Title:

An Alveolata secretory machinery adapted to parasite-host cell invasion

Authors

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Apicomplexa are unicellular eukaryotes and obligate intracellular parasites, including *Plasmodium*, the causative agent of malaria and *Toxoplasma*, one of the most widespread zoonotic pathogens. Rhoptries, one of their specialized secretory organelles, undergo regulated exocytosis during invasion\(^1\). Rhoptry proteins are injected directly into the host cell to support invasion and subversion of host immune function\(^2\). The mechanism by which they are discharged is unclear and appears distinct from those in bacteria, yeast, animals or plants. Here we show that rhoptry secretion in Apicomplexa shares structural and genetic elements with the exocytic machinery of ciliates, their free-living relatives. Rhoptry exocytosis depends on intramembranous particles in the shape of a rosette embedded in the plasma membrane of the parasite apex. Formation of this rosette requires multiple Non-discharge (Nd) proteins conserved and restricted to Ciliata, Dinoflagellata, and Apicomplexa, that together constitute the superphylum Alveolata. We identified Nd6 at the site of exocytosis in association with an apical vesicle. Sandwiched between the rosette and the tip of the rhoptry, this vesicle appears as a central element of the rhoptry secretion machine. Our results describe a conserved secretion system that was adapted to provide defense for free-living unicellular eukaryotes and host cell injection in intracellular parasites.

Apicomplexan parasites are invasive and defined by the presence of an apical complex used to recognize and gain entry into host cells. It includes two secretory organelles: micronemes and rhoptries\(^3\). Microneme proteins are secreted to the parasite surface and mediate motility, host cell recognition and invasion\(^4\). Rhoptry proteins are injected directly into the host cell\(^2\), where they anchor the machinery propelling the parasite into the host cell\(^5\), facilitate nutrient
import interfer with the immune response, and modulate gene expression to promote infection. Rhopty secretion requires a trigger in the form of microneme proteins binding to host cell receptors. Importantly, rhopty proteins not only cross the plasma membrane of the parasite (exocytosis) but also that of the host. Rhopty exocytosis factors identified to date are mostly specific to Apicomplexa, suggesting that these cells depend on unique lineage-restricted secretory mechanisms. Consistent with this, no eukaryotic SNAREs, the main drivers for fusing vesicles to target membrane in eukaryotic system, have so far been associated with rhopty exocytosis. Apicomplexa also lack genes encoding prokaryotic secretion systems. Thus, how rhopty effectors are delivered into the host cytoplasm remains unclear.

To explore the mechanisms of rhopty secretion, we searched for examples of regulated secretion in organisms phylogenetically closely related to Apicomplexa. All Alveolata bear alveolar sacs beneath the plasma membrane, which give the superphylum its name. They contain elaborate membrane-bounded secretory organelles with shared evolutionary origin but different morphologies and functions. Known as trichocysts in the Ciliata *Paramecium*, they function in defense—docking at the plasma membrane and discharging in response to predation. Exocytic membrane fusion occurs at plasma membrane sites consisting of rosettes of 8-9 intramembranous particles (IMPs). Previous electron microscopy (EM) studies have described similar rosettes at the apex of several apicomplexan parasites. Though suggested to be involved in exocytic fusion, the function of the IMPs rosette has never been experimentally addressed in Apicomplexa. Analysis of *Paramecium tetraurelia* mutants defective in both trichocyst exocytosis and rosette assembly led to identification of *nd* (non-discharge) genes (Supplementary Table 1). To determine if similar factors could be involved in exocytosis of secretory organelles in Apicomplexa, we first searched for Nd homologs in the tree of life. Genome
mining for \textit{nd} genes and phylogenetic analyses revealed that \textit{nd6} and \textit{nd9} are conserved in Ciliata, but also in Dinoflagellata, Chromerida and Apicomplexa (Extended data Fig. 1 and Extended Data 1). We also found Nd9 homolog in the Perkinsozoa, a sister group of Dinoflagellata. Altogether, this analysis suggested a conserved evolution and common function of \textit{nd} genes across Alveolata.

