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ORIGINAL ARTICLE



The roles of Centrin 2 and Dynein Light Chain 8a in apical secretory organelles discharge of *Toxoplasma gondii*

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Abstract

To efficiently enter host cells, apicomplexan parasites such as *Toxoplasma gondii* rely on an apical complex composed of tubulin-based structures as well as two sets of secretory organelles named micronemes and rhoptries. The trafficking and docking of these organelles to the apical pole of the parasite is crucial for the discharge of their contents. Here, we describe two proteins typically associated with microtubules, Centrin 2 (CEN2) and Dynein Light Chain 8a (DLC8a), that are required for efficient host cell invasion. CEN2 localizes to four different compartments, and remarkably, conditional depletion of the protein occurs in stepwise manner, sequentially depleting the protein pools from each location. This phenomenon allowed us to discern the essential function of the apical pool of CEN2 for microneme secretion, motility, invasion and egress. DLC8a localizes to the conoid, and its depletion also perturbs microneme exocytosis in addition to the apical docking of the rhoptry organelles, causing a severe defect in host cell invasion. Phenotypic characterization of CEN2 and DLC8a indicates that while both proteins participate in microneme secretion, they likely act at different steps along the cascade of events leading to organelle exocytosis.

KEYWORDS

annuli, apicomplexa, centrin, conoid, dynein light chain, egress, exocytosis, invasion, microneme, motility, rhoptries, *Toxoplasma gondii*

1 | INTRODUCTION

Microtubules are filamentous tubulin heteropolymers found in all eukaryotic cells. They provide a framework that is essential for the maintenance of cellular architecture and shape. In addition to their structural role, microtubules are also involved in a wide variety of cellular processes such as chromosome segregation, intracellular transport of organelles and vesicles, as well as ciliary and flagellar motility (for a review see Reference 1). Microtubules originate from the

microtubule organizing center (MTOC) which is the site for coordinated microtubule nucleation. Remarkably, the obligate intracellular parasite *Toxoplasma gondii* harbors two MTOCs from which emerge distinct populations of microtubules. The centrocone is a unique spindle compartment of the nuclear envelope that is associated with centrosomes and is responsible for chromosome segregation during closed mitosis.² It has been implicated in daughter cell budding, division of the Golgi apparatus and the inheritance of a nonphotosynthetic plastid-like organelle called the apicoplast.³⁻⁵ The

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second MTOC is noncentrosomal and consists of a ring-shaped tubulin structure, positioned at the apex of the parasite, referred to as the apical polar ring (APR). Twenty-two subpellicular microtubules emerge from the APR and elongate along two-thirds of the parasite body. They form a cytoskeletal basket that supports the pellicle of the parasite and maintains its shape and rigidity. Present in all alveolates, the pellicle is composed of a unique triple lipid bilayer, which is made of the plasma membrane apposed to flattened membranous vesicles called the inner membrane complex (IMC). This is underlined by an elaborate network of intermediate filament-like alveolins.⁶ Anchored to the APR, the conoid is composed of a tightly packed spiral of atypical tubulin polymer fibers⁷ with a pair of intraconoidal microtubules at its center and two preconoidal rings at the top. The conoid is a motile structure which is retracted inside the cell body in intracellular parasites but extrudes through the APR in extracellular parasites due to a rise in intracellular calcium (Ca²⁺) levels.⁸

Besides these tubulin structures, the apical complex of *T. gondii* harbors two types of regulated secretory organelles, the micronemes and the rhoptries, that are secreted sequentially during host cell invasion. Micronemes contain, among other proteins, adhesins that are associated with the parasite plasma membrane postexocytosis and play a role in motility, invasion and egress.⁹ Rhoptries are elongated organelles composed of a bulbous region and a neck region containing ROPs and RONs proteins, respectively. Once secreted, a complex of RON proteins is inserted into the cytoplasmic face of the host cell plasma membrane and interacts with the microneme protein AMA1 (apical membrane antigen 1) at the parasite surface. This establishes a firm anchor, termed the moving junction, through which the parasite can propel itself into the host cell.¹⁰

Proper apical positioning and secretion of these two secretory organelles are crucial for successful invasion and hence the survival of *T. gondii*. Micronemes are typically organized like a crown around the APR and are associated with the subpellicular microtubules.^{11,12} The rhoptries are clustered together just beneath the conoid except for the one or two that are primed for secretion and are observed with their neck inserted into the conoid, possibly by docking to the intraconoidal microtubules.^{13,14} The acylated ARO protein is anchored to the cytoplasmic face of the rhoptry membrane and is essential for the apical positioning and clustering of the organelles.¹⁵⁻¹⁸ The integrity of the APR and subpellicular microtubules is crucial for microneme trafficking and anchoring to the apical pole of the parasite.¹²

Both sets of secretory organelles appear to be secreted at the apex of the parasite¹⁹ in response to elevated intracellular Ca²⁺. Microneme exocytosis depends on a pleckstrin-homology domaincontaining protein (APH) that is anchored by acylation to the surface of micronemes and is able to sense phosphatidic acid (PA).²⁰⁻²² Ultimately, binding of APH to PA triggers a membrane fusion event that releases the microneme content via a hypothetical SNARE-like complex involving DOC2.1.²³ The signaling cascade leading to rhoptry secretion is still unknown, nevertheless a recent study showed that parasites conditionally depleted for FER2, a Ca²⁺ sensor belonging to the ferlin family, are unable to discharge the rhoptries. This suggests that rhoptry secretion might also be driven by Ca²⁺ signaling.²⁴ Even though valuable advances have been made toward a better understanding of the rhoptry and microneme membrane fusion mechanisms, little is known about the docking and transport of these secretory organelles to the site of fusion at the tip of the parasite. Secretion of both organelles is thought to occur via the conoid, but most of the conoidal proteins studied so far have been implicated in conoid stability.^{12,25,26} Only depletion of the polar ring-associated protein, RNG2, has been reported to impact modestly on microneme exocytosis.²⁷

Both T. gondii centrin 2 (CEN2) and dynein light chain 8a (DLC8a) localize to the apical complex and are predicted to be essential based on the fitness score obtained from a CRISPR-Cas9 genome wide knockout screen.²⁸ Centrins are a group of highly conserved eukaryotic Ca²⁺-binding phosphoproteins that mediate Ca²⁺-dependent contraction of the ring structures typically associated with centrioles or flagellum.^{29,30} In T. gondii, CEN2 is found at the centrosome, the peripheral annuli, the preconoidal ring and the basal cup of the parasites.³¹ Intriguingly, the apical complex of the *T. gondii* tachyzoite is believed to originate from a repurposed flagellum.³² This could suggest that the apically localized CEN2 has flagellar origins. For example. in the flagellated green algae Chlamydomonas reinhardtii, centrins have been documented to contribute to Ca²⁺-mediated contraction and flagellum excision.³³ Intriguingly, the peripheral annuli are enigmatic structures that have been described only in T. gondii. They consist of six circles positioned at the lower edge of the apical cap. The functional relevance of these annuli is presently unknown and only two markers have been described, namely, CEN2 and the peripheral annuli protein 1 (PAP1).³⁴ The pool of CEN2 localizing to the basal cup is involved in the constriction of the basal pole, a nonessential process that occurs at the end of division.³⁵ The role of CEN2 at the other locations is unknown.

