between changes in acetylation or overall microtubule organization, we costained cells with the anti-acetylated α -tubulin antibody 6-11B-1 and the anti- α -tubulin antibody DM1A. Total α -tubulin and acetylated tubulin localization appeared qualitatively similar in Nek8445 knockdown cells (Figure 4A). Immunoblotting with the two tubulin antibodies indicated that there is not a significant difference in the ratio of acetylated to nonacetylated α -tubulin when comparing control morpholino and Nek8445 knockdown cells (Figure 4D). This confirmed that Nek8445 does not regulate acetylation. However, consistent with overall smaller cells that lack median bodies, we found that tubulin levels are approximately 25% lower in Nek8445-depleted cells compared with control morpholino-treated cells.

Nek8445-depleted cells have defective ventral disk organization

We previously reported that Nek8445-depleted cells were defective in their ability to attach to the sidewalls of culture tubes (Hennessey et al., 2016). Attachment is largely attributed to the microtubule-based suction cup-like organelle known as the ventral adhesive disk (Dawson and Nosala, 2017). Because the ventral disk failed to stain with anti-tubulin antibodies in many of the Nek8445-depleted cells, we examined localization of the ventral disk marker δ -giardin to verify that a ventral disk was still present (Jenkins et al., 2009; Halpern et al., 2017). We did not find any cells lacking δ -giardin-3HA staining, indicating that the ventral disk was present (Figure 5). However, we observed abnormal disk organization that correlated with abnormal cell shape and flagella length. Of the Nek8445-depleted cells that consequently had short flagella, we observed 38% ($n = 108$) of these cells to have an altered disk morphology compared with the control morpholino-treated cells ($n = 63$). Defects included a larger area devoid of δ -giardin staining in the center of the disk (bare area),

frayed, and open disk conformations (Figure 5). Similar defects have been observed in cells depleted of disk-associated proteins that have a structural role in the ventral disk (McInally *et al.*, 2019a); therefore, Nek8445 may have a role in regulating ventral disk assembly.

Nek8445-depleted cells have axoneme exit defects

In contrast to the typical half-pear shape of trophozoites, a subset of Nek8445-depleted cells were observed to have a rounded cell shape that lacked a portion of the posterior end of the cell. Tubulin staining (6-11B-1) indicated that the caudal flagella of these cells were either tucked under the cell body or the axonemes were trapped within the cell body, but this could not be resolved with light microscopy (Figures 4 and 5). To resolve whether the axonemes were trapped in the trophozoite cell body we turned to scanning electron microscopy (SEM). Control morpholino ($n = 26$) and translation-blocking anti-Nek8445-treated cells ($n = 50$) were processed for SEM 12 h after morpholino electroporation. As observed by light microscopy, a subset of the Nek8445-depleted cells was much smaller and had short flagella (Figure 6). While the anterior, ventral, and posterolateral flagella were visible, the caudal flagella were frequently missing. In some cases, we could see abnormal membrane protrusions near where the caudal flagella were expected to exit, with successful exit from an atypical position (Figure 6, arrowhead). The defect in axoneme exit site specification is expected to contribute to the cytokinesis defects observed with Nek8445 depletion as the flagella must be properly oriented in order to collaboratively generate forces for cytokinesis (Mariane *et al.*, 2005; Paredez *et al.*, 2011; Hardin *et al.*, 2017).

The defective axoneme exit defect provided us with the opportunity to investigate whether the exit site is predetermined or established after the axonemes are in position. Unlike other eukaryotes where basal body appendages known as transition fibers dock the basal body to the plasma membrane (Goncalves and Pelletier, 2017), *Giardia* basal bodies are positioned many microns from where the cytoplasmic axoneme exits the cell body to become sheathed by plasma membrane. *Giardia* lacks all known transition zone proteins, so how the axoneme exit site is specified remains an open question (Avidor-Reiss and Leroux, 2015; Barker *et al.*, 2014). A recent study of *Giardia* intraflagellar transport (IFT) suggests the exit sites are pore-like diffusion barriers that regulate IFT (McInally *et al.*, 2019b). The Hehl lab, University of Zurich, identified GL50803_8855 as a protein that forms a ring at the axoneme exit site, where the axonemes transition from cytoplasmic to membrane sheathed (GiardiaDB). We endogenously tagged exit site protein GL50803_8855 (ESP8855) with a triple HA tag to test whether the exit site was altered in the Nek8445 knockdown cells. As previously reported, we observed distinct localization of ESP8855-HA to the exit site of each axoneme set in cells treated with the control morpholino ($n = 23$; Figure 7A). In Nek8445 knockdown cells, ESP8855 localization varied. In cells where the caudal axoneme exited the cell body, ESP8855-HA was present where axonemes met the plasma membrane (Figure 7B). This included several cells with abnormally oriented caudal cytoplasmic axonemes (6 of 58) that mispositioned their caudal exit sites (Figure 7C). However, ESP8855-HA did not mark the posterior exit site in 59% ($n = 58$) of rounded cells with short caudal axonemes that in some cases were too short to reach the cell cortex (Figure 7D). Together we interpret these findings to indicate that the exit site is specified upon axoneme contact with the plasma membrane.