To define the parasite localization of Nd proteins, we tagged \textit{nd6} (TGGT1\_248640) and \textit{nd9} (TGGT1\_249730) at the endogenous loci in \textit{Toxoplasma gondii}, an experimentally tractable apicomplexan (Extended data Fig. 2). Both proteins displayed a punctuate signal throughout the cytoplasm, but in addition \textit{TgNgd6} accumulated at the apical tip of the parasite (Fig. 1a, b). A closer look at the apical tip by immuno-EM localized \textit{TgNgd6} at the site of rhoptry exocytosis, in association with the parasite plasma membrane and an underlying membranous spheroid known as the “apical vesicle (AV)” (Fig. 1c). Despite previously observed by EM in \textit{Toxoplasma}\textsuperscript{21,30} and suggested to be linked to rhoptries\textsuperscript{22,30}, the composition and function of the AV remain unknown.

Because \textit{TgNgd6} and \textit{TgNgd9} were predicted to be fitness-conferring genes\textsuperscript{31}, in order to investigate their function we generated inducible knockdown mutants using an auxin-inducible degron for \textit{TgNgd6}\textsuperscript{32} and tetracycline-induced repression for \textit{TgNgd9}\textsuperscript{33,34} (Extended data Fig. 2). Parasites conditionally depleted of \textit{TgNgd6} or \textit{TgNgd9} showed reduced plaque formation on fibroblast monolayers, indicating the inability of both mutants to efficiently complete the lytic cycle (Extended Data Fig. 3a). \textit{Tgnd6-}\textit{iKD} and \textit{Tgnd9-}\textit{iKD} mutants showed no detectible defects in their intracellular replication, egress, conoid protrusion, motility, or host cell attachment (Extended Data Fig. 3b-g). In contrast, invasion was severely impaired for both mutants (Fig. 1d).

To understand the mechanistic basis of this invasion defect, we evaluated microneme and rhoptry secretion in the \textit{Tgnd6-}\textit{iKD} and \textit{Tgnd9-}\textit{iKD} mutants. Release of the micronemal
protein AMA1 into the supernatant was unimpeded by the absence of TgNd6 or TgNd9 (Fig. 1e). In contrast, rhoptry secretion was significantly impaired by the loss of TgNd6 or TgNd9 as revealed by quantifying the release of rhoptry protein ROP1 into host cells using immunofluorescence assays (IFA) (Extended Data Fig. 3h), or the delivery of Cre recombinase fused to the rhoptry protein toxofilin into the nucleus of a suitable reporter cell (Fig. 1f). This was not due to defects in rhoptry biogenesis, as the mutants showed normal organelle formation and apical positioning, as judged by IFA and EM (Extended Data Fig. 4). Analysis by freeze fracture EM of the apex of TgNd6-iKD mutants showed no significant reduction in the presence of the apical rosette (Fig. 1g). In contrast, T. gondii depleted of TgNd9 displayed a strong decrease in cells with a rosette at the parasite apex (Fig. 1g). This last result link rosette formation with rhoptry secretion and support an evolutionarily conserved mechanism of regulated exocytosis in Alveolata.

To determine whether other Apicomplexa require nd genes for rhoptry function, we analysed Nd9 in Plasmodium falciparum, the causative agent of the deadliest form of malaria. We confirmed that P. falciparum intracellular merozoites possess a fusion rosette of 8 IMPs (Fig. 2a). As nd9 is also predicted to have a fitness cost in P. falciparum, we used the rapamycin-inducible dimerizable Cre recombinase (DiCre) system to conditionally excise the low expressed Pfnd9 gene (PF3D7_1232700) (Extended Data Fig. 5a-d). DiCre mediated ablation of Pfnd9 resulted in substantial reduction in parasite proliferation (Fig. 2b), which was due to the inability of Pfnd9-iKO mutants to reinvaode host cells while their intracellular development and egress were unaffected (Fig. 2b, c; Extended Data Fig. 5e). As in T. gondii, microneme secretion was unaltered in Pfnd9-iKO parasites (Fig. 2d) but rhoptry secretion was affected (Fig. 2e).