DLC8a belongs to a family of cytoskeletal motor proteins that form a functional complex with the cytoplasmic dynein heavy chains, intermediate chains and light intermediate chains. This complex transports cargoes, such as vesicles and organelles, along microtubules over long distances in the cytoplasm (for a review see Reference 36). DLC8 is found in all eukaryotes studied so far including those lacking cytoplasmic dynein heavy chains such as higher plants or red algae.³⁷ Among the four DLC8 proteins expressed by *T. gondii*,³⁸ DLC8a is the only one that has been proposed to be important for the tachyzoite lytic cycle.²⁸ One of the cytoplasmic dynein heavy chains conserved across the phylum of Apicomplexa is also important for parasite fitness suggesting that it might form a motor complex with DLC8a (Table S1, Supporting Information).³⁷ When expressed as a second copy, DLC8a localizes to the apical tip, spindle poles, centrosome and basal ring of the parasites,³¹ whereas endogenously C-terminal tagged DLC8a localizes only to the apical cap.³⁸

Here, we have embarked on a phenotypic analysis of these two conserved apicomplexan proteins typically associated with microtubules. By conditionally controlling CEN2 expression, we have identified its subcellular localization-dependent role in microneme secretion that ensures motility, invasion and egress. CEN2 also appears to be implicated in the integrity of the peripheral annuli. In contrast, DLC8a contributes predominantly to invasion by sustaining successive rounds of microneme secretion and by properly anchoring the rhoptries to the apical complex.

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2 | RESULTS

2.1 | Gradual depletion of CEN2 results in distinct phenotypes linked to its multiple localization

C-terminal tagging of CEN2 with yellow fluorescent protein (YFP) at the endogenous locus revealed punctate structures associated with the centrosome, the preconoidal ring, the peripheral annuli and the basal cup of the parasite.³⁹ CEN2 at the posterior pole participates in the basal constriction of the parasite in association with the myosin motor MyoJ, a function that has been shown to be dispensable for parasite survival.³⁵ Considering the highly negative fitness score for CEN2 mutants reported in a *T. gondii* genome-wide screen,²⁸ we postulated that CEN2 carried important additional functions at its other locations. To address this point, we dissected its role using a Tetinducible CEN2-YFP parasite line (CEN2-iKD).³⁵ Immunofluorescence



FIGURE 1 CEN2 is essential for the tachyzoite lytic cycle. A, Immunofluorescence time-course of the downregulation of CEN2-YFP upon ATc treatment. IMC1 stains the periphery of the parasites. The white arrows point to the centrosomes while the yellow arrowheads point to the enlarged basal pole of the parasites. Scale bars: 2 μ m. B, Scheme of the 48 hours lytic cycle of tachyzoite. Freshly egressed parasites are able to glide and attach to an uninfected host cell before reorientation and invasion. Multiple rounds of division occur within the parasitophorous vacuole (PV) before the parasites egress, lyse the cell and re-infect the neighboring uninfected cells. C, Plaque assay performed over 7 days with the parental (Δ KU80) and the CEN2 inducible (CEN2-iKD) strains in the presence or absence of ATc. CEN2-iKD parasites show a strong defect in their lytic cycle. Scale bars: 0.5 cm. D, Intracellular growth assay performed after 0, 48 and 72 hours of ATc treatment. No defect was detected for the CEN2-depleted parasites

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assays (IFA) revealed that CEN2 depletion occurred in a stepwise manner (Figure 1A). First, a loss of CEN2 was observed 24 hours after ATc treatment at the peripheral annuli and the conoid. Depletion of CEN2 from the basal cup occurred only at 48 hours (Figure 1A, yellow arrowheads). The pool of CEN2 associated with the centrosomes was the last to be affected and was still weakly detectable after 72 hours of ATc treatment (Figure 1A, white arrows).

We assessed the effect of long-term depletion of CEN2 by evaluating the viability of the parasites on plaque assay (Figure 1B). Every 48 hours, the parasites accomplish an entire lytic cycle ending by egressing from the infected cells and invading the neighboring ones. Over a period of 7 days, areas of lysis became visible on the monolayer, recapitulating several successful lytic cycle rounds. The monolayer infected with CEN2-depleted parasites appeared almost intact,







FIGURE 2 CEN2 is critical for microneme secretion. A, Invasion assay performed after 48 hours ± ATc on the parental (Δ KU80) and the CEN2 inducible strains (CEN2-iKD) revealing a severe defect in the invasion efficiency of the CEN2-depleted parasites. B, Egress assay performed after 48 hours ± ATc using BIPPO as inducer. The CEN2-depleted parasites exhibit a strong defect in their capacity to egress from the host cell. C, Motility assay performed after 48 hours ± ATc using BIPPO as inducer. The different types of gliding motility have been quantified by video-microscopy (see Figure S1B and C). D, Western blot analysis of microneme protein MIC2 secretion after induction with 2% ethanol for 30 minutes. The dense granule protein 1 (GRA1) shows the constitutive secretion of dense granules and catalase (CAT) shows the integrity of the parasites. ESA, excreted secreted antigens; pMIC2, processed MIC2. E, Attachment assay performed after 48 hours ± ATc. The attachment efficiency of the CEN2-depleted parasites is strongly affected. For A, B, C and E, results are presented as mean ± SD. The significance of the results has been assessed using an unpaired t test. The two-tailed P-values are written on the graphs. F, The localization and integrity of the apical secretory organelles were checked after 0 and 48 hours of ATc treatment using the markers MIC2, which stains the micronemes and ARO, anchored at the surface of the rhoptries. GAP45 stains the periphery of the parasites and actin (ACT1) shows the cytosol. The integrity of the apical cap of the IMC was verified using the marker ISP1. Scale bars: 2 µm

reflecting the deleterious consequence of CEN2 depletion on parasite survival (Figure 1C). Taking advantage of the sequential loss of CEN2, we evaluated the impact of CEN2 depletion from its various locations over time. To assess the role of CEN2 in division with regards to its centrosomal localization, CEN2-iKD parasites were grown in the presence of ATc for 72 hours. These parasites replicated in a manner comparable to the treated parental strain (control) (Figure 1D). We then evaluated the synchronicity of division at 48 hours of ATc treatment when the basal pool of CEN2 was depleted. In this assay, all the parasites within the same vacuole divided in a synchronous fashion, indicating that CEN2 does not play a role in the intravacuolar connection between parasites (Figure S1A). These observations reveal that CEN2 at the centrosome and basal pole is not critical for parasite proliferation.

The function of CEN2 at the apical pole of the parasite was assessed by phenotyping the parasites after 48 hours of ATc treatment. At that time, the pools of CEN2 at the apical tip and annuli are undetectable. CEN2-depleted parasites were impaired in their ability to invade host cells by 85% compared to control parasites (Figure 2A). We observed a comparable defect in the ability of the parasites to egress from the infected host cell (Figure 2B). Analysis of parasite motility by video microscopy revealed that 80% of CEN2-depleted parasites were non-motile vs 40% in the control strain (Figures 2C and S1B). Among the remaining motile parasites, the three types of movement, helical, circular and stationary twirling, were observed (Figure S1C). Successful invasion, egress and motility rely on adhesins that are released to the parasite surface upon microneme exocytosis (Figure 1B). As CEN2 depletion led to a defect in these three processes, we evaluated the ability of these parasites to secrete their micronemes. CEN2-depleted parasites showed a lower basal level of microneme secretion, compared to both control and untreated parasites, and a complete inability to respond to ethanol, a chemical inducer of microneme exocytosis (Figure 2D). In accordance with the defect in microneme exocytosis, parasite attachment to host cells was severely impaired, with attachment efficiency dropping by 70% between the control and CEN2-depleted parasites (Figure 2E). The absence of CEN2 did not affect the morphology and the subcellular distribution of micronemes and rhoptries (Figure 2F). Moreover, induced conoid protrusion was not impaired upon CEN2 depletion (Figure S1D). Overall, these results point to a key role for CEN2 in microneme exocytosis that ensures motility, attachment, invasion and egress.