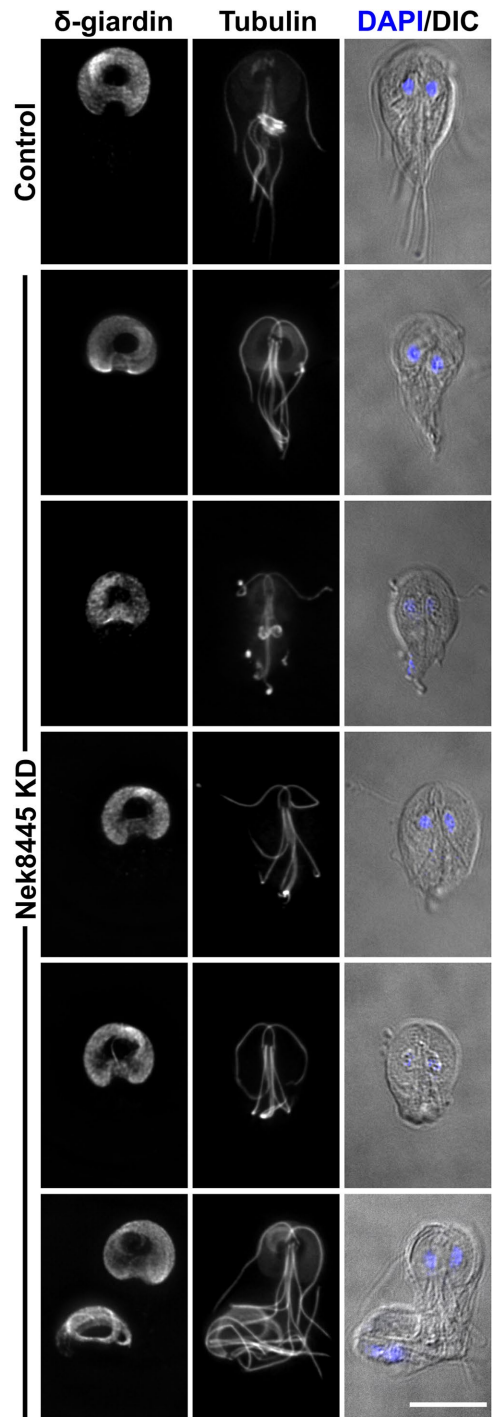


FIGURE 5: Nek8445-depleted cells have ventral disk organization defects. Control morpholino-treated cells and Nek8445 knockdown cells stained for δ -giardin-3HA and tubulin (6-11B-1 antibody). Note the altered disk morphology such as increased width, larger bare area, and some striated staining of δ -giardin-3HA not observed in control cells. Of rounded Nek8445-depleted cells, 38% ($n = 108$) had altered disk morphology not observed in the control morpholino-treated cells ($n = 63$). Scale bar = 5 μ m.

Superresolution imaging reveals Nek8445-depleted cells lack a funis

Nek8445 depletion alters the lengths of all flagella, but the caudal axonemes were specifically susceptible to being trapped in the cell

