TgCep250 is dynamically processed through the division cycle and is essential for structural integrity of the Toxoplasma centrosome

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ABSTRACT The apicomplexan centrosome has a unique bipartite structure comprising an inner and outer core responsible for the nuclear cycle (mitosis) and budding cycles (cytokinesis), respectively. Although these two cores are always associated, they function independently to facilitate polyploid intermediates in the production of many progeny per replication round. Here, we describe the function of a large coiled-coil protein in Toxoplasma gondii, TgCep250, in connecting the two centrosomal cores and promoting their structural integrity. Throughout the cell cycle, TgCep250 localizes to the inner core but, associated with proteolytic processing, is also present on the outer core during the onset of cell division. In the absence of TgCep250, stray centrosome inner and outer core foci were observed. The detachment between centrosomal inner and outer cores was found in only one of the centrosomes during cell division, indicating distinct states of mother and daughter centrosomes. In mammals, Cep250 processing is required for centrosomal splitting and is mediated by Nek phosphorylation. However, we show that neither the nonoverlapping spatiotemporal localization of TgNek1 and TgCep250 nor the distinct phenotypes upon their respective depletion support conservation of this mechanism in Toxoplasma. In conclusion, TgCep250 has a tethering function tailored to the unique bipartite centrosome in the Apicomplexa.

INTRODUCTION The phylum Apicomplexa consists almost exclusively of obligate intracellular parasites that have significant impact on public health. Among these single-celled organisms, Toxoplasma gondii is one of the most successful zoonotic pathogens that can be transmitted and replicate in many different host species. Toxoplasma gondii has developed flexible replication strategies allowing efficient survival in distinct tissue types. Tachyzoites are the acute replication form of T. gondii and are responsible for most of the clinical burden in infected individuals. They undergo a binary replication mode termed endodyogeny, wherein two daughter cells assemble within and emerge from the mother cell. The cell cycle follows the sequence of G1, S, and M phases, but the G2 phase is absent or too short to be detected (Gubbels et al., 2008) and is driven by a set of unconventional cyclin-dependent kinase-related kinases and cyclins (Alvarez and Suvorova, 2017). Mitosis is (semi)closed and spatiotemporally coordinated with cytokinesis, as these two events occur concomitantly. The 13 chromosomes are clustered at the centromeric region throughout the cell cycle at the “centrocone,” a unique membranous structure that houses the spindle microtubule during mitosis (Brooks et al., 2011). The tight organization of chromosomes is thought to ensure that each daughter cell receives a full set of genetic material when the parasite undergoes a more complex division cycle in other developmental stages (Francia and Striepen, 2014). In its definitive feline host, endopolygeny consists of several rounds of DNA synthesis and mitosis in the same cytoplasmic mass followed by synchronized cytokinesis in accordance with the last round of mitosis, producing 8–16 uninucleated progeny per replication cycle (Ferguson et al., 1974).
The Toxoplasma centrosome is divergent from mammalian cells in architecture and composition. For example, the centrioles are composed of nine singlet microtubules, smaller in size than mammalian centrioles, and the centriole pair is arranged in parallel rather than perpendicular (Francia and Striepen, 2014). Orthologues of many key components in the mammalian centrosome cannot be found in the Toxoplasma genome (Morlon-Guyot et al., 2017). Even without canonical protein orthologues, recent studies have shown that a group of coiled-coil proteins localize to the centrosome and support proper karyokinesis and cytokinesis (Suvorova et al., 2015; Courjol and Gisso, 2018). This work identified a distinctive architecture of two centrosomal cores: the outer core (distal to the nucleus) and the inner core (proximal to the nucleus). Independent activation of the inner and outer cores has been proposed as the mechanism underlying the uncoupling of S/M from daughter budding during polyplody intermediate stages across apicomplexan division modes, including endopolygeny in Toxoplasma and schizogony in the malaria-causing Plasmodium spp. (Chen and Gubbels, 2015; Suvorova et al., 2015).