2.2 | Depletion of CEN2 destabilizes the peripheral annuli

At the parasite apex, CEN2 localizes to the peripheral annuli, a structure of unknown function for which only one other marker has been described to date, the peripheral annuli protein 1 (PAP1).³⁴ Using super-resolution microscopy by stimulated emission depletion (STED), we confirmed that endogenously tagged PAP1 specifically localizes to the peripheral annuli (Figure 3A). Moreover, 3D reconstruction of PAP1 and CEN2 staining indicated that PAP1 forms a

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ring-shaped structure surrounding CEN2-YFP (Figure 3B). To evaluate if the loss of PAP1 affected CEN2 localization, we generated PAP1-KO transgenic parasites in the CEN2-iKD background (PAP1-KO/CEN2-iKD, Figure S2A). In the absence of PAP1, CEN2-YFP still localized correctly at the peripheral annuli (Figure 3C) and did not appear to be degraded (Figure 3D). Moreover, deletion of PAP1 did not affect the fitness of the parasites which were still able to accomplish their lytic cycle (Figure 3E). PAP1-KO parasites were able to invade, replicate and egress as efficiently as the control (Figure S2B-D).

Reciprocally, we asked if the loss of CEN2 could affect the structure of the peripheral annuli. Despite several attempts, we were unable to visualize PAP1 in CEN2-iKD parasites. Instead, we used a novel marker of the peripheral annuli, TGGT1 230340, which we identified by mass spectrometry on a biotin-based proximity-labelling (BioID) experiment using SAS6L, a protein found above the conoid³² (Dataset S1). Epitope tagging of TGGT1_230340 at the endogenous locus revealed that this protein specifically localized to the peripheral annuli and we therefore named it PAP2. PAP1 and PAP2 are largely conserved in the coccidian-subgroup of Apicomplexa (Table S1) but the two proteins do not share any obvious sequence homology. PAP1 contains two coiled-coil domains and a putative intermediate filament-binding trichohyalin-like domain with 11 repeats of a 33 amino-acid stretch (Figure S3A). PAP2 is predicted to possess armadillo repeats and a coiled-coil domain (Figure S3B,C).

PAP2 localization by STED microscopy revealed a labelling that resembles that of PAP1, with a ring-shape structure localized exclusively to the six peripheral annuli of the parasite (Figure 3F). A 3D surface reconstruction of PAP2 labelling indicated that it surrounds CEN2-YFP at each annulus, suggestive of a multiple layered structure at the peripheral annuli (Figure 3G and Movie S1). The peripheral annuli have been shown to be resistant to detergent extraction, probably because of their association with the subpellicular cytoskeleton.³¹ To test if these structures are connected with the subpellicular microtubules, we treated parasites for 24 hours with 0.5 nM oryzalin, an herbicide known to bind and inhibit plant microtubule polymerization without affecting mammalian microtubules. In T. gondii, 0.5 nM oryzalin is known to efficiently disrupt the subpellicular microtubules without causing aberrant cell division⁴⁰ (Figure 4A, Δ KU80). Under these conditions, the three markers of the peripheral annuli remained unaffected suggesting that peripheral annuli protein localization is not dependent on the subpellicular microtubules (Figure 4A).

In order to follow the peripheral annuli in the absence of CEN2, we endogenously tagged PAP2 at its C-terminus in CEN2-iKD parasites (CEN2-iKD/PAP2-3Ty). PAP2 distribution was clearly affected by the depletion of CEN2. Up to 70% of the parasites presented either a fainter PAP2 signal, an aberrant PAP2 localization, or a reduced number of PAP2 positive annuli per parasite. This contrasted with the six peripheral annuli homogenously observed in control parasites (Figure 4B). Concordantly, the level of PAP2-3Ty protein assessed by western blot, was significantly decreased upon depletion



FIGURE 3 CEN2, PAP1 and PAP2 colocalize to the annuli and form a complex structure. A, Stimulated emission depletion (STED) microscopy 3D projection of intracellular tachyzoites of *T. gondii* expressing CEN2-YFP/PAP1-3Ty. PAP1-3Ty staining is visualized using an anti-Ty antibody (red). Scale bar: 1 µm. A scheme depicts the parasite organization in the imaged vacuole. P1 and P2 = parasite 1 and parasite 2. B, 3D STED surface reconstruction of the apical part of parasite 1 (P1) showing that CEN2-YFP staining is surrounded by PAP1-3Ty staining. Scale bar: 0.4 µm. C, IFA of CEN2-YFP in wild-type and PAP1-KO parasites. GAP45 stains the periphery of the parasites. Scale bar: 2 µm. In the absence of PAP1, CEN2 still localizes to the annuli (C) and is not destabilized as demonstrated by western blot using anti-GFP antibodies and catalase (CAT) as a loading control (D). E, Plaque assay performed over 7 days with the CEN2-YFP-iKD and CEN2-YFP-iKD/PAP1-KO strains in presence or absence of ATc. No deleterious effect on the tachyzoite lytic cycle was monitored upon PAP1 deletion. F, STED microscopy 3D projection of intracellular tachyzoites expressing CEN2-YFP/PAP2-3Ty. PAP2-3Ty staining is visualized using an anti-Ty antibody (red). Scale bar: 1 µm. A scheme depicts the parasite organization in the imaged vacuole. P1-4 = parasite 1-4. G, 3D STED surface reconstruction of the apical part of parasite 3 (P3) and parasite 4 (P4) showing the CEN2-YFP surrounded by PAP2-3Ty. Scale bar: 0.4 µm

(A) (B) ∆KU80 CEN2-iKD CEN2-iKD/PAP2-3Ty - ATc CEN2-iKD/PAP2-3Ty + ATc DMSO DMSO 0.5nM oryzalin 0.5nM oryzalin tuh c-tub FN2-YF P<0.0001 100 % of vacuoles with normal PAP2 staining 80 60 40 merge merge merge merge 20 CEN2-iKD -ATc CEN2-iKD +ATc PAP1-3Ty PAP2-3Ty (C) CEN2-iKD/PAP2-3Tv 0.5nM oryzalin DMSO DMSO 0.5nM oryzalin ATc (h) 0 24 48 α-Τν α-Τν 170 130 ♦ PAP2-3Ty 100 70 55 40 35 -25. kDa 70 ▲ CAT 55 merge merge merge merge α-Τγ/α-CAT

FIGURE 4 CEN2 is critical for the architecture of the peripheral annuli. A, ΔKU80 (control), CEN2-YFP-iKD, PAP1-3Ty and PAP2-3Ty parasites have been treated with DMSO or 0.5 nM oryzalin. Oryzalin treatment disturbs the cortical microtubules as shown in the control but not the apical localization of CEN2, PAP1 or PAP2. GAP45 stains the periphery of the parasites. Scale bars: 2 μm. B, Immunofluorescence performed on intracellular CEN2-iKD parasites stably expressing the endogenously tagged peripheral annuli protein 2 (PAP2). Upon ATc treatment, PAP2 staining is affected in its intensity and localization but is not homogenous within a vacuole. The number of vacuoles exhibiting a normal PAP2 staining has been quantified and is strongly decreased in CEN2-depleted parasites. Results are presented as mean ± SD and their significance has been assessed using an unpaired *t* test. The two-tailed *P*-values are written on the graph. C, The expression and stability of PAP2 has been assessed by western blot using the anti-Ty antibodies to detect the endogenously tagged protein and anti-catalase (CAT) as a loading control

of CEN2 (Figure 4C). Despite the fact that PAP2 depletion was predicted to have a low impact on parasite fitness (fitness score of -1.14^{28}), we were unable to generate a PAP2-KO parasite line to determine if the absence of PAP2 had an impact on the localization of CEN2 or PAP1 at the peripheral annuli. In conclusion, the peripheral annuli are stable in the absence of the subpellicular microtubules, whereas the depletion of CEN2 appears to be important for the localization of PAP2.