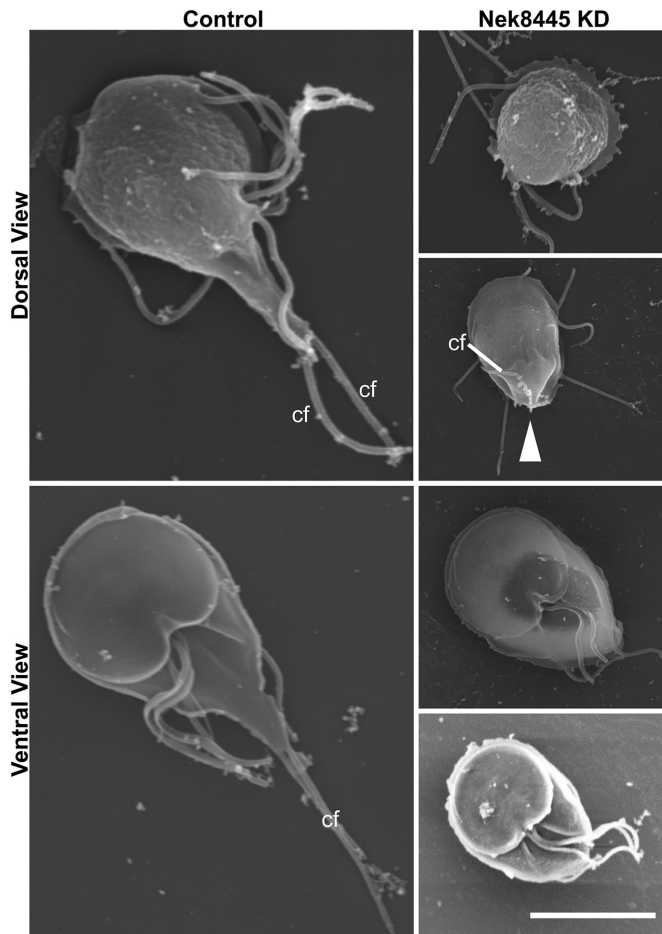


FIGURE 6: SEM analysis reveals Nek8445-depleted cells have caudal axoneme exit defects. Control morpholino and Nek8445 knockdown cells processed for SEM 12 h after morpholino treatment. Note the shorter cell length and absence of caudal flagella (cf) at the anterior apex of the cell. White arrowhead indicates an abnormal membrane protrusion associated with a caudal flagellum that eventually exits the cell body at an abnormal position. Scale bar = 5 μ m.

body without an exit site. Many of the trapped axonemes are long enough to exit the cell body; therefore, the failure to exit cannot simply be attributed to length. The caudal cytoplasmic axonemes make the longest run through the cell body and are uniquely sandwiched between the two halves of the funis (see Figure 1). Because the cytoplasmic caudal axonemes often stained with higher contrast than in controls, and the exit sites were sometimes mispositioned (Figures 4–7), we questioned whether the funis was lost or disrupted as a consequence of Nek8445 depletion. The funis is difficult to observe with standard resolution microscopy because it has overlapping localization with the caudal cytoplasmic axonemes and spacing of the rib-like microtubules is near the resolution limit of standard light microscopy. We turned to structured illumination super-resolution microscopy to compare control and Nek8445-depleted cells. We were able to resolve the funis in every control morpholino-treated cell examined ($n = 12$) while none of the rounded Nek8445 knockdown cells ($n = 12$) had any vestige of a funis (Figure 8). The lack of a funis is consistent with the abnormal posterolateral axoneme/flagella positioning observed in Nek8445 knockdown cells. Previously observed connections between the funis microtubules and the posterolateral cytoplasmic axonemes were anticipated to be important for caudal cell movement (Benchimol

et al., 2004); however, the funis also appears to have a role in stabilizing the position of the posterolateral axonemes. Another unique feature of the rounded cells is that we observed several cells with frayed axoneme tips, further suggesting an axoneme stability defect in Nek8445-depleted cells (Wloga et al., 2006; Bower et al., 2013; Meng and Pan, 2016). The correlation of round cell shapes, mispositioned axonemes/flagella, defective axoneme exit, and the lack of a funis is consistent with the funis having a role in establishing *Giardia* cell shape and guiding caudal axoneme exit. Together this study illustrates the overall importance of Nek8445 in regulating the microtubule cytoskeleton of *Giardia*.

DISCUSSION

We set out to study the primary cause of cytokinesis defects in Nek8445-depleted cells. Recent advances in live cell imaging have made it possible to visualize *Giardia* mitosis and cytokinesis with fantastic spatial-temporal resolution providing insights into *Giardia*'s myosin-independent mechanism of cytokinesis (Hardin et al., 2017). Characterization of the Nek8445 knockdown phenotypes illustrates a requirement for Nek8445 in coordinating microtubule organization. This study reinforces the view that the microtubule cytoskeleton has expanded function for *Giardia* cell division, which includes force generation necessary for cytokinesis (Benchimol, 2004; Correa and Benchimol, 2006; Hardin et al., 2017).

Our working model for myosin-independent cytokinesis (Hardin et al., 2017) posits that at the onset of cytokinesis, the caudal cytoplasmic axonemes, each connecting to a daughter ventral disk, orients the daughter disks so that the anteriormost regions face away from one another. Major reorganization of the flagella occur as the cells initiate cytokinesis such that each daughter cell will inherit four full-length flagella and grow four new flagella (Nohynkova et al., 2006; Sagolla et al., 2006; Tumova et al., 2007; Hardin et al., 2017). As the furrow expands, the newly duplicated axonemes, which are completely contained within the cytoplasm of the dividing cell, are used as tracks to guide additional membrane into the cleavage furrow (Hardin et al., 2017). Ultimately, the four mature flagella assume the position and identities of the anterior and caudal flagella (Nohynkova et al., 2006; Hardin et al., 2017). Power strokes from the anterior flagella drive the two daughter cells in opposite directions generating the membrane tension required for division (Zhang and Robinson, 2005; Ralston et al., 2006; Poirier et al., 2012; Hardin et al., 2017).