The Toxoplasma centrosome is the key organelle coordinating mitotic and cytokinetic rounds. The centrosome resides at the apical end of the nucleus during interphase and remains closely associated with a specialized nuclear envelope fold, the centrocroce, which houses the spindle microtubules during mitosis (Gubbels et al., 2006). The centrosome also plays a critical role in partitioning the single-copy Golgi apparatus and the chloroplast-like organelle, the apicoplast, during cell division (Nishi et al., 2008). Late in G1, the centrosome rotates to the distal end of the nucleus, where it duplicates and then returns to the proximal end (Chen et al., 2015). At this point, the spindle microtubules assemble and are stabilized within the centrocroce, whereas subsequently the centrosome serves as a scaffold for assembly of daughter cell cytoskeletal components (Anderson-White et al., 2012). In mitosis, a striated fiber assembly (SFA) protein emanates from the centrosome and connects the daughter apical conoid to the centrocroce. SFA polymerization is thought to maintain the stoichiometric count of daughter cells and replicating genomic materials. In SFA-depleted cells, cell division is completely disrupted (Francia et al., 2012). Little is known about centrosome biogenesis or regulation during these tightly regulated events in Toxoplasma. Although orthologues in charge of new centriolar biogenesis, Polo-like kinase PLK4 and its substrate STIL/Ana2/SASS, are absent from the Toxoplasma genome, previous work has shown that a conserved NIMA-kinase, TgNek1, is required for centrosomal splitting (Chen and Gubbels, 2013), and a TgMAPK-L1 plays a role in inhibiting centrosomal overduplication (Suvorova et al., 2015). However, exactly what aspects of the centrosome these kinases act upon and what their substrates are remains unclear.

Here, we asked whether the previously described TgCep250 (Suvorova et al., 2015) could be the substrate of TgNek1 required for centrosomal splitting orthologous to the human HsNek2–C-NAP1/Cep250 relationship in centrosome splitting (Agircan et al., 2014). In our interpretation of the data, this hypothesis is rejected, but it is instead revealed that TgCep250 is required to connect the inner core with the outer core and to maintain overall centrosomal integrity. Moreover, these data represent the first experimental observation directly hinting at a differential status of mother versus daughter centrosome in the Apicomplexa. In conclusion, although TgCep250 is not the critical TgNek1 substrate, its function in keeping centrosomal complexes connected appears to be parallel to the role of human C-NAP1/Cep250 in managing structural connections in the centrosome.

RESULTS
TgCep250 has a dynamic localization and is associated with proteolysis

To start charting the function of the previously reported Toxoplasma centrosomal protein TgCep250 (TGGT1_212880) (Suvorova et al., 2015), we first confirmed the subcellular localization of TgCep250 throughout the parasite’s tachyzoite division cycle. TgCep250 was tagged endogenously with a C-terminal yellow fluorescent protein (YFP) tag (Huynh and Carruthers, 2009), and the TgCep250-YFP parasite line was transfected with a fosmid expressing TgCep250L1-hemagglutinin (HA) as an inner core centrosomal marker (Suvorova et al., 2015). During the G1 phase of the cell cycle, TgCep250-YFP is closely associated with the centrosomal inner core (highlighted by TgCep250L1), whereas during cytokinesis, we observed four TgCep250-YFP foci (Figure 1). The upper two foci were in close apposition to the outer core, and these results are in agreement with previous observations (Suvorova et al., 2015).

To dissect the function of TgCep250, we generated a conditional knockdown line (TgCep250-cKD) using a tetracycline-regulatable promoter. Simultaneously, we inserted a single-Myc tag at the N-terminus of the gene (Figure 2A). Correct integration of the construct promoter. Simultaneously, we inserted a single-Myc tag at the N-terminus of the gene (Figure 2A). Correct integration of the construct through single homologous recombination was validated by PCR (Figure 2B). In the presence of anhydrotetracycline (ATc), Myc-TgCep250 protein becomes undetectable within 6 h of incubation by immunofluorescence assay (IFA) and Western blot (Figure 2, C and D). Interestingly, the N-terminal Myc-tagged TgCep250 only formed two foci colocalizing with outer core marker α-Centrin throughout the cell cycle. This suggested that the placement of the tag on TgCep250 either alters the subcellular localization or that proteolytic processing is associated with this differential localization pattern. To test this hypothesis, we added a C-terminal 3xTy tag to the Myc-TgCep250-cKD locus (Figure 3A) and analyzed TgCep250-3xTy by IFA and Western blot. As shown in Figure 3, B and C, C-terminally 3xTy-tagged TgCep250 forms four foci like TgCep250-YFP. Moreover, we observed multiple protein bands smaller than the full-length protein on the blot using α-Ty antibody (Figure 3C; note that full-length TgCep250 is a large coil-coiled protein with a molecular weight