2.3 | DLC8a is associated with tubulin-rich structures

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DLC8a has been previously reported to localize to structures that appose with structures exhibiting CEN2 labelling, suggesting that these two proteins might, in part, share a similar function. When expressed as a second copy, DLC8a localized to the polar ring, the conoid, the spindle poles, the centrosome and the basal ring.³¹ In another study, a C-terminal-tagged version of DLC8a was reported to

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FIGURE 5 DLC8a localizes upstream of the secretory organelles and is critical for tachyzoite lytic cycle. A, IFA performed on intracellular parasite expressing the inducible N-terminally tagged DLC8a (MycDLC8a). MycDLC8a is found at the apical pole of the parasites as well as at the centrosome where it partially colocalizes with CEN1 and CEN2. Costaining of MycDLC8a with markers of the micronemes (MIC6) and rhoptries (ARO) revealed that MycDLC8a localization is upstream of these secretory organelles. IFAs with CEN1 and MIC6 have been fixed with cold methanol while IFAs with CEN2 and ARO have been fixed with PFA/GA. Scale bars: 2 μ m. B, In extracellular parasites, MycDLC8a staining is found at the apical end of the conoid in nonstimulated conditions and covering the conoid in protruded conditions (+A23187). IMC1 stains the periphery of the parasite. Scale bars: 1 μ m. C, Immunofluorescence of Myc-DLC8a-iKD parasites treated either with DMSO or 0.5 nM oryzalin for 24 hours showing that DLC8a remains apical during cortical microtubule depolymerization. Scale bars: 2 μ m. D, E, The regulation of MycDLC8a-iKD was assessed by IFA after 48 hours ± ATc (D) and by western blot (E) using anti-Myc antibodies. MycDLC8a reached undetectable levels after 48 hours of ATc treatment. Scale bars: 2 μ m. F, Plaque assay performed over 7 days with the parental (Δ KU80) and the MycDLC8a inducible (MycDLC8a-iKD) strains in the presence or absence of ATc. MycDLC8a-iKD parasites do not form plaques in the presence of ATc confirming a strong growth defect. Scale bars: 0.5 cm. G, Intracellular growth assay performed with Δ KU80 and MycDLC8a-iKD after a 60 hours pretreatment ± ATc. No significant defect was noticed between the parental and the mutant strains. Results are presented as mean ± SD

localize solely at the apical cap.³⁸ To investigate DLC8a function, we generated an inducible knockdown cell line, MycDLC8a-iKD, by exchanging the endogenous promoter with the ATc regulatable promoter that expressed an N-terminal Myc epitope-tagged DLC8a protein (Myc-DLC8a; Figure S4A,B). In intracellular parasites, Myc-DLC8a localized to the parasite centrosome and at the apical cap with a stronger signal at the APR (Figure 5A). At the centrosome, DLC8a colocalized with the CEN1 and CEN2 labelling (Figure 5A–top panels). At the apical pole, Myc-DLC8a was detected at a site more apical than the micronemes and the rhoptries (Figure 5A–lower panels). In extracellular parasites, the protein localized on the top of the retracted conoid, while upon treatment with the Ca²⁺ ionophore A23187 (an inducer of conoid protrosion), Myc-DLC8a labelling was detected on the entire extruded conoid (Figure 5B) as previously reported by immuno-electron microscopy.³¹

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In intracellular parasites treated with 0.5 nM oryzalin, Myc-DLC8a was still detected at the apical cap of the parasite, indicating that its localization is unaffected by the absence of the subpellicular microtubules (Figure 5C).

2.4 | TgDLC8a drives microneme secretion, required for parasite attachment

As with CEN2, assessing the functional roles of DLC8a is challenging because it resides at multiple sites and thus several phenotypes might overlap. In MycDLC8a-iKD parasites, Myc-DLC8a levels were undetectable after 48 hours of ATc treatment using both IFA and western blot (Figure 5D,E). Parasites depleted of DLC8a generated smaller lysis plaques than control parasites, indicative of a defect in one or multiple steps of the lytic cycle (Figure 5F). The role of DLC8a



FIGURE 6 DLC8a is critical for microneme secretion. A, Red/green invasion assays performed with freshly egressed Δ KU80 and MycDLC8a-iKD parasites after a 60 hours pretreatment ± ATc. DLC8a depleted parasites are severely impacted in their invasion capacity. B, Microneme secretion assay carried out with freshly egressed Δ KU80 and MycDLC8a-iKD parasites pretreated for 60 hours ± ATc and analyzed by western blot. Neither ethanol nor BIPPO induce MIC2 secretion in DLC8a-depleted parasites. GRA1 attests to the constitutive secretion of dense granules. ESA, excreted secreted antigens; pMIC2, processed MIC2. C, IFA performed on intracellular parasites treated ± ATc for 60 hours to assess the integrity and positioning of the micronemes. No defect was observed for the two proteins MIC2 and MIC6. Scale bars: 2 µm. D, Attachment assay performed with freshly egressed Δ KU80 and MycDLC8a-iKD parasites pretreated for 60 hours ± ATc. A drop in the attachment efficiency of the parasites depleted in MycDLC8a can be observed. E, Induced egress assay performed on parasites treated for 60 hours ± ATc. C, BIPPO-stimulated condition. F, Gliding assay performed by video-microscopy on parasites pretreated for 60 hours ± ATc. No defect in motility was recorded. For A, D, E and F, results are presented as mean ± SD. The significance of the results has been assessed using an unpaired *t* test and the two-tailed *P*-values are written on the graphs

at the centrosomes was evaluated by scoring the number of parasites per vacuole after 60 hours of ATc treatment. In this assay, no delay in

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replication was observed (Figure 5G).

In contrast, DLC8a plays a critical role at the apical pole of the parasite, with less than 10% of DLC8a-depleted parasites being able to invade compared to the control (Figure 6A). To further dissect the phenotype, microneme secretion was assayed following triggering by either ethanol or BIPPO, a phosphodiesterase inhibitor.⁴¹ DLC8adepleted parasites failed to secrete MIC2 above basal levels in response to both stimulations (Figure 6B). This defect was also observed for the secretion of two other microneme proteins. AMA1 and MIC8 (Figure S4C). However, upon DLC8a depletion, the micronemes were still properly positioned at the apical pole, suggesting a role for DLC8a in the cascade of events leading to microneme exocytosis and not in the trafficking or apical positioning of the organelles (Figure 6C). The basal level of microneme secretion observed in DLC8a-depleted parasites seemed insufficient to promote efficient parasite attachment and invasion into the host cell (Figure 6D). although it supported egress and motility (Figures 6E,F and S4D). Overall, depletion of DLC8a strongly impacts invasion and microneme secretion but only moderately affects motility, egress and attachment.

2.5 | Depletion of TgDLC8a results in aberrant positioning of the rhoptries impacting on their discharge

DLC8a-depleted parasites exhibited a severe invasion defect, with motility and egress being unexpectedly normal. This suggested an additional role for DLC8a in rhoptry function. In these parasites, IFAs performed with antibodies to the rhoptry membrane marker, ARO, demonstrated a typical apical signal for the rhoptries, but also revealed the unexpected presence of filamentous structures all over the parasite body—as previously reported in ARO-depleted parasites¹⁶ (Figure 7A). This abnormal staining for the rhoptries was confirmed with anti-RON9 and anti-ROP7 antibodies. ROP7 labelling showed a similar filamentous pattern as ARO labelling, whereas RON9 appeared more punctiform. Both ROP7 and RON9 overlap with ARO, indicating that they are still associated with rhoptry material despite the fact that the rhoptry subcompartments cannot be clearly delineated.