To gain a more complete understanding of the molecular composition of the rhoptry secretion machinery, we searched for Nd interacting proteins in T. gondii. TgNd9 displays
two Armadillo repeats while TgNd6 shows homology with GDP/GTP exchange factors (GEFs) known to activate GTPases (Fig. 3a). We used TgNd9 for immunoprecipitation (IP) experiments as we found TgNd6 to be largely insoluble (Extended data Fig. 2g). Mass spectrometry analysis revealed robust interaction of TgNd9 with TgNd6 and with TgFER2 (TGGT1_260470), a member of the ferlin calcium sensor family, known to be essential for Toxoplasma rhoptry secretion13 (Fig. 3a and Supplementary Table 2). The TgNd9 IP also enriched TGGT1_222660, a protein harboring Armadillo repeats and Leucine Rich Repeats, named hereafter TgNdP1 (Nd Partner 1) and TGGT1_316730 (TgNdP2: Nd Partner 2), a protein with a C2 calcium lipid binding domain (Fig. 3a). Both genes are broadly shared among Alveolata (Extended Data Fig. 1c, d and Supplementary Data 1) and predicted to be fitness-conferring in Toxoplasma31 and Plasmodium38. In contrast, TgNd9-IP interactor TGGT1_277840, a GTPase, is restricted to Apicomplexa and Dinoflagellata, and TGGT1_253570 is only found in the apicomplexan subgroup Coccidia. Importantly, all these proteins are also recovered when using reverse IP with tagged-TgNdP1 protein (Supplementary Table 3).

To learn more about the new conserved partners of TgNd9, we generated T. gondii lines in which TgNdP1 or TgNdP2 was tagged by an epitope and could be ablated conditionally (Extended Data Fig. 6). Both proteins appear as punctate cytoplasmic staining (Fig. 3b) but TgNdP2—like TgNd6—also appears as a dot at the apical tip of the parasite, although consistently with lower intensity. Depletion of TgNdP1 or TgNdP2 resulted in a profound growth defect (Extended Data Fig. 7a) that we linked to impairment of host cell invasion and rhoptry secretion (Fig. 3c, d and Extended Data Fig. 7). Again, loss of rhoptry secretion went hand-in-hand with loss of the rosette in Tgndp1-iKD and Tgndp2-iKD parasites (Fig. 3e).
To further validate our conservation data and broaden our discoveries back to Ciliata, we generated knockouts for orthologues of both NdP1 (TTERM_01287970, TnΔndp1) and NdP2 (TTERM_00498010; TnΔndp2) in the free-living Ciliata *Tetrahymena thermophila* (Extended Data Fig. 8a, b). The homologous organelles to *Paramecium* trichocysts in *Tetrahymena* are called mucocysts and are non-essential for laboratory growth. We found that TnΔndp1 and TnΔndp2 cells were defective in mucocyst secretion, which was triggered by exposure to dibucaine (Fig. 3f). We further showed that the impairment of exocytosis was not due to defects in mucocyst biogenesis, given that mucocyst maturation (as measured by processing of mucocyst pro-proteins) and trafficking (monitored by IFA) remained unaltered (Extended Data Fig. 8c, d). Taken together, we identified a complex of proteins essential for organellar exocytosis and rosette assembly, conserved across Alveolata.