FIGURE 1: Dynamic localization of TgCep250 throughout the cell cycle. (A) IFA of TgCep250-YFP parasites costained with α-GFP (Cep250, shown in green), α-Centrin (centrosomal outer core marker, shown in magenta), α-HA (centrosomal inner core marker shown in red), and DAPI (in blue). Images acquired by superresolution microscopy (SIM). (B) Single-color panels show dynamic localization of TgCep250-YFP during asexual division cycle, essentially as previously reported (Suvorova et al., 2015). Cell cycle stages as indicated.
FIGURE 2: Generation and validation of a TgCep250-cKD line. (A) Schematic representation of TgCep250 conditional knockdown (cKD) construction by single homologous recombination. (B) PCR validation using TATi-ΔKu80 (parent) and Myc-TgCep250-cKD genomic DNA as templates. Primer pairs as indicated in A. (C) IFA using α-Myc and α-Centrin showed that the Myc-tagged TgCep250 is depleted within 6-h incubation of ATc. Boxed areas are magnified in the 4× zoom panels. (D) Western blot using α-Myc showed that the protein expression level is down-regulated in the presence of ATc. The purple arrowhead marks the full-length Cep250 protein (note that TgCep250 has a predicted molecular weight of 762 kDa); the smaller bands present in both the −ATc and +ATc samples are considered aspecific and serve as loading controls.

FIGURE 3: Differential localization of TgCep250 is regulated by proteolysis. (A) Schematic representation of Myc-TgCep250-3xTy-cKD construction using CRISPR/Cas9. (B) Myc-TgCep250-cKD localized only to the outer core (stained with α-Myc and α-Centrin) and TgCep250-3xTy localized to both inner and outer core (stained with α-Ty and α-HA on endogenously tagged TgCep250L1-HA as inner core marker). Left, conventional wide-field microscopy; right, superresolution (SIM) microscopy. (C) Multiple protein bands were detected in Western blot using α-Ty antibody suggestive of extensive proteolytic processing. α-Tubulin antiserum (12G10) was used as a loading control. Orange arrowheads mark the specifically detected TgCep250 protein fragments. (D) Schematic interpretation of Western blot and IFA data representing the relationship between TgCep250 proteolysis and subcellular localization.

TgCep250 is essential and links the centrosomal inner and outer cores

Next, we probed the function of TgCep250 with a series of experiments using the TgCep250-cKD parasite line. In the presence of ATc, TgCep250-cKD parasites failed to form plaques, suggesting TgCep250 is essential for parasite survival (Figure 4A). Phenotypic analysis by IFA showed that, in the absence of TgCep250, the nucleus failed to partition to the daughter cells (Figure 4B, white arrow), and as a result, anuclear parasites were observed (Figure 4B, white arrowheads). After 24 h of ATc induction, ~90% of all vacuoles harbored a parasite with a mislocalized nucleus (Figure 4C). Furthermore, we noticed that the stoichiometric balance of 1:1 daughter to (outer)centrosome count was also disrupted, resulting in multiple Centrin foci being accumulated in a single cell boundary, as outlined by IMC3, a cortical cytoskeleton marker (Figure 4B, double arrowhead). To examine
TgCep250 stabilizes bipartite centrosome

However, the TgCep250 in the mother centrosome is stably incorporated, thereby maintaining inner and outer core adhesion in this centrosome. Indeed probing Myc-TgCep250-3xTy-cKD parasites with Ty antiserum confirmed that Cep250 was still present in the normal-looking, mother centrosome, whereas it was either strongly depleted or undetectable in the daughter centrosome (Supplemental Figure S1).

**FIGURE 4:** Loss of TgCep250 causes lethal centrosomal defects. (A) Plaque assay of TgCep250-cKD and the TATI-Δku80 parent line ± ATc for 7 d shows that TgCep250 is essential. (B) IFA of TgCep250-cKD stained with α-Centrin (in red) and α-IMC3 (in green) ± ATc for 24 h. Arrowheads mark parasites with nuclear loss; arrow marks a nucleus outside a parasite; double arrowhead marks a parasite with two nuclei and three outer cores; asterisk marks a parasite with daughter buds out of sync with the rest of the vacuole. (C) Quantification of nuclear loss as shown in B. Vacuoles were scored for loss of at least one nucleus from the present parasites. At least 100 vacuoles harboring four or more parasites per experiment and condition were counted; error bars represent SD from three independent experiments. (D) TgCep250-cKD parasites coexpressing TgCep250L1-HA were stained with α-Centrin (in red) and α-HA (in green) marking the outer and inner cores, respectively. Images are representative of observed phenotypes; gray box: control wild-type nuclei displaying duplicated centrosomes; green box: stray inner core complex in presence of two normal appearing inner/outer pairs; blue box: outer core duplication independent of inner core duplication; orange box: complete detachment of inner and outer cores. (E) Quantification of defective centrosomal phenotypes; bar colors match the colored boxes in D. At least 100 nuclei with a duplicated centrosome pair per experiment and condition were counted; error bars represent SD from three independent experiments.