The ultrastructure of the DLC8a-depleted parasites in several vacuoles was analyzed by transmission electron microscopy (TEM) (Figure S5A-C) and focused ion beam milling combined with scanning electron microscopy (FIB-SEM) (Figure 7B-D and Movie S2). No noticeable defect on the morphology of the apical pole was observed. The conoid appeared normal with rhoptries properly docked to the apex (Figure S5A,B) and the typical crown of micronemes at its base (Figure S5C). However, in several sections, some mature rhoptries did not cluster with the apical ones but rather were dispersed in the cytoplasm (one to five rhoptries per parasite in the serial sections analyzed) (Figures 7B-D and S5B, yellow-colored letters). FIB-SEM serial sections were used to perform 3D reconstruction of one representative DLC8a-depleted parasite (Figure 7E). This 3D image confirmed that several mature rhoptries were dispersed within the parasite, while others were properly clustered apically with one docked into the conoid.

Considering the presence of apically docked rhoptries and yet the severe invasion defect observed in DLC8a-depleted parasites, we investigated the ability of the rhoptries to discharge their content. ROP16 is a parasite effector protein secreted into the host cell to modulate the host immune response by phosphorylating the protein STAT6.42 Here, the secreted ROP16-dependent level of STAT6 phosphorylation (STAT6-P) was used to quantify rhoptry discharge. As DLC8a-depleted parasites already presented an attachment defect, STAT6-P positive cells were normalized to the number of attached parasites instead of the number of host cell nuclei. In the absence of DLC8a, the parasites lost 50% of their ability to discharge the rhoptries compared to control parasites (Figures 7F). Distinctly, DLC8a-depleted extracellular parasites showed no defect in BIPPOstimulated conoid protrusion (Figure S5D). Taken together these results highlight a dual role of DLC8a in stimulating microneme exocytosis and in proper apical positioning of the rhoptries, both of which are critical for successful host cell invasion.

2.6 | CEN2 and DLC8a are distinct mediators of organelle exocytosis

We showed that CEN2 and DLC8a were found in apposed structures and that both proteins were involved in microneme exocytosis. This led us to investigate a potential interaction between them. We endogenously tagged CEN2 in MycDLC8a-iKD parasites (CEN2-YFP/-MvcDLC8a-iKD). Depletion of DLC8a following ATc treatment did not impact the localization (Figure 8A) or the expression level of CEN2-YFP (Figure 8B). Reciprocally, we transiently expressed eGFP-DLC8a³¹ in CEN2-iKD parasites. In absence of CEN2-YFP (following ATc treatment), the level of eGFP-DLC8a remained unaffected (Figure 8C). To rule out any physical interaction between CEN2 and DLC8a, we performed coimmunoprecipitation followed by mass spectrometry analysis of either CEN2-YFP or Myc-DLC8a immunoprecipitates (Dataset S2). In these experiments, CEN2 was not found in Myc-DLC8a immunoprecipitates and DLC8a was not found in CEN2-YFP immunoprecipitates. These experiments confirmed that the two proteins do not directly interact and likely act independently on the cascade of events leading to microneme exocytosis. Relevantly, PAP2 was found in the mass spectrometry results of CEN2-YFP pull-down, supporting our previous observations and suggesting that the two proteins are associated at the peripheral annuli (Figure 4B,C).

3 | DISCUSSION

In this study, we dissected the roles of DLC8a and CEN2, two proteins conserved across the Apicomplexa that are linked to several microtubule-based structures (Table S1). We showed that both proteins fulfill multiple, essential tasks in *T. gondii* parasites. While DLC8a deletion was achieved with concomitant disappearance of the protein from the centrosomes, the apical cap and the polar ring, the

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FIGURE 7 Mature rhoptries are found dispersed in the cytoplasm upon DLC8a depletion. A, IFA on intracellular parasites treated \pm ATc for 60 hours to assess the integrity and positioning of the rhoptry. In MycDLC8a-depleted parasites, the staining of the rhoptry surface protein ARO appears elongated, the staining of the rhoptry neck protein RON9 appears more punctate as well as the pattern of rhoptry bulb protein ROP7. Scale bars: 2 µm. B-D, A parasitophorous vacuole of the MycDLC8a-iKD cell line treated for 78 hours with ATc was imaged by FIBSEM. Selected slices throughout the imaged volume are presented and show tachyzoites with disperse rhoptrie localizations (yellow r). C and D are images 590 and 750 nm deeper relative to image B. c, conoid; d, dense granules; m, mitochondrion; n, nucleus; r, rhoptry; tvn, tubulovesicular network; scale bars: 1 µm. E, 3D model of the tachyzoite highlighted with asterisk in B-D. Tachyzoite plasma membrane is in cyan, conoid in black, nucleus in light blue and rhoptries in orange. F-G, The ability of the Δ KU80 and MycDLC8a-iKD parasites to secrete their rhoptries was assessed using a Phospho-STAT6 test. It revealed a significant defect in the ROP16 secretion for the MycDLC8a depleted parasites. ARO-iKD parasites were used as a control because they present dispersed rhoptries under ATc treatment. F, Left panel: representative IFA of HFF monolayers infected with ATc-treated parasites. Parasites were stained with an anti-ACT1 (actin) antibody, host cell nuclei were stained with DAPI and cells in which rhoptry have been discharged are stained with an anti-STAT6-P antibody. Right panel: graph representing the percentage of STAT6 positive nuclei associated with a parasite. Results are presented as mean \pm SD. The significance of the results has been assessed using an unpaired t test and the two-tailed *P*-values are written on the graph



FIGURE 8 CEN2 and DLC8a are two independent actors of organelle exocytosis. A, IFA on intracellular parasites MycDLC8a-iKD expressing CEN2-YFP showing that depletion of DLC8a with ATc does not affect CEN2-YFP labelling. Scale bars: 2 μ m. B, Western blot analysis of CEN2-YFP level (using anti-GFP antibodies) in CEN2-YFP/MycDLC8a-iKD parasites in untreated or ATc-treated parasites. Catalase (CAT) is used as a loading control. C, Western blot analysis of eGFP-DLC8a level in either Δ KU80 or CEN2-iKD parasites transiently transfected with eGFP-DLC8a and treated \pm ATc for 48 hours. The level of eGFP-DLC8a remains the same even when CEN2-YFP is downregulated by the addition of ATc

knockdown of CEN2 gene expression resulted in a stepwise disappearance of the protein from its multiple locations. The pools of CEN2 associated with the preconoidal rings and the peripheral annuli were the first to be depleted, whereas depletion of CEN2 from the basal cup and the centrosomes occurred later. The differential kinetics of CEN2 depletion illustrates the peculiar mechanism of division of *T. gondii*, with the apical complex made de novo for the two daughter cells at each division cycle, while the centrosome is a highly stable structure that duplicates and is inherited from the mother.⁴³ The pool of CEN2 associated with the basal cup, a structure also made de novo for each daughter cell, is lost at an intermediate rate. The fact that CEN2 at this location disappears later than in the other *de novo* structures could be partly due to a recycling process from the basal pole of the mother, as previously shown for IMC proteins.⁴⁴

The parasites depleted of CEN2 divide normally, suggesting that either the residual amount of CEN2 observed at the centrosome is sufficient to support division or that CEN2 carries a nonessential function at this location. In favor of the latter hypothesis, the presence of two other centrins, CEN1 and CEN3, which localize to the centrosome³¹ and are likely essential for tachyzoite survival,²⁸ might assure proper eukaryotic chromosome segregation. Both CEN2 and MyoJ are involved in the posterior constriction of the parasites, while MyoI is responsible of the intravacuolar connection.³⁵ In contrast to MyoJ/MyoI depletion, the absence of CEN2 does not impact the intravacuolar connection, because CEN2-depleted parasites still divide synchronously. Consequently, the intravacuolar connection is independent of the basal constriction but requires the presence of MyoJ and MyoI.