The observation that Nek8445 knockdown cells with typical cell shape and organization fail cytokinesis (Figure 2), suggests that the flagella may have improper function before the extensive shortening observed in rounded cells. Cytokinesis in these cells appears to fail because the caudal axonemes are not able to orient the daughter disks in opposition to each other (Figure 2 and Supplemental Video S1). This reorientation requires both coordinated force generation and the delivery of plasma membrane to provide slack for the disks to move away from each other (Hardin et al., 2017). The similarity between the failed cytokinesis phenotype of brefeldin A treatment and Nek8445-depleted cells is striking (Hardin et al., 2017), which leads us to suspect that membrane trafficking could be impaired.

While most 12-h post knockdown cells had typical cell shape and organization as assessed with DIC optics, tubulin staining revealed that a significant portion of cells (~45%) lack median bodies. The lack of median bodies is an indicator of reduced tubulin reservoirs. During mitosis, microtubules flux from the median body, to the spindle, nascent disks, and finally to the nascent axonemes (Nohynkova et al., 2006; Hardin et al., 2017). Growth of new axonemes could be

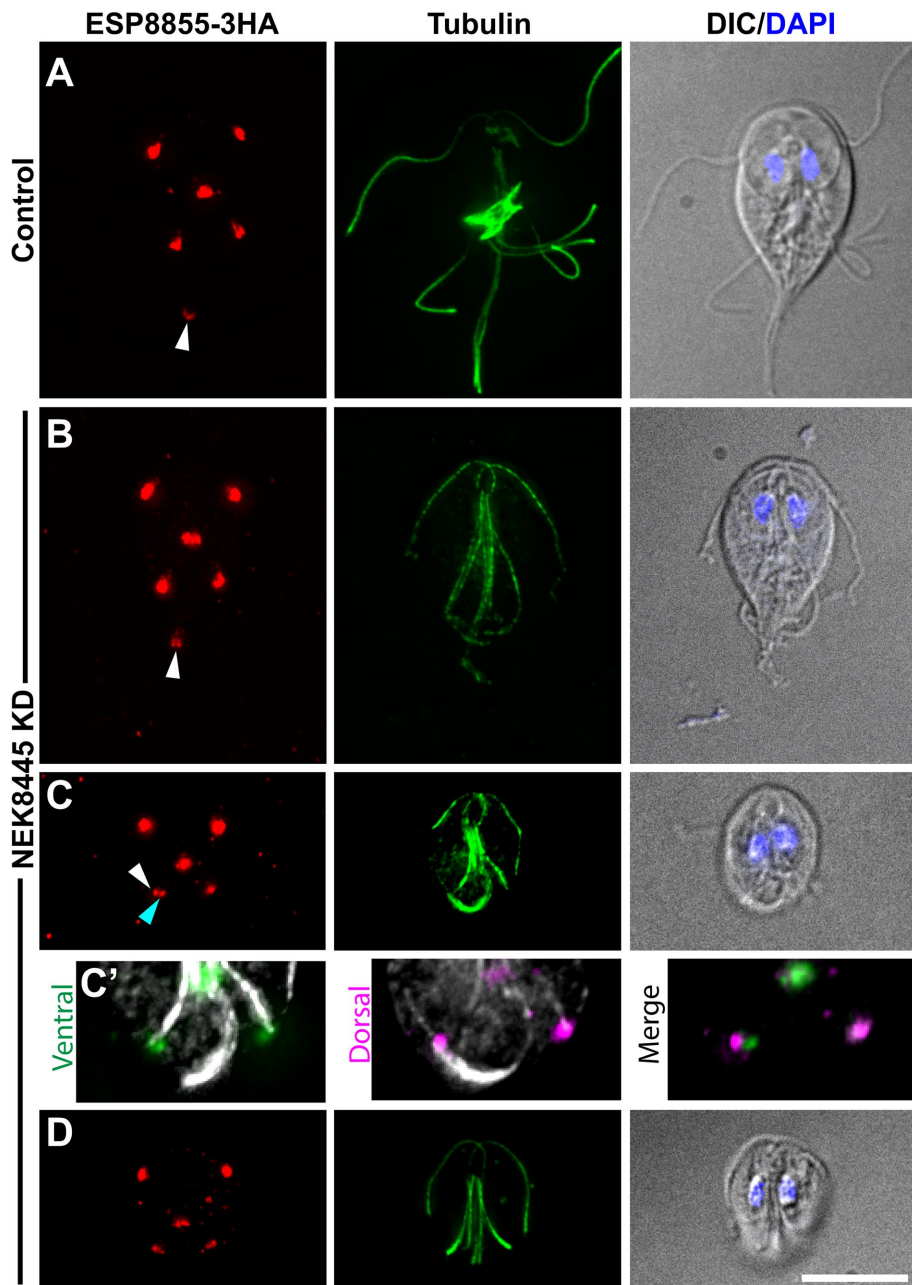


FIGURE 7: Nek8445-depleted cells misposition or lack exit sites. Control morpholino and Nek8445 knockdown cells were stained for tubulin (6-11B-1 antibody; green) and ESP8855-3HA (red). White arrowheads mark the caudal axoneme exit site and are oriented with the trajectory of the exiting axoneme. (A) Typical exit site organization in control cells. (B) Knockdown cell with typical teardrop shape retains exit sites at the expected position. (C) Mispositioned caudal axoneme exit site observed in 10% of rounded knockdown cells ($n = 58$). Note that these images are maximum projections; the caudal axoneme exit site in this cell is clearly positioned above the adjacent posterolateral axoneme exit site (cyan arrowhead) when viewed in the 3D image stack. (C') Shows maximum projections of the ventral and dorsal halves of the Z-stack to show that these exit sites are displaced in Z similar to the cell with the arrowhead in Figure 6. (D) Complete lack of a caudal axoneme exit site is observed in 59% of rounded knockdown cells ($n = 58$). Scale bar = 5 μm .

impaired due to a limited tubulin supply. Importantly, the nascent axonemes point into the cleavage furrow during cytokinesis and are used to direct new membrane into the furrow. Thus, short or improperly organized nascent axonemes could potentially reduce the availability