the nuclear partitioning defect in more detail, we used different sets of antibodies to analyze TgCep250-depleted parasites. We defined three different classes of centrosomal defects in TgCep250-depleted cells using centrosomal inner (TgCep250L1) and outer core (Centrin) markers (Figure 4, D and E). Detachment of the inner core from the outer core was observed in the majority of mitotic, mutant parasites (marked in orange). Furthermore, we observed parasites in which the outer core split into two foci, while only two inner cores were present, suggestive of the parasite undergoing an additional replication cycle (marked in blue; 10% of the mitotic parasites). Finally, nearly 35% of the mitotic parasites displayed stray inner core foci (marked in green). Detachment of inner and outer cores likely results in the nuclear-partitioning defect observed in Figure 4B, which therefore is a secondary defect. An additional observation is that detachment of inner and outer cores occurred in most instances on only one of the two centrosomes. The most logical explanation is that the detached inner and outer core pair represents the daughter centrosome and the intact pair represents the mother centrosome (Bornens and Gonczy, 2014). In this scenario, during centrosomal biogenesis, TgCep250 cannot be incorporated into the newly formed daughter centrosome, resulting in disconnection of the inner and outer cores.

Disconnected inner and outer cores retain their function

The anuclear phenotype observed in TgCep250-cKD parasites was very similar to the anuclear daughters observed in parasites depleted in kinetochore component TgNuf2 (Farrell and Gubbels, 2014). We tested how these two observations relate to each other and whether the inner core is still functional when it is separated from the outer core. IFA assays using mitotic markers revealed the interactions between centrosomal outer core (α-Centrin), the kinetochore (α-TgNdc80 or α-TgNuf2) (Farrell and Gubbels, 2014), and the spindle microtubules (EB1-YFP or α-TgEB1) (Chen et al., 2015) (Figure 5A). In the absence of TgCep250, both kinetochore complex and spindle microtubules became detached from the centrosomal outer core (Figure 5A, white arrowhead). Using an EB1 and Centrin costain, we determined that, while the spindle retained its normal morphology, the disconnect from the outer core occurred in roughly 50% of the vacuoles going through the first division following phenotype induction (Figure 5B). To determine whether TgCep250 plays a role in interaction between the inner core and the kinetochore,
FIGURE 5: The inner and outer cores remain functional upon dissociation in TgCep250-depleted parasites. Assessment of inner core (A–D) and outer core (E and F) function in TgCep250-depleted parasites. (A) IFA of TgCep250-cKD ± ATc for 18 h stained with α-Centrin (outer core; red) and costained with either α-TgNdc80 (kinetochore; green; left) or coexpression of TgEB1-YFP (spindle microtubules; green; right). Arrowheads mark nuclei displaying loss of outer core association with the kinetochore; double arrowheads mark nuclei displaying loss of outer core association with the spindle. No spindle or kinetochore defects were observed, indicating the inner core is fully functional. (B) Quantitation of the Centrin and EB1 staining represented in A using TgCentrin1 and TgEB1 antisera on TgCep250-cKD parasites induced ± ATc for 18 h. The incidence of loss of the connection between these structures in at least one nucleus per vacuole was scored. At least 100 nuclei with a spindle EB1 signal per experiment and condition were counted; error bars represent SD from three independent experiments. (C) IFA of TgCep250-cKD ± ATc for 18 h expressing TgCep250L1-HA (inner core: α-HA; green) and costained α-TgNdc80 (kinetochore; red) showed that the interaction between the kinetochore and inner core centrosome remains intact in the absence of TgCep250. (D) Quantitation of the phenotype observed in C using HA and TgNuf2 antisera on TgCep250-cKD parasites induced ± ATc for 18 h. The incidence of loss of the connection between these structures in at least one nucleus per vacuole was scored for loss of no, one, or two lost connections. Also scored, but not plotted, were incidences of kinetochores outside the nucleus in absence (1.16 ± 0.48%) and presence of ATc (0.76 ± 0.67%). At least 100 nuclei with a duplicated kinetochore per experiment and condition were counted; error bars represent SD from three independent experiments. (E) Superresolution (SIM) IFA of TgCep250-cKD ± ATc coexpressing TgCep250L1-HA stained with α-ISP1 (green: early daughter scaffold marker), α-Centrin (red: outer core), and α-HA (cyan: inner core). Numbered, boxed areas are magnified on the right. Note that outer cores are normally associated with daughter scaffolds (asterisk), even if the inner core connection is lost (arrowhead). (F) Superresolution (SIM) IFA of TgCep250-cKD coexpressing TgCep250L1-HA stained with α-HA (green: inner core), α-Centrin (blue: outer core), α-TgSFA (magenta: daughter tether), and α-TgNdc80 (red: kinetochore). Dotted lines outline parasites. (G) Summarizing schematic representation of data interpretation illustrating that inner and outer cores function normally upon their separation due to TgCep250 depletion.
FIGURE 6: The kinetochore is required to anchor centrosomal cores to the nuclear periphery. (A) TgNuf2-cKD parasites treated ± ATc for 18 h stained with centrosomal outer core (α-Centrin, in green) and inner core (TgCep250L1-HA; α-HA, in red) markers. Asterisks mark intact nucleus–centrosome connections; arrowheads mark intact inner and outer core pairs stayed from the nucleus (the associated small DAPI spot is the plastid genome); arrows mark missegregated nuclei. (B) Quantification of data in A. At least 100 vacuoles with a duplicated centrosome pair in each parasite were counted per experiment and condition and scored for loss of no, one, or two centrosomes per nucleus (maximum observed per vacuole was scored); error bars represent SD from three independent experiments. (C) The centrosomes counted in B were further differentiated for inner and outer core separation; error bars represent SD from the three independent experiments as in B. Statistical analysis by two-tailed t test.