Remarkably, both CEN2 and DLC8a are implicated in microneme exocytosis, although the phenotypic consequences of their depletion are distinct. CEN2 ensures sufficient microneme secretion to support motility, attachment, invasion and egress. Microneme exocytosis likely occurs after the passage of the organelles through the conoid, and we postulate a role for CEN2 at the preconoidal rings in this process. Alternatively, CEN2 could play a more active role either by triggering the contraction of the preconoidal rings that might be needed for microneme secretion or, as a phosphoprotein, by functioning in the signaling cascade leading to microneme exocytosis⁴⁵ without affecting conoid protrusion. In this context, centrins have been reported in *Paramecium* to buffer Ca²⁺ and to downregulate the signal after stimulation of exocytosis.⁴⁶ Additionally, in vertebrate photoreceptor cells, a centrin complex is believed to block protein and small molecule passage in the presence of Ca^{2+.47}

In contrast to CEN2 depletion, the absence of DLC8a impacts microneme secretion in a subtler manner, because motility, attachment and egress were not abrogated. While the existence of at least two distinct subpopulations of micronemes has been reported,48 DLC8a-depleted parasites are uniformly unable to secrete MIC2, AMA1 and MIC8, ruling out a selective role of DLC8a on one subpopulation of organelles. Microneme secretion occurs in waves in response to fluctuation in intracellular Ca²⁺, suggesting that small groups of micronemes might be transferred sequentially to the conoid.⁸ In accordance with this model, subsets of micronemes are localized underneath the APR of the parasite while others are found more basally. DLC8a-depleted parasites appear unable to sustain the successive waves of exocytosis that may be necessary to support the ultimate step of the entry process. In such a scenario, a population of micronemes that are already apically localized and primed for secretion would be secreted, while microneme replenishment for subsequent secretion would be impaired due to a defect in peripheral microneme transport along the microtubules. Among the dynein heavy chains found in T. gondii, TGME49_294550 is present in the genome of all the apicomplexans (Table S1) and might be involved in this task, as this family is known to be involved in the long distance transport of vesicles and organelles.49

Remarkably, DLC8a depletion also affected the apical positioning of the rhoptries, leading to a few mis-positioned organelles disseminated in the cytoplasm. Although some rhoptries were still apically positioned with their neck properly inserted into the conoid, organelle discharge and invasion were significantly impaired. The invasion phenotype might result from an insufficient number of properly positioned rhoptries, given that the parasite is known to discharge rhoptries in uninfected-host cells prior to successful invasion.⁵⁰ Alternatively, the impairment in rhoptry discharge could result from a microneme exocytosis defect at the time of invasion, possibly due to the absence of MIC8, which was previously reported to be important for rhoptry discharge.⁵¹

CEN2 is a marker for the lower boundary of the apical cap that is delineated by a ring of approximately six annuli. These peripheral annuli form a mysterious structure of unassigned function. The annuli have been described only in T. gondii, but are likely to be present in the coccidian subgroup of Apicomplexa. We report here a new marker of this structure, PAP2, which was identified by mass spectrometry during biotin-based proximity-labelling (BioID) of SAS6L and shown here to be associated with CEN2. It is found in most coccidians (Dataset S1). The previously described PAP1 protein³⁴ is conserved in some coccidian parasites (Table S1) and possesses a trichohyalin-like domain in addition to two coiled-coil domains. Trichohyalin is an intermediate filament-associated protein made of two EF-hand domains and a 28 aa-sequence repeat region⁵² comparable to the one we observed in PAP1. Interestingly, a network of intermediate filamentlike proteins, the alveolins, underlies the cytoplasmic face of the IMC and might interact with PAP1.⁵³ Supporting this hypothesis, PAP1 has been found in the proteome of the subpellicular cytoskeleton of T. gondii⁵⁴ and the annuli are resistant to oryzalin-mediated microtubule depolymerization. Regarding PAP2, its sequence is predicted to include ARM repeats and a coiled-coil domain. Comparison of aminoacid sequences between PAP1 and PAP2 reveals no similarity. However, in silico protein modeling (Phyre2 server) suggests that both proteins share structural similarities with their central core forming a double stranded coiled-coil rod domain usually found in intermediate filament proteins.

Super-resolution microscopy revealed that peripheral annuli exhibit a complex and organized structure with distinct protein content. More precisely, CEN2-YFP appears as a dot surrounded by a ring of PAP1 and PAP2 proteins. In the absence of CEN2, we observed that PAP2 was lost and degraded. Currently, the peripheral annuli can only be observed by IFA, hence we cannot formally conclude that loss of annuli-associated proteins leads to the loss of the underlying structure. PAP1 and PAP2 are the only two specific markers of this structure and are predicted to be dispensable for tachyzoite survival.²⁸ Nevertheless, the PAP2 locus seems refractory to disruption despite several attempts using two different knockout strategies. The annuli have been hypothesized to participate in endocytosis or in dense granule secretion; however, no defect in dense granule secretion was observed in CEN2-depleted parasites or in PAP1-KO parasites (data not shown). The multiple roles of CEN2 and the concomitant disappearance of PAP2 from the annuli, together with the inability to obtain PAP2-KO parasites, hampers a functional assignment for these structures. In this context, we cannot formally conclude that the loss of CEN2 at the apical tip per se is responsible for the observed phenotype. Further investigation will be needed to unravel the contribution of these enigmatic structures.

In conclusion, we report the role of two conoidal proteins, CEN2 and DLC8a, in microneme secretion that do not appear to play a role in conoid protrusion. However, the two proteins seem to act at different steps along the signaling cascade leading to microneme exocytosis. These findings support the current model that microneme secretion is dependent on Ca²⁺ pulses, while microtubules provide a track for the transport of these organelles to the conoid as suggested by the localization of DLC8a in extracellular protruded parasites. In addition, our study suggests a role for a microtubule-associated complex in rhoptry docking at the apical pole of the parasite. Importantly, both CEN2 and DLC8a might fulfill similar functions across the phylum as they do in *T. gondii*.

4 | MATERIALS AND METHODS

4.1 | Molecular cloning

The sequence of the primers used for the cloning can be found in Table S2.

4.1.1 | Myc-DLC8-iKD (inducible knock-down)

To generate the Tet-repressive knockdown of DLC8a, a PCR fragment encoding the TATi trans-activator, HXGPRT selection cassette, the TetO7S1 promoter and the c-Myc tag was generated using the KOD DNA polymerase (Novagen, Merk) with the vector 5'COR-pT8TATi1-HX-tetO7S1myc⁵⁵ as template and the primers DLC8a-7214/DLC8a-7215 that also carry 30 bp homology with the 5' end of DLC8a. To direct the insertion of the PCR product at the start of DLC8a locus, a specific sgRNA vector has been generated using the Q5 site-directed mutagenesis kit (New England Biolabs), the vector pSAG1::CAS9-GFPU6::sgUPRT as template⁵⁶ and the primer pair DLC8a-7378/gRNA-4883.