of membrane for furrowing. Due to the lack of robust cell synchronization and the very short window during which cells use these tracks to deliver membrane, it is technically challenging to find knockdown cells actively dividing in order to examine the positioning and length of the nascent axonemes.

Short flagella may indicate a defect in IFT. Such a role for Nek8445 would not be surprising as Nek family kinases are known to regulate IFT (Lin *et al.*, 2003; Surpili *et al.*, 2003; Wloga *et al.*, 2006; Meng and Pan, 2016). We did not measure axoneme or flagella growth rates, but the presence of short flagella with swollen tips is consistent with an IFT defect. Indeed, the short flagella phenotype observed in Nek8445-depleted cells is reminiscent of the *Giardia* kinesin-2 mutants that impair IFT (Hoeng *et al.*, 2008; Carpenter and Cande, 2009).

The caudal axoneme exit defect provided a unique opportunity to probe the mechanism of exit site specification/flagella pore formation in *Giardia*. ESP8855 marks axoneme exit sites. When Nek8445 was depleted, cells with extremely short cytoplasmic axonemes were observed to lack ESP8855 staining at the posterior apex of the cell, which is where the caudal axonemes exit in wild-type interphase cells. Additionally, some cells had mispositioned posterolateral and caudal axonemes with improperly positioned exit sites (Figures 6 and 7C). These findings are consistent with axoneme contact with the plasma membrane being the specifier for exit site/flagella pore formation rather than the axoneme being targeted to a predetermined position on the membrane. Because only the caudal flagella were observed to lack exit sites, it is likely a secondary defect of Nek8445 depletion.

The caudal cytoplasmic axonemes make the longest run through the cell body and are unique in their association with the funis, a ribbon of microtubules that runs along the length of the cytoplasmic axoneme of each caudal flagellum (Benchimol *et al.*, 2004). The ribbon originates near the basal bodies and as it progresses toward the posterior, individual microtubules splay out to form rib-like structures that match the shape of the cell (Figures 1 and 8; Benchimol *et al.*, 2004). A defect in axoneme guidance or stabilization could lead to defects in specifying the exit site if, for example, the axoneme must remain in place for the exit site to be assembled. Indeed, the axoneme exit defect was

specifically associated with rounded Nek8445-depleted cells, which we found also lack funis microtubules (Figure 8). This suggests that the funis is responsible for establishing posterior cell morphology and additionally acts as a scaffold for guiding caudal axoneme exit.