summarized in Figure 5G, although we saw inner core fragmentation in the absence of Cep250, the inner core fragment retained at the nucleus remained associated with the kinetochore and was fully functional in spindle formation, whereas the detached outer core was still fully able to nucleate daughter scaffolds.

TgNuf2 anchors the centrosome to the nuclear periphery
To further dissect the association between the inner core and the kinetochore, we expressed the inner core marker TgCep250L1 in a kinetochore-depleted parasite line, TgNuf2-cKD (Farrell and Gubbels, 2014). It has been reported that, in the absence of TgNuf2, the association between the (outer)centrosome and spindle pole was abolished and that the (outer)centrosome pulled away from the nucleus (Farrell and Gubbels, 2014). With the inner core marker in the Nuf2-cKD background, we observed that the connection between the inner and outer core remained intact and that the whole centrosome dissociated from the nucleus (Figure 6A). We observed an approximately equal number of parasites in which either one or both centrosomes were dissociated (Figure 6B), which suggests there is not a mother and daughter distinction in this scenario. Although the majority of centrosomes did not fragment, we observed a statically relevant increase in inner and outer core dissociation from 2.4 to 10% (Figure 6C).

TgNek1 promotes separation of only the outer cores
It has been shown that a Toxoplasma orthologue of NIMA kinases, TgNek1, facilitates centrosomal splitting in tachyzoites (Chen and Gubbels, 2013). To dissect the regulation of centrosomal core splitting in a TgNek1-depleted background, we constructed a conditional knockdown line of TgNek1 (TgNek1-cKD) (Supplemental Figure S2A). Plaque assays showed that, in the presence of ATc, TgNek1-cKD parasites display a severe growth defect (Supplemental Figure S2B) consistent with the temperature-sensitive TgNek1 phenotype previously reported (Chen and Gubbels, 2013). Moreover, IFA with α-TgNek1 confirmed that TgNek1 expression is depleted in the presence of ATc (Supplemental Figure S2C). Interestingly, upon depletion of TgNek1, the duplicated outer cores remain connected, whereas the inner cores separate along with the kinetochores (Figure 7A). The distance between separated kinetochores in TgNek1-cKD and wild-type parasites is comparable, which in turn is comparable with previously reported distances (Suvorova et al., 2015). Quantification of kinetochore separation relative to outer core separation revealed that this is a very strong phenotype (Figure 7, B–D) and indicates that the spindle is able to separate the kinetochores, along with the inner cores, when outer core splitting is blocked.

DISCUSSION
In mammalian cells, duplicated centrosomes are tethered together by a fiber complex composed of C-NAP1/Cep250 (Fry et al., 1998), rootletin (Bahe et al., 2005), and LRRC45 (He et al., 2013). Phosphorylation of C-NAP1/Cep250 by NIMA kinase Nek2 promotes degradation of C-NAP1/Cep250 and results in splitting of the linked centrosomes (Fry et al., 1998). Reciprocal BLAST analysis revealed a family of Toxoplasma proteins displaying features of the C-NAP1/Cep250 and rootletin orthologues, of which two localize to the centrosomes: TgCep250 (763 kDa) and TgCep250L1 (296 kDa) (Suvorova et al., 2015; Morlon-Guyot et al., 2017). Because TgCep250 showed dynamic changes associated with centrosomal replication (Suvorova et al., 2015), we asked whether regulatory and/or structural features might be conserved as well between mammals and Toxoplasma.