4.1.2 | PAP1-3Ty-HXGPRT (knock-in)

To generate the PAP1-3Ty-HXGPRT vector, a gDNA fragment corresponding to the C-terminus of PAP1 (TGGT1_242790A and B) was amplified by PCR using the primer pair PAP1_5505/PAP1_5506 and cloned into the *Kpn*I and *Nsi*I sites of the pTUB8-MIC13-3Ty-HX.⁵⁷

4.1.3 | PAP2-3Ty-DHFR (knock-in)

To generate the PAP2-3Ty-DHFR vector, a gDNA fragment of the C-terminal part of PAP2 (TGGT1_230340) was amplified by PCR using the primers PAP2_6308/PAP2_6309 and cloned into the *Apal* and *Nsil* sites of the Ct-ASP5-3Ty-DHFR.⁵⁸

4.1.4 | PAP1-KO-DHFR (knockout)

To generate the PAP1-KO, a PCR amplicon of the DHFR resistance cassette was generated using the KOD DNA polymerase (Novagen,

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Merk), the vector p2854-DHFR⁵⁹ as template and the primers PAP1_8995/PAP1_8996 that encode 30 bp overhangs corresponding to the start and the 3' end of PAP1. A double CRISPR-Cas9 guide strategy was used to drive the insertion of the amplicon into the PAP1 locus. Two specific gRNAs were made as previously described. Then a PCR amplicon of one guide amplified using the primers 2nd-gRNA-F_6147/2nd-gRNA-R_6148 was subcloned into the other guide between the *KpnI* and *XhoI* restriction sites, thereby generating a dgRNA plasmid with two targeted sgRNAs.

4.2 | Parasite culture and transfection

The *T. gondii* Δ KU80 RH strain was used throughout this study and tachyzoites were propagated in confluent Human Foreskin Fibroblasts (HFFs) with Dulbecco modified Eagle's medium supplemented with 5% fetal bovine serum. Transfection of tachyzoites were performed as previously described⁶⁰ either with 15 to 20 µg of linearized plasmid for endogenous tagging or with a mix of 15ug Cas9-sgRNA plasmid and 10 to 20 µg of PCR product. Clonal populations were isolated by limited dilution and the presence of the recombinant locus was verified by PCR performed on genomic DNA with the primers outlined in Table S3, Figures S2 and S4 and the GoTaq polymerase (Promega). For the transient expression of eGFP-DLC8a, 100 µg of pmin-eGFP-DLC8a plasmid³¹ was transfected into CEN2-iKD. Immediately after the electrical pulse, the transfected parasites were split into two dishes containing HFF either in the absence or in the presence of ATc. Extracellular parasites were collected 2 days post-transfection.

4.3 | Immunofluorescence assay

For IFA, parasite-infected HFF cells coated on coverslips were fixed with 4% paraformaldehyde (PFA)/0.05% glutaraldehyde (GA) in PBS for 10 to 12 minutes (for most of the antigen to be labeled) or with cold methanol for 8 minutes at -20° C (for α -Myc to detect MycDLC8a and for α -acetylated tubulin) and then processed with and without Triton X-100 respectively, as previously described.²⁰ Confocal and STED microscopy images were collected with a confocal laser scanning microscope LSM700 (Zeiss) and a TCS SP8 STED ×3 microscope (Leica) respectively, at the Bioimaging Core Facility of the University of Geneva Medicine Faculty. For confocal data, the images were analyzed using ImageJ and the z-stacks were projected using the maximum projection tool. For the STED microscopy, 3D reconstruction and projection were performed using Imaris 9.0 (Bitplane) and LasX (Leica) softwares, respectively. Some images were contrast enhanced for figure presentation.

4.4 | Western blot

For immuno-blot analysis, pellets from extracellular tachyzoites were resuspended in PBS buffer, mixed with loading buffer under reducing condition (DTT), sonicated in a bioruptor (diagenode) and then boiled for 10 minutes before loading onto a SDS-PAGE gel. After running, wet-transfer was performed onto nitrocellulose membranes before probing with appropriate antibodies in 5% nonfat milk powder in PBS-0.05% Tween20. Bound secondary conjugated antibodies were visualized using the ECL system (for horseradish peroxidase) or detected directly (for the Alexa-Fluor-680-conjugated).

4.5 | Plaque assay

Freshly egressed parasites were used to infect confluent HFF cells. These plates were incubated for 7 days \pm ATc and then fixed with PFA/GA and stained with a crystal violet solution (Sigma).

4.6 | Intracellular growth assay

Parasites pretreated for 24 hours (Δ KU80 and CEN2-iKD) or 36 hours (Δ KU80 and DLC8a-iKD) ± ATc were allowed to grow on new confluent HFFs for 24 hours ± ATc before fixation with PFA/GA. IFA was then performed using anti-GAP45 antibodies and the number of parasites per vacuole was counted in 100 vacuoles from three independent experiments. The results are presented as mean ± SD and the significance of the results has been assessed using an unpaired *t* test.

4.7 | Red/green invasion assay

Parasites pretreated \pm ATc for 48 hours (Δ KU80 and CEN2-iKD) or 60 hours (Δ KU80 and DLC8a-iKD) were used to perform the invasion assay as previously described.⁶¹ The number of extracellular and intracellular parasites was determined by counting 100 parasites in duplicate for three independent experiments. Results are presented as mean \pm SD and the significance of the results was assessed using an unpaired *t* test.

4.8 | Induced egress assay

Parasites were pretreated for 18 hours (Δ KU80 and CEN2-iKD) or 30 hours (Δ KU80 and DLC8a-IKD) ± ATc. Freshly egressed parasites were then allowed to grow on new confluent HFFs for 30 hours ± ATc before adding either BIPPO (50 μ M), A23187 from *Streptomyces chartreusensis* (3 μ M) or DMSO for 8 minutes. IFAs were performed using α -GRA3 to stain the parasitophorous vacuole and α -GAP45 to stain the parasites. The average number of egressed vacuoles was determined by counting at least 100 vacuoles in duplicate for each condition and for three biological experiments. The results are presented as mean ± SD and the significance of the results has been assessed using an unpaired *t* test.

4.9 | Host cell attachment assay

Extracellular parasites expressing GFP and mutant cell lines were pretreated \pm ATc for 48 hours (Δ KU80 and CEN2-iKD) or 60 hours (Δ KU80 and DLC8a-iKD) and then mixed at a 50/50 ratio. The parasite mixture was added to HFF-coated coverslips ("assay") and centrifuged for 1 minute at 1000 rpm, washed once and then fixed with PFA/GA for 10 minutes. In parallel, an aliquot of this mixture was stained with Hoechst for 20 minutes and fixed with PFA/GA for 10 minutes to determine the initial ratio on a Beckman Coulter Gallios flow cytometer ("control"). For the "assay," immunofluorescence (IFA) was performed using α -GAP45 antibodies to determine the ratio of attached parasites. The experiments have been performed in triplicates, 1000 parasites were counted by flow cytometry and 100 parasites by IFA. Percentage of GFP positive (internal control) and negative parasites (strain of interest) were calculated for both flow cytometry and microscopy methods. Efficiency of attachment was measured by calculating the ratio between the proportion of parasites attached to the host cell ("assay" %) and the proportion of parasite that were initially present in the mix ("control" %).

4.10 | Microneme secretion assay

Freshly egressed parasites pretreated ± ATc for 48 hours (Δ KU80 and CEN2-iKD) or 60 hours (Δ KU80 and DLC8a-iKD) were used to performed microneme secretion assay with 2% ethanol or 50 μ M BIPPO as previously described.²⁰ Secretion was assessed by western blot.