The exact position of the rosette relative to the apical tip of the rhoptry and the enigmatic AV remained elusive, since freeze-fracture technique—used to image the rosette on the membrane—does not capture the internal structures at the same time. To overcome this limitation, we imaged the *Toxoplasma* apex by cryo-electron tomography (cryo-ET)—a technique combining the advantages of 3D imaging with molecular resolution to reveal ultrastructure *in situ* in its native biological context. We were able to simultaneously visualize and define three linked elements — 1) the rosette (dark blue), 2) the AV (magenta), and 3) the apical tip (cyan) of the rhoptry (orange) (Fig. 4a and b, Extended Fig. 9). The rosette showed an 8-fold rotational symmetry around a central axis (Fig. 4c and d) and extended under the parasite plasma membrane (light blue) to interact with the AV (Fig. 4c). Thus, the rosette is tightly sandwiched between the AV and the plasma membrane and extensively interacts with both membranes. The AV in turn sits over the rhoptry tip (Fig. 4a and b). The contiguity of all elements from the rhoptry tip to the plasma membrane supports the idea that the rosette and the AV are integral part of the rhoptry secretion machinery. Interestingly, the
connection between AV and rosette is sometimes observed even in the absence of a docked rhoptry (Fig. 4e) suggesting that they assemble independently of rhoptry docking. Supporting this hypothesis, when we imaged the *Toxoplasma* ARO mutant in which rhoptries fail to dock and are dispersed in the cytoplasm, both the rosette (seen by freeze-fracture; Fig. 4f) and the AV (seen by EM; Fig. 4g) were still present at the apex of the parasite. Altogether, these results positioned the rosette and AV at the heart of the exocytic machinery in Apicomplexa and implied that rhoptries do not make contact and fuse directly with the plasma membrane.

Our work breaks ground on the molecular and structural mechanisms for rhoptry exocytosis in Apicomplexa. We defined the apical rosette of IMPs as the site for rhoptry exocytosis in Apicomplexa, and we characterized an Alveolata-specific Nd complex necessary for the assembly of the rosette. In *Paramecium*, physiological studies predicted *PtNd6* to be active at the plasma membrane while *PtNd9* appeared to be a diffusible cytoplasmic component interacting with both trichocyst and plasma membranes.

However, both the localization and identity of partners in Ciliata remained unknown. We localized *TgNd9* to the cytoplasm and *TgNd6* to the site of exocytosis in *Toxoplasma*. We found both proteins to form a complex that included *TgNdP1* and *TgNdP2*, which we demonstrated to be essential for organelle exocytosis in both Ciliata and Apicomplexa. This complex further includes proteins with C2 domains (*TgFER2* and *TgNdP2*), a homolog of the membrane fusion Ferlin family (*FER2*), a GTPase (*TGME49_277840*), and a putative GEF protein (*TgNd6*). These proteins and their domains yield a regulatory model in which calcium signaling and nucleotide binding and hydrolysis constitute key steps that control rosette assembly and/or organelle discharge (more discussion in Supplementary text).

In this study, we also shed new light on the enigmatic AV. Described in *Toxoplasma*, it is also visible in early micrographs from other apicomplexan parasites (Extended Data Fig. 10). We show that the AV is on one side tightly
connected with the plasma membrane via the rosette, and from the other sits on the tip of the rhoptry, precluding a direct link and fusion between the rhoptry and the plasma membrane. In Ciliata, where no comparable AV is present, the docking and fusion of the trichocysts to the plasma membrane shapes the rosette. Diverging from this scenario, we also showed that the docking of the rhoptries in *Toxoplasma* is dispensable for rosette formation (Fig. 4e-g), which suggests that the assembly of the rosette in *Toxoplasma* might be instead induced by docking of the AV.