We showed that the functional orthologue of human Nek2 in Toxoplasma, TgNek1, is essential for outer core but not inner core splitting (Figures 7 and 8). This observation implies that the sub-strate of TgNek1 only tethers the outer core and the mechanism that physically separates the inner and outer core is different. Both TgNek1 and TgCep250 are closely associated with the centrosomal outer core; however, their timing is different. TgNek1 is only transiently recruited to the centrosome in G1/S-phase transition and disappears after the cell cycle progresses into mitosis (Chen and Gubbels, 2013), which is just when TgCep250 shows up in the outer core (summarized in Figure 8). Thus, although we cannot formally exclude the possibility that TgCep250 is a TgNek1 substrate, the proteolytic event of TgCep250 that leads to both inner and outer core localization is likely independent from TgNek1 activity, because TgNek1 is already absent from the centrosome during cytokinesis. Phenotypic analysis of the TgCep250-cKD and TgNek1-cKD phenotypes also showed different cell cycle defects, indicating the roles of TgCep250 and TgNek1 are functionally distinct (Figure 8). It is worth noticing that TgCep250 is phosphorylated at Ser-252 in whole-organism phosphoproteome data (Treek et al., 2011). However, the function of this TgCep250 phosphorylation and the kinase that phosphorylates it remain to be determined. HsNek2 phosphorylation of HsCep250 triggers proteolytic processing of HsCep250 and results in splitting of the duplicate centrosome. Notably, TgCep250 is also proteolytically processed and provides another parallel across
species. However, we do not know the exact function of TgCep250 in Toxoplasma, but we do know it is associated with the outer core at the onset of cytokinesis. It is tempting to speculate this could be related to activation of the outer core toward cytokinesis, or alternatively, it could be a step in centrosomal maturation. This question cannot be answered in Toxoplasma tachyzoites, as every mitotic round is connected with daughter budding.

In mammalian cells, the differential decoration of centriolar appendages defines the difference in age and ability to nucleate microtubules between the centriolar pairs (Bornens and Gonczy, 2014). Toxoplasma’s annotated genome does not contain orthologues of mammalian centriolar appendage genes, and no clear appendage structure was observed with transmission electron microscopy. Functionally, however, differential states of mother and daughter centrosomes have been postulated to originate the nongeometric daughter expansion numbers observed during schizogony in the Plasmodium spp. red blood cell cycle (Arnott et al., 2011). In the proposed model, the mother centrosome is ready for another round of duplication, associated with S phase and mitosis, sooner than the daughter centrosome, which still has to mature, and this is the mechanistic basis for nongeometric daughter numbers (Arnott et al., 2011). In direct support of this model is our observation that the detachment of outer and inner cores occurs mostly in one pair of centrosomes upon TgCep250 depletion. We show that TgCep250 is retained in the intact centrosome (mother) but is depleted from the destabilized daughter centrosome (Supplemental Figure S1). Our results therefore provide the first experimental evidence that daughter and mother centrosomes are different in apicomplexan parasites.

Overduplication of outer cores has been reported in parasites expressing a temperature-sensitive allele of the peripheral centrosomal matrix (PCM) component MAPK-L1 (Suvorova et al., 2015), whereas outer core fragmentation was observed in TgCep530-depleted parasites (TgCep530 resides in between the inner and outer cores) (Courjol and Gissot, 2018). In TgCep250-depleted cells, we observed multiple outer cores, which could be the result of either overduplication or fragmentation (Figure 4, D and E). The TgCep250 phenotypes differs from the MAPK-L1–defective cells, as an accumulation of inner and outer cores aligned in a “dumbbell form.”
was not observed. On the other hand, the TgCep250 phenotype is also dramatically different from the multiple outer cores of various intensity observed in TgCep530-deficient parasites. The multiple outer cores observed upon TgCep250 depletion appear clearly organized and are of equal intensity, which is more consistent with an overduplication than with fragmentation. In addition, we observed normal initiation of daughter buds in the absence of TgCep250 (Figures 5, E and F, and 8), suggestive of fully functional outer cores.