4.11 | Rhoptry secretion assay

A phospho-STAT6 test was used to measure the ability of parasites to successfully secrete rhoptry proteins into the host cells. Briefly, freshly egressed DLC8-iKD parasites pretreated \pm ATc for 60 hours were resuspended in cold DMEM at a concentration of 20.10⁵ parasites/mL and 5.10^5 parasites were used to infect HFF-coated coverslips. The plates were centrifuged 30 seconds at 1100 g to allow parasites to attach and incubated for 20 minutes on ice to synchronize the invasion. Cells were then incubated at 37°C for 20 minutes and fixed with ice-cold methanol for 8 minutes at -20° C. Following blocking, immuno-detection was done using anti-PSTAT6 (Cell signaling 9361; 1/400) and the number of parasites associated to a PSTAT6 positive nucleus was determined. The experiments were done in triplicate and 100 parasites were counted each time.

4.12 | Induced gliding assay

Motility was performed by video microscopy on a Nikon eclipse Tiinverted microscope at 37°C with either Δ KU80 and CEN2-iKD parasites pretreated for 48 hours ± ATc or Δ KU80 and DLC8a-iKD parasites pretreated for 60 hours ± ATc as previously described.⁶² The type of gliding was evaluated in 100 parasites, in three biological replicates. The results are presented as mean ± SD and the significance of the results has been assessed using an unpaired *t* test. Tracking of parasites was performed using the Manual Tracking plugin on ImageJ software.

4.13 | Electron microscopy

For transmission electron microscopy, the *T* gondii infected HFF cells were grown as monolayers on 13 mm round glass coverslips. After 78 hours treatment with ATc, cells were fixed with 2.5%

glutaraldehyde (Electron Microscopy Sciences) and 2% paraformaldehyde (Electron Microscopy Sciences) in 0.1 M phosphate buffer (PB) at pH 7.4 for 1 hour at room temperature. Cells were then washed 5×5 minutes with 0.1 M sodium cacodylate buffer, pH 7.4 and postfixed with 1% osmium tetroxide (Electron Microscopy Sciences) and 1.5% potassium ferrocyanide in 0.1 M sodium cacodylate buffer, pH 7.4 for 1 hour immediately followed by 1% osmium tetroxide (Electron Microscopy Sciences) in 0.1 M sodium cacodylate buffer pH 7.4 for 1 hour. Coverslips were then washed in double distilled water 2×5 minutes and *en block* stained with aqueous 1% uranyl acetate (Electron Microscopy Sciences) for 1 hour. After 5 minutes wash in double distilled water cells were dehydrated in graded ethanol series $(2 \times 50\%, 70\%, 90\%, 95\%$ and $2 \times$ absolute ethanol) for 3 minutes each wash. Dehydrated cells were infiltrated with Durcupan resin (Electron Microscopy Sciences) diluted with ethanol at 1:2, 1:1, 2:1 for 30 minutes each, and twice with pure Durcupan for 30 minutes each. Fresh Durcupan resin was added for additional 2 hours. Coverslip with grown cells faced down was placed on 1 mmthick silicone ring filled with fresh resin which was placed on a glass slide coated with mold-separating agent (Gloorex). Finally, samples were polymerized at 65°C overnight in oven until resin was cured. The glass coverslip was removed from the resin disk by immersing alternately into hot water and liquid nitrogen.

Targeted parasitophorous vacuole inside HFF cells was marked on the resin surface using laser microdissection microscope (Leica Microsystems). Under stereo-microscope, marked area was cut out from the disk and glued with superglue to a blank resin block. Leica Ultracut UCT microtome (Leica Microsystems) and a glass knife were used to trim the cutting face. Seventy nanometer ultrathin serial sections were cut with a diamond knife (DiATOME) and collected onto 2 mm single slot copper grids (Electron Microscopy Sciences) coated with Formvar plastic support film.

Sections were examined with Tecnai 20 TEM (FEI) operating at an acceleration voltage of 80 kV and images were collected by sidemounted MegaView III CCD camera (Olympus Soft-Imaging Systems) controlled by iTEM acquisition software (Olympus Soft-Imaging Systems).

4.14 | Focused ion beam scanning electron microscopy and 3D reconstruction

Sample for FIBSEM imaging were prepared as described above. Either whole resin block or large cut out area, containing the region of interest marked by laser microdissection microscope (Leica Microsystems) was glued onto a flat SEM stub with superglue and on each side of the block the silver conductive paste was applied. The mounted sample was gold coated with 20 nm thick layer of gold.

The imaged volume was acquired by FEI Helios NanoLab G3 UC DualBeam microscope (FEI). Ion beam was used in conjunction with a gas injection system to deposit a thick (\sim 1.5 µm) layer of platinum on the top surface of the sample above the region of interest to reduce the FIB milling artifacts. The imaging surface was exposed by creating

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the front trench and subsequently two side trenches were created using 21 nA of focused ion beam current at 30 kV voltage.

AutoSlice and View G3 software (FEI) was used to acquire the serial SEM images. Focused ion beam at current of 2.5 pA and 30 kV of acceleration voltage was applied to mill 10 nm layer from imaging face and freshly exposed surface was imaged with back scattered electron beam current of 400 pA at acceleration voltage of 2 kV, the dwell time of 9 µs/pixel and at the resolution of 4 nm/pixel. The size of the final imaged volume in *x*-, *y*- and *z*-dimension was $5751 \times 1503 \times 810$ pixels corresponding to $23 \times 6 \times 8.1$ µm for DLC8a-iKD (+78 hours ATc). The final volume of cropped parasit-ophorous vacuole was 10.99 × 6 × 8.1 µm.

Serial images were combined into single image stacks and aligned using the FIJI program (fiji.sc/). After alignment, images were scaled down to have volume with isotropic voxel properties of the 10 nm/pixel in all x-, y- and z-dimension.

Three-dimensional reconstruction was generated by semiautomated approach using Ilastik software (ilastik.org) and 3D model was visualized using the Blender program (v.2.79; blender.org).

4.15 | Antibody list

Primary antibodies used in this study: monoclonal mouse α -Ty (hybridoma BB2, 1:10 IFA, 1:10 WB or ascyte 1:1000 IFA⁶³), α -Myc (hybridoma mAb 9E10, 1:10 IFA, 1:10 WB), α-GFP (Roche, 1:1000 WB), α-actin (hybridoma, 1:10 IFA, 1:10 WB,⁶⁴ α-MIC2 (hybridoma, generous gift from Dr V. carruthers, 1:10 IFA, 1:10 WB), α-ISP1 (1:1000 IFA,⁶⁵ a generous gift from Dr P. Bradley), α -GRA1 (1:3000 WB, Anawa), α-ROP7 T4-3H1, α-RON9 (hybridomas, generous gift from Dr J-F. Dubremetz, 1:10 IFA, 1:10 WB) and polyclonal rabbit α-catalase (1:2000,⁶⁶), α-CEN1 (1:1000 IFA, kerafast), α-ARO (1:1000 IFA,15), α-GAP45 (1:10.000 IFA,67), α-IMC1 (1:1000 IFA,61), α-MIC6 (1:1000 WB,⁶⁸) and α-PSTAT6 (1:400 IFA, Cell signaling 9361). Secondary antibodies used in this study: anti-mouse and anti-rabbit HRP (Sigma), Alexa-Fluor-680-conjugated goat anti-rabbit and anti-mouse IgG antibodies, Alexa-Fluor-488-conjugated goat anti-rabbit and antimouse IgG antibodies and Alexa-Fluor-594-conjugated goat antirabbit and anti-mouse IgG antibodies (Thermofisher).

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CONFLICT OF INTEREST

The authors declare no conflict of interest.

AUTHOR CONTRIBUTIONS

D.S.-F., K.F. and G.L. conceived the project. K.F., G.L., D.J.D. and B.M. designed, performed and interpreted the experimental work. D.S.-F. and K.F. supervised the research. G.L., D.J.D., K.F. and D.S.-F. wrote the paper.

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SUPPORTING INFORMATION

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