While *Paramecium* trichocysts are discharged into the environment to thwart predators, apicomplexan rhoptries translocate their contents directly into the cytoplasm of host organisms to infect and parasitize. The difference in terms of presence or absence of the AV (Fig. 4h) suggests that the AV is an adaptation in Apicomplexa for parasitism and cell invasion. Its presence may reflect additional complexity of the secretory machinery in apicomplexans, in which exocytosis must be coupled with injection of rhoptry content through the barrier of a host cell membrane. In support of this hypothesis, a similar vesicle is present at the apex of *Perkinsus marinus* (previously named *Dermocystidium marinum*). This organism possesses rhoptries and invades and parasitizes the cells of oysters. Phylogenetically, *Perkinsus* falls between Dinoflagellata and Apicomplexa and is viewed as a basal taxon and example of early adaptation to parasitism.

The rosette and the set of Nd proteins highlight a common ancestry for the fusion machinery linked to secretory organelles in two groups of protists that diverged hundreds of millions of years ago and have adopted radically different lifestyles. The architecture and molecular composition of the rhoptry system that enables the intracellular parasitism of Apicomplexa is now revealed. Future studies combining high resolution structural biology approaches with the genetics and cell biology of *Toxoplasma* will allow a full understanding
of how this fascinating mechanism unfolds and is regulated. Such an understanding may find application in designing strategies against malaria, cryptosporidiosis and toxoplasmosis.

Methods

Parasite Immunofluorescence microscopy

Immunofluorescence assays (IFAs) on intracellular T. gondii parasites were conducted as previously described\(^46\). Briefly, cell monolayers were washed and fixed in 4% paraformaldehyde (PFA) in Phosphate Saline Buffer (PBS) for 20 min. After three washes with PBS, cells were permeabilized with 0.1% (v/v) Triton X-100 or with saponin 0.1% (v/v) (for invading parasites) in PBS for 5 min, blocked with 10% fetal bovine serum in PBS (PFBS) for 45 min, incubated with primary Abs diluted in 2% PFBS, washed three times, and then incubated with secondary antibody. The coverslips were mounted onto microscope slides using Immunomount (Calbiochem).

For IFAs on P. falciparum, thin blood smears of highly synchronized DMSO and rapamycin-treated PfNd9-iKO schizonts were air-dried, fixed in 4% (w/v) PFA and permeabilized in 0.1% (v/v) Triton X-100. After blocking in 1.5% BSA, samples were probed for 1 h with the suitable antibodies: rabbit anti-PfAMA1\(^47\) antibody (1:1000); mouse anti-PfMSP1\(^48\) antibody (1:1000); mouse anti-PfRAP2\(^49\) antibodies (1:500). The secondary antibodies used were Alexa Fluor 488 and 594-conjugated antibodies against mouse or rabbit IgG (highly cross-adsorbed) both diluted 1:4000 (Molecular Probes). Samples were then counterstained with Hoechst and slides mounted with Vectashield® antifade mounting medium.

Except when specified, observations were performed with a Zeiss Axioimager Z1 epifluorescence microscope equipped with a Zeiss Axiocam MRm CCD camera and 100X/1.4 Oil Plan Apochromat objective. Images were processed using Zen Blue 2.3 pro
(Zeiss) software. Imaging of TgNd6, TgNd9, TgNdP1 and TgNdP2 co-localization with rhopty protein TgARO (Fig. 1 and Fig. 3) was performed on a ZEISS confocal LSM880, equipped with an Airyscan detector and a 63X/1.4 Oil Plan Apochromat objective. Zen Black (Zeiss) was used for image airyscan processing. Z-stack images and z-projection images were denoised, adjusted in brightness and contrast, and colored with the program Fiji.

Adjustments for brightness and contrast were applied uniformly on the entire images and when required, matching pairs of images were recorded with the same exposure time and processed identically. All optical images were collected at the Montpellier Ressources Imagerie facility of the University of Montpellier (MRI, www.mri.cnrs.fr).