This observation provides further support for the hypothesis of independent function and regulation of inner and outer cores as the basis to uncouple mitosis from daughter budding throughout the different multiplication strategies in the Apicomplexa (Chen and Gubbels, 2015; Suworova et al., 2015). In addition, the formation of spindle microtubules on disconnected inner cores (Figures 5, A and B, and 8) is also in line with the independent functionality of the bifunctional centrosomal cores. Further support for this model is provided in the IFA assay combining SFA, Centrin, TgCep250L1, and TgNdc80 (Figure 5, F and G), as the integrity of each of these structures appears normal, which is essential and critical for the endowment of each daughter cell with a single copy of genetic material (Beck et al., 2010; Francia and Striepen, 2014). Indeed, we observed a hierarchical organization of these critical structures in wild-type parasites (Figure 5F). However, in TgCep250-cKD cells, the connection between the inner (green) and outer core (blue) is lost, while association between the SFA and outer core, as well as between the kinetochore and the inner core, remains intact (Figure 8). Taken together, our data are consistent with the previously hypothesized model of independent regulation of inner and outer cores in the bipartite centrosome (Suworova et al., 2015).

Our previous work showed that the kinetochore is required for outer core association with the nucleus (Farrell and Gubbels, 2014), because it locks the microtubules emanating from the centrosome onto the nuclear envelope (Chen et al., 2015). Using the novel tools developed here, we now expanded these insights by showing that, in TgNuf2-cKD parasites, the intact inner and outer core pair dissociates in its entirety from the nucleus (Figure 6). Thus, we can conclude that the kinetochore serves as a molecular anchor to lock the intact bipartite centrosome and that spindle microtubules anchoring to the nuclear periphery are not required to maintain the connection between the inner and outer cores.

Overall, we conclude that the centrosomal outer core, inner core, and kinetochore are hierarchically organized and that TgCep250 is essential for structural stability of the connection between the centrosomal cores embedded at the time of centrosomal duplication. Two different forms of Cep250 are present in tachyzoites: a full-length form localizing to the outer core during the transition between mitosis and cytokinesis and a proteolytically processed form localizing to the inner core throughout the tachyzoite cell cycle. It is unlikely that TgNek1 is the kinase triggering proteolytic cleavage. The tethering function of TgCep250 appears to be maintained between human and Toxoplasma orthologues, but the structures connected are tailored to each organism. The phosphorylation controls and the proteolytic mechanisms in Toxoplasma remain unknown, but provide an exciting area for future research.

MATERIALS AND METHODS
Parasite strains
Toxoplasma gondii tachyzoites of RHΔHX (Donald and Roos, 1998), RHΔKu80ΔHX (Huynh and Carruthers, 2009), TAT1ΔKu80 (Sheiner et al., 2011), and their transgenic derivatives were grown in human foreskin fibroblast as previously described (Roos et al., 1994). Pyrimethamine (1 μM), chloramphenicol (20 μM), or mycophenolic acid (25 μg/ml) in combination with 50 μg/ml xanthine was used to select stable transgenic parasite lines and cloned by serial dilution. We used 1.0 μg/ml anhydrotetracycline to repress specific gene expression of conditional knockout lines. Cep250L1-HA fosmid was kindly provided by Michael White, University of South Florida (Vinayak et al., 2014). The tub-EB1-YFP/sagCAT plasmid has been described before (Chen et al., 2015).

Plasmid constructs
All primers used are listed in Supplemental Table S1. Endogenously tagged parasite strains were generated as previously described (Sheiner et al., 2011). Briefly, 1- and 1.9-kb regions upstream of the stop codon of the TgCep250 gene were PCR amplified using genomic DNA from the Toxoplasma RH strain as a template and were cloned by ligation-independent cloning (LIC) into pYFP-LIC-DHFR (kindly provided by Vern Carruthers, University of Michigan). NarI was used to linearize pCep250-YFP-LIC-DHFR for site-specific homologous recombination.

Tetracycline-regulatable parasite strains were generated by single-crossover homologous recombination using a plasmid kindly

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**FIGURE 8:** Schematic summarizing the spatiotemporal roles of TgCep250, TgNuf2, and TgNek1 throughout centrosomal replication, mitosis, and onset of daughter budding. Top sequence represents normal development, with the appearance of Nek1 and Cep250 specifically indicated. Note that Nek1 and Cep250 do not overlap in timing or location, thus making it unlikely Cep250 is a Nek1 substrate. Bottom schema represent the phenotypes upon depletion of the factors as labeled at the bottom. Note that Cep250 is retained on the mother (M) centrosome in the Cep250 knockdown line but results in a destabilized daughter (D) centrosome. For simplicity, daughter bud formation is represented by the appearance of ISP1 (see Figure 5E). SFA, striated fiber assembly; PCM, peripheral centrosomal matrix.
provided by Wassim Daher, University of Montpellier, France (Morlon-Guyot et al., 2014). The plasmid was redesigned by adding a single Myc-tag between the minimum promoter and the N-terminal genomic region of the target genes. All the inserts were amplified by PCR and digested with the enzyme pairs BglII and NotI. For construction of the Myc-Cep250-cKD line, the 5’ end of the TgCep250 was amplified, digested with BglII and NotI enzyme pairs, and ligated into the DHFR-TetO7sa94-Myc vector. The resulting construct was linearized using NcoI, electroporated into the TAT1ΔKu80 line by electroporation, and selected for pyrromethamine resistance. The Myc-NeK1-cKD line was generated following the same strategy.