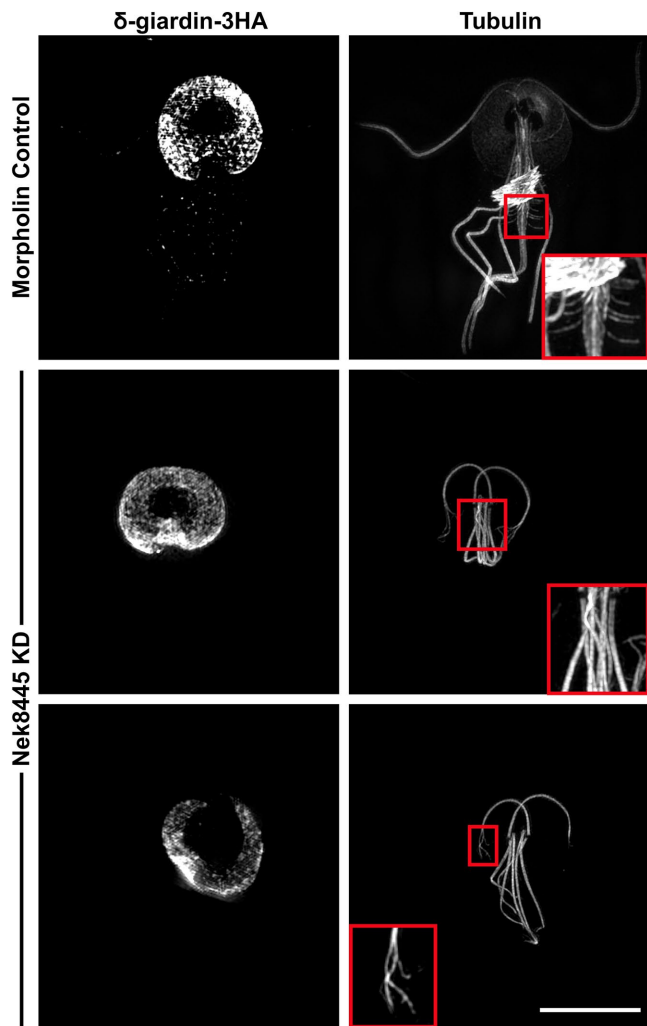


FIGURE 8: Structured Illumination superresolution imaging of δ -giardin-3HA and tubulin (6-11B-1 antibody) in morpholino-treated cells. Nek8445-depleted cells lack median bodies, a funis, and the axonemes have a fraying defect. Control inset shows a magnified view of funis microtubules. Middle inset shows that the Nek8445-depleted cells lack both a median body and funis. The bottom image shows an example of a Nek8445-depleted cell with abnormal disk morphology and splayed axoneme tips. Scale bar = 5 μ m.

Although many Nek8445-depleted cells had short flagella, we interestingly observed other cells with longer cytoplasmic axonemes trapped within the cell. This is consistent with the idea that *Giardia* can employ IFT-independent cytoplasmic axoneme growth as cytoplasmic axonemes do not necessarily need active IFT for growth (Briggs *et al.*, 2004; McInally *et al.*, 2019b). The importance of this observation remains to be determined. IFT has been assessed in interphase *Giardia*, where it has been shown that IFT train assembly occurs at the flagella pore/axoneme exit site (McInally *et al.*, 2019b). Regulation of new axoneme growth in mitotic cells remains unstudied. IFT-independent growth is likely important for the growth of nascent axonemes, which begin completely within the cell cytoplasm. We previously noted that after cytokinesis the ventral flagella, which have the shortest run through the cell body, regrow more quickly than the posterolateral pair (Hardin *et al.*, 2017). Whether this is due to regulated growth or the fact that the ventral axoneme would be the first to initiate an exit site and establish regulated IFT remains to be determined.

For an otherwise minimalistic parasite, the hugely expanded family of Nek kinases is intriguing. *Giardia* has 198 members compared with just 1 nonessential Nek kinase in yeast, 7 in *Arabidopsis*, 11 in humans, 13 in *Chlamydomonas*, and 39 in *Tetrahymena* (Wloga *et al.*, 2006; Manning *et al.*, 2011; Takatani *et al.*, 2015). Each of the three *Giardia* Nek kinases that have been studied appear to regulate different cellular processes. Depletion studies of *G/Nek1* and *G/Nek2* have not been performed, so whether they are required for cell division remains to be determined. However, localization studies did reveal that these proteins relocated during mitosis. In general, Nek kinases are activated during the G2/M transition by mitotic kinases (Fry *et al.*, 2017), but Nek kinases have also been shown to function during interphase (Chang *et al.*, 2009; Cohen *et al.*, 2013; Govindaraghavan *et al.*, 2014). While we cannot rule out interphase function, relocalization of Nek8445, *G/Nek1*, and *G/Nek2* during mitosis suggests these kinases likely regulate the microtubule cytoskeleton in a cell cycle-dependent manner (Smith *et al.*, 2012). The anaphase promoting complex, which canonically regulates cell cycle events through proteolytic degradation, has not been identified in *Giardia* (Gourguechon *et al.*, 2013). It has been speculated that kinases could play an expanded role in regulating cell cycle-related processes (Gourguechon *et al.*, 2013); the expanded Nek kinase family potentially fills this role. Future studies aimed at identifying pathways regulated by additional Nek kinases will be needed to further support this hypothesis.