Invasion assays

Toxoplasma plaque assays, replication assays, and two-color invasion assays were performed as previously described. Briefly, to synchronize invasion, freshly egressed tachyzoites (5 × 10^6), pre-treated for 48 h ± ATc (Tgnd9-iKD and Tgndp1-iKD) or 24 h ± IAA (Tgnd6-iKD and Tgndp2-iKD), were harvested and settled on ice for 20 min on HFF monolayer grown on coverslips in 24-well plates. Invasion was allowed for 5 min at 38 °C and stopped by fixation with 4% PFA in Hank's Balanced Salt Solution (HBSS) for 20 min at room temperature. To detect extracellular parasites, immuno-detection was performed with the mouse mAb T4 1E5 anti-SAG1 antibody (dilution 1:1000) in 2% FCS/HBSS, without previous permeabilization. After permeabilization with 0.01% saponin for 15 min, a second IFA was performed using rabbit anti-ROP1 antibody (dilution 1:1000) to label the parasitophorous vacuole of intracellular parasites. Extracellular and intracellular parasites were counted by microscopic examination of at least 20 fields per coverslip (n=3 coverslips). Graphs show the mean of three independent invasion assays.
Microneme secretion assay

Microneme secretion in *T. gondii* was assayed by monitoring the release into the culture medium. Freshly egressed *T. gondii* tachyzoites, pre-treated for 48 h ± ATc (*Tgnd9*-iKD and *Tgndp1*-iKD) or 24 h ± IAA (*Tgnd6*-iKD and *Tgndp2*-iKD), were harvested by centrifugation at 600 g, and washed twice in intracellular buffer (5 mM NaCl, 142 mM KCl, 1 mM MgCl₂, 2 mM EGTA, 5.6 mM glucose and 25 mM HEPES, pH 7.2), prewarmed to 37°C. Parasites were resuspended in DMEM (supplemented with 2 mM glutamine) ± propranolol 500 µM, and incubated at 37°C for 20 min to induce microneme secretion. Parasites were pelleted at 1000 g for 5 min at 4°C, washed once in PBS and stored at -20°C. Supernatants were centrifuged at 2000 g for 5 min, at 4°C and used as ESA (excreted/secreted antigen). Pellets and ESA samples were analysed for micronemal protein (AMA1) by Western blot.

Microneme secretion in *P. falciparum* was assayed by monitoring the surface translocation of AMA1. Highly synchronized DMSO and rapamycin-treated *Pfnd9*-iKO schizonts were exposed to 10 µM E64 to block egress. IFAs were then performed with anti-*PfAMA1* antibodies to analyse AMA1 translocation to the merozoite surface.

Rhoptry secretion assays

To evaluate the efficiency of rhoptry secretion in *T. gondii*, we expressed in the *Tgnd*-iKD and *Tgndp*-iKD lines a rhoptry secretion reporter protein consisting of the rhoptry protein toxofilin fused with Cre-recombinase, a nuclear localization signal (NLS), and a myc tag, collectively called Secreted Cre, Epitope-tagged (SeCrEt)

36. The reporter strains generated were called *Tgnd9*-iKD_SeCrEtUPRT, *Tgndp1*-iKD_SeCrEtUPRT, and *Tgnd6*-iKD_SeCrEtHXGPRT. Ds Red cells, constitutively expressing DsRed and able to switch to eGFP expression upon Cre-mediated recombination, were used as reporter cells

36. DsRed
Cells were maintained at < 50% confluence (regardless of the flask size) and plated the day before parasite infection, at a density of 2 x 10^5 cells /ml (in T25 flasks). Tachyzoites pre-treated for 48 h ± ATc (Tgnd9-iKD and Tgndp1-iKD) or 24 h ± IAA (Tgnd6-iKD) were collected and used to infect the DsRed cells at a multiplicity of infection (MOI) of 3. 24 hours post-invasion (hpi) infected DsRed cells were trypsinized and examined by fluorescence-activated cell sorting (FACS) to assess the percentage of Ds-Red (rhoptry secretion impairment) and GFP (successful rhoptry secretion) expressing cells. Number of GFP positive cells were normalized as the percentage compared to 100% in the control (-IAA or -ATc).