The Myc-Cep250-3xTy-cKD line was generated by site-specific insertion of a PCR product carrying the tag and an HXGPRT selectable marker using CRISPR/Cas9. The pU6 plasmid expressing the guide RNA and Cas9 nuclease was kindly provided by Sebastian Lourido from the Whitehead Institute (Sidik et al., 2014). The CRISPR construct targeted the 3’UTR of TgCep250 near the stop codon. A primer pair was designed to amplify the 3xTy tag and the selectable marker (HXGPRT) flanking it with 40 base pairs of homologous region before the stop codon and after the protospacer-adjacent motif site. Ammplicon generated from the primer pair was gel purified and cotransfected with the protospacer and Cas9 encoding plasmid.

IFA and microscopy
Parasites were seeded overnight (~16–18 h) and fixed with methanol as previously described (Gubbels et al., 2006). The following antibodies were used in this study: Myc (MAb 9E10, mouse, 1:50; Santa Cruz); HA (3F10, rat, 1:3000; Roche); TgIMC3 (rat, 1:2000; Anderson-White et al., 2011); TgNuf2 and TgNdc80 (guinea pig, 1:2000; Farrell and Gubbels, 2014); SFA (rabbit, 1:1000; kindly provided by Boris Srienden, University of Georgia; Francia et al., 2012); TgNeK1 (1:1000; Chen and Gubbels, 2013); HsCentrin (rabbit, 1:1000; kindly provided by Iain Cheeseman, Whitehead Institute); Tg (mouse, 1:1000; kindly provided by Sebastian Lourido); TgEB1 (rat, 1:3000; this work).
Alexa Fluor A488, A568, A594, A633, and A647 secondary antibodies were used. We used 4’,6-diamidino-2-phenylindole (DAPI) to stain nuclear material. Images were acquired using a Zeiss Axiovert 200M wide-field fluorescence microscope equipped with a Plan-Fluo 100×1.4 NA oil objective and a Hamamatsu Orca-Flash 4.0 LT camera. For superresolution structured illumination microscopy (SIM), a Zeiss Eliza S.1 microscope equipped with a Plan-Apochromat 63×/1.40 oil objective and a CO-PO-Tech pco.edge 4.2 sCMOS camera in the Boston College Imaging Core was used in consultation with Bret Judson. Images were acquired and processed in Zeiss ZEN v. 2.3 software using the standard mode.

EB1 antiserum generation
For generation of N-terminal His6-tagged fusion protein, 600 base pairs from TgEB1 cDNA (corresponding to the N-terminal 200 amino acids) were PCR amplified using primers EB1-F/R-LUC-His and cloned into the pAVAA0421 plasmid (Alexandrov et al., 2004) by ligation-independent cloning. The fusion protein was expressed in BL21 STAR (DE3)pLysS Escherichia coli using 0.5 mM isopropyl β-D-1-thiogalactopyranoside in 2×YT broth overnight at 37°C and purified under native conditions over Ni-NTA agarose (Invitrogen). Polyclonal antiserum was generated by immunization of a rat (Covance, Denver, PA). Antiserum were affinity purified as previously described (Gubbels et al., 2006) against recombinant His6-TgEB1.

Western blot analysis
Freshly lysed parasites were collected after 3-μm filtration by centrifugation and washed twice in 1X phosphate-buffered saline. Parasite pellets were lysed by resuspension in 50 mM Tris-HCl (pH 7.8), 150 mM NaCl, and 1% SDS containing 1X protease inhibitor cocktail (Sigma-Aldrich) and heating at 95°C for 10 min. Parasite lysates were analyzed by SDS-PAGE on a 3–8% Tris-acetate gradient gel. Before polyvinylidene fluoride membrane transfer, the gel was subjected to 100 mM acetic acid for 5 h to break up the large TgCep250 protein. Transfer buffer contained 25 mM Tris (pH 8.3), 195 mM glycine, 0.025% (wt/vol) SDS, and 15% methanol. Blots were hybridized with α-Myc-HRP, α-Ty, or α-α-tubulin (MAb 12G10; Developmental Studies Hybridoma Bank).

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É. Ch.-T. Chen and M.-J. Gubbels Molecular Biology of the Cell
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