Given the total number of Nek kinases in *Giardia*, we anticipated that each kinase would have a limited role in cytoskeletal regulation, yet Nek8445 depletion disrupts nearly all of the major microtubule structures in *Giardia*. The exception is that Nek8445 does not appear to regulate centrosome separation or spindle function as mitosis appears to progress normally. The other microtubule defects could result from a role for Nek8445 in regulating a key microtubule-binding protein with a role in regulating *Giardia*'s microtubule-based structures. For instance, EB1 has important roles in regulating microtubule dynamics and flagella length (Pedersen *et al.*, 2005; Schroder *et al.*, 2007). Several studies have localized Nek kinases to centrosomes, the tips of microtubules and axonemes, and implicated a relationship with EB1 (Mahjoub *et al.*, 2004; O'Regan and Fry, 2009; Govindaraghavan *et al.*, 2014; Prosser *et al.*, 2015). The relationship between Nek kinases and EB1 is not fully characterized, but *Aspergillus nimA* has been shown to require EB1 for its localization (Govindaraghavan *et al.*, 2014). Intriguingly, EB1 knockdown in *Giardia* has been shown to reduce median body volume, flagella length, and cause loss of axoneme central pairs (Kim and Park, 2019). Although less severe, these phenotypes are similar to what we observe in Nek8445-depleted cells. It will be interesting to identify Nek8445 phosphorylation targets to gain insight into microtubule array regulation in *Giardia*.

Whether many other Nek kinases play as critical a role as Nek8445 in cell division and microtubule array biogenesis remains to be determined, but the presence of a small gatekeeper residue in Nek8445 makes this protein kinase a particularly attractive drug target as this structural feature renders this kinase sensitive to BKIs. Importantly, we have already shown this class of inhibitor is effective at killing parasites resistant to metronidazole (Hennessey *et al.*, 2016), the frontline treatment for giardiasis.

METHODS AND MATERIALS

Parasite culture

G. lamblia wild-type strain WBC6 (ATCC 50803) cells were grown in TYI-S-33 medium supplemented with 10% bovine serum and 0.05 mg/ml bovine bile (Keister, 1983). Cells were cultured at 37°

under hypoxic conditions using 15 ml polystyrene screw-cap tubes (Corning Life Sciences DL).

Vector construction and transfection

With the exception of Rab11, the constructs used in this study were C-terminal fusions. All constructs were integrated into the genome using single-site recombination to maintain endogenous expression levels (Gourguechon and Cande, 2011). Primer sequences and restriction enzymes are shown in Supplemental Table S1. All PCR amplifications were performed using iProof DNA polymerase (Bio-Rad). Typically, an amplicon of ~1 kb in length that lacked the start codon was cloned in frame to a C-terminal triple-hemagglutinin epitope tag (3× HA) of the pKS_3HA_Neo or PAC plasmid (Gourguechon and Cande, 2011). The plasmids were linearized at a unique restriction site in the coding region of the gene of interest and ~5 µg of DNA was used to transfect wild-type *Giardia*. Transfected cells were selected with G418 at 40 µg/ml or puromycin at 45 µg/ml.

To follow Nek8445 localization throughout mitosis and cytokinesis, the C-terminus of GL50803_8445 was tagged with an 11-amino acid flexible linker and the fluorescent protein mNeonGreen (Allele Biotechnology). The mNG-N11-Neo vector was constructed for this purpose by amplifying mNeonGreen using the primers listed in Supplemental Table S1. The PCR product was cloned into the pKS 3HA Neo vector (Gourguechon and Cande, 2011) after excising the 3HA tag with *Bam*H1 and *Eco*R1.

Live cell imaging

Cells were imaged in 35-mm glass bottom dishes overlaid with agarose growth medium at 37°C under hypoxic conditions as described in Hardin *et al.* (2017).

Immunofluorescence microscopy analysis

Fixed cell imaging was performed as described in Paredes *et al.* (2011). In cases where a higher mitotic index was desired, the starve and release method was utilized before fixation (Sagolla *et al.*, 2006). Briefly, 4-day-old past confluent cultures were inverted to loosen dead cells and then the dead cells and spent medium were poured off and replaced with fresh medium. Four hours later, the cells were pelleted at 500 × g at room temperature. The pellet and remaining attached cells were fixed in PME (100 mM Pipes, pH 7.0, 5 mM EGTA, 10 mM MgSO₄) plus 2% paraformaldehyde, 0.025% Triton X-100, 100 µM MBS, and 100 µM EGS for 30 min at 37°C. Cells were again pelleted, washed with PME, resuspended with PME, and adhered to poly-L-lysine- (Sigma-Aldrich) coated coverslips. Cells were permeabilized in PME + 0.1% Triton X-100 for 10 min, then washed twice with PME + 0.05% Triton X-100 and blocked for 30 min in PME-BALG (PME + 1% bovine serum albumin, 0.1% NaN₃, 100 mM lysine, 0.5% cold water fish skin gelatin ; Sigma-Aldrich, St. Louis, MO).