Additionally, rhoptry secretion has been also quantified by classical e-vacuole assay. Freshly egressed parasites, pre-treated for 48 h ± ATc (Tgnd9-iKD and Tgndp1-iKD) or 24 h ± IAA (Tgnd6-iKD), were preincubated with 1 µM of cytochalasin D (cytD) for 10 min and then incubated with HFF cells in the presence of cytD for 15 min. IFAs were then performed with anti-ROP1 (rhoptry secretion) and anti-SAG1 (parasite surface), and the number of ROP1 stainings per field was determined by microscopic examination of at least 20 fields per coverslip (n = 3 coverslips).

To quantify rhoptry secretion in P. falciparum, the same amount of DMSO and rapamycin-treated Pfnd9-iKO purified schizonts were arrested with 1.5 µM C2 for 4 h before being washed twice in order to allow them to egress. Parasites were then incubated for 30 min in complete medium with 1 µM cytD in the presence of red blood cells (RBCs). IFAs were then performed with anti-PfRAP2 antibodies to visualise rhoptry secretion events ('spits’ of RAP2 export into the RBC). The number of secretion events were counted over the total of RBCs by microscopic examination of 11 fields (~3000 events analysed).

**Freeze-fracture**
Cells used for the freeze-fracture were harvested, pelleted and fixed in 2.5% glutaraldehyde in 0.1 M phosphate buffer at room temperature for 2 h. Following primary fixation, the samples were rinsed in the same buffer, and kept overnight in 30% glycerol 0.1 M phosphate solution. The cells were then quickly frozen by immersion in liquid nitrogen under vacuum. The frozen samples were fractured in a BAL-TEC BAF 060 apparatus; subsequently the fracture face was shadowed by evaporating platinum 45° (3.2nm) and Carbon 90° (25nm). The replicas were then washed in 6.5% sodium hypochlorite, rinsed in a chloroform (2/3) - (1/3) solution, rinsed in distilled water and mounted on copper grids. In order to interpret the replicas obtained in freeze-fracture technique one must keep in mind that the portion of the lipid bilayer associated with the exterior of the cell is termed the E-face, or extracellular face. The portion associated with the interior of the cell is termed the P-face, or protoplasmic face.

In vitro growth assay of P. falciparum parasite asexual development

The growth capability of Pfnd9-iKO mutant parasites was assessed by comparing it with the p230p DiCre parental line ± rapamycin in biological triplicates. DMSO (vehicle control) and rapamycin-treated cultures were synchronized at ring stages by sorbitol synchronization and parasitaemia adjusted to 0.2% to follow growth over 3 cycles. Parasites around 30 hpi were collected at each cycle to determine parasitaemia by counting Giemsa-stained parasites on at least 1000 RBCs per culture.

P. falciparum induced egress and time-lapse microscopy

P. falciparum egress was imaged using C2 compound to tightly synchronize egress. DMSO and rapamycin-treated Pfnd9-iKO parasites were highly synchronized and blocked 40 hpi (mature schizonts) with 1.5 µM C2 compound to prevent egress. Four hours later parasites
were washed twice with warm RPMI 1640 medium (Gibco), to remove C2, and placed into a 35 mm Dish (MatTek). The parasites suspension was sealed by adhering a 22 x 22 mm square coverslip to the Dish and the preparation was introduced on a temperature-controlled microscope stage at 37°C. Bright field images were collected 5 min after washing off the C2 at 1 s intervals over a total of 15 min using a ZEISS AxioObserver Microscope fitted with a coolsnap HQ2 digital camera and 63X/1.4 Oil Plan Apochromat objective. Images were exported to MOV movies using ZEN 2 (blue edition) software. Number of egress events in 15 min were normalized as the percentage compared to 100% in the control.