Cells were stained with rabbit anti-*Gi*Actin antibody 28PB+1 (Paredes *et al.*, 2011) and mouse monoclonal anti-HA (Clone HA7; Sigma-Aldrich) both diluted 1:125 in PME-BALG and incubated overnight. After three subsequent washes with PME + 0.05% Triton X-100, cells were incubated with Alexa-488 goat anti-mouse and Alexa-555 goat anti-α-rabbit (Sigma-Aldrich, St. Louis, MO; diluted 1:150 in PME-BALG) for 1 h. Cells were washed three times with PME + 0.05% Triton X-100. The coverslips were mounted with ProLong Gold or Prolong Diamond anti-fade plus DAPI (ThermoFisher Scientific, Rockford, IL). Fluorescence deconvolution microscopy images were collected as described in Hardin *et al.* (2017). For all standard resolution microscopy experiments, a minimum of 200 cells were examined with at least 20 imaged per experiment. Structured illumination images were acquired on an OMX SR (GE Healthcare).

Morpholino knockdown and growth assays

Trophozoites were cultured until 85–100% confluence, placed on ice for 30 min to detach cells, pelleted at 500 × g for 5 min, and medium was replaced with 1.0 ml fresh ice-cold *Giardia* growth medium. Cells and cuvettes were chilled on ice. Lyophilized morpholinos listed in Supplemental Table S1 (Gene Tools, Philomath, OR) were resuspended in sterile water and 30 µl of a 100 mM morpholino stock was added to 300 µl of cells in a 4-mm cuvette. We used Gene Tools standard control morpholino as a negative control. Cells were electroporated (375 V, 1000 µF, 750 ohms; GenePulser Xcell; Bio-Rad, Hercules, CA) and returned to growth medium. See Krtkova and Paredes (2017) for a detailed protocol.

Western blot analysis

Giardia trophozoites were chilled on ice for 30 min to detach them from the culture tube side walls. Cells were pelleted at 700 × g, washed once in HEPES buffered saline, then resuspended in 300 µl of lysis buffer (50 mM Tris, pH 7.5, 150 mM NaCl, 7.5% glycerol, 0.25 mM CaCl₂, 0.5 mM dithiothreitol, 0.5 mM phenylmethylsulfonyl fluoride, 0.1% Triton X-100, Halt 100X Protease Inhibitor Cocktail (ThermoFisher Scientific), then sonicated. The lysate was cleared by centrifugation at 10,000 × g for 10 min at 4°C and then boiled in 2× Laemmli Sample Buffer (Bio-Rad). After SDS-PAGE, samples were transferred to Immobilon-FL membrane (Millipore) following the manufacturers' directions. Primary polyclonal rabbit anti-*Gi*Actin 28PB+1 (Paredes *et al.*, 2011), monoclonal anti-HA mouse HA7 antibodies (IgG1; Sigma-Aldrich), 6-11B-1 anti-acetylated tubulin (IgG2b), and DMA1 anti-α-tubulin (IgG1, Sigma-Aldrich) were diluted 1:2500 in blocking solution (5% dry milk, 0.05% Tween-20 in TBS). Secondary isotype-specific anti-mouse Alexa-555 or anti-rabbit Alexa-647 antibodies (Molecular Probes) were used at 1:2500. Horseradish peroxidase-linked anti-mouse or anti-rabbit antibodies (Bio-Rad) were used at 1:7000. Multiplexed immunoblots were imaged on a Chemidoc MP (Bio-Rad). Digital images were exported to 16-bit TIFF files and protein levels were determined by measuring integrated density using ImageJ (Schneider *et al.*, 2012). Protein levels in Figure 2A were normalized using actin as a loading control. For equal loading in Figure 4D the cells were counted using a MoxiZ Coulter counter (ORFLO) and the total cell number was adjusted to match the tube with the lowest cell number before making lysates. Reported values are the mean of three independent experiments.

SEM

For SEM analysis, trophozoites were pelleted at 700 × g for 7 min, fixed with 2.5% glutaraldehyde in 0.1 M phosphate-buffered saline (PBS), pH 7.4 and incubated for 1 h. After fixation, cells were pelleted again at 700 × g for 7 min and washed twice and resuspended in PBS. Cells were adhered to poly-L-lysine- (Sigma-Aldrich) coated coverslips for 15 min, then washed three times for 5 min with PBS and postfixed in 1% OsO₄ for 1 h. Cells were washed three times for 5 min with dH₂O, dehydrated in alcohol, dried to critical point with CO₂, sputter coated with gold, and analyzed on a JEOL JSM-840A scanning electron microscope.

Accession numbers

Genes/proteins. Nek8445: GL50803_8445; Axoneme Exit Site Protein: GL50803_8855; Rab11: GL50803_1695; Actin: GL50803_40817; δ-giardin: GL50803_86676.

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