**Tetrahymena dibucaine assay to quantify mucocyst secretion**

*Tetrahymena* cells were grown to stationary phase (10^6 cells/ml) in 25 ml SPP (2% proteose peptone, 0.1% yeast extract, 0.2% dextrose and 0.003% ferric-EDTA supplemented with 250 μg/ml penicillin G, 250 μg/ml streptomycin sulfate, and 0.25 μg/ml amphotericin B fungizone) for 48 h, and then concentrated by centrifugation into a loose 1.5ml pellet. Cells were stimulated with 2.5 mM dibucaine, vigorously mixed for 30 s and diluted to 15 ml with 10 mM Hepes, pH 7.5, and 5 mM CaCl2. After gently mixing, the culture was centrifuged at 1,200 g for 2 min, resulting in the formation of a cell pellet/flocculent bilayer. Quantification of mucocysts secretion has been done by weighing the flocculent layer overlying the pellet of cells.

**Transmission electron microscopy**

*Tgnd9*-iKD and *TgARO*-iKD, together with the *Δku80*-TATi parental strain, were treated with ATc for 72 h. *Tgnd6*-iKD parasites, and *Δku80*-Tir1 parental strain were treated with IAA for 24 h. Extracellular parasites were collected in the culture medium after natural egress and fixed by adding and equal volume of phosphate buffer 0.1 M containing 5% of
glutaraldehyde for one hour at RT. Parasites were then centrifuged and resuspended in 1 ml
of fresh buffer with 2.5% glutaraldehyde for 2 h before being kept at 4°C until further
processing. Each of the following steps were performed in suspension followed by
centrifugation steps in a tabletop microcentrifuge. Sample were post-fixed with 1% OsO₄ and
1.5% potassium ferricyanide in 0.1 M phosphate buffer for 1 hour at RT, washed in water and
then incubated in 2% uranyl acetate in water overnight at 4°C. Tachyzoites were then
dehydrated in growing concentration of acetonitrile, followed by impregnation in Epon118:
acetonitrile 50:50 for 2 hours, 90:10 for 2 additional hours and then overnight in pure epon.
Pellets were polymerized in fresh Epon for 48 h at 60°C. 70 nm ultrathin sections were cut
with a Leica ultracut (Leica microsystems), counterstained with uranyl acetate and lead
citrate.

Immunoelectron microscopy

*T. gondii*-HA₃ parasites, infected fibroblast monolayers were trypsinized and fixed with equal
volume of 8% formaldehyde (FA) in phosphate buffer overnight at 4°C and resuspended in
4% fresh FA until further processing. Cells were then incubated with 0.1 % glycine in
phosphate buffer, pelleted and embedded in 12% gelatin, cut in small blocks (< 1 mm) and
infused in 2.3 M sucrose on a rotating wheel for 24 h at 4°C. Gelatin blocks were mounted on
specimen pins and frozen in liquid nitrogen. Cryo-sectioning was performed on a Leica UC7
cryo-ultramicrotome, 70 nm cryosections were picked-up in a 1:1 mixture of 2.3 M sucrose
and 2% methylcellulose in water and stored at 4°C. For on-grid immunodetection, grids were
floated on 2% gelatin in PBS for 30 min at 37°C to remove methylcellulose/sucrose mixture,
then blocked with 1% skin-fish gelatin (SFG, Sigma) in PBS for 5 min. Successive
immunolabeling steps were performed on drops as follows: 1) rat monoclonal anti-HA
antibodies (clone 3F10, Roche) in 1% BSA, 2) rabbit polyclonal anti-rat IgG antibody
(Sigma) in 1% BSA, 3) Protein A-gold (UMC Utrecht) in 1% BSA. Four washes (2 min each) with 0.1% BSA were performed between steps. After Protein A treatment, grids were washed four times for 2 min each with PBS, fixed 5 min in 1% glutaraldehyde in water, and washed six times for 2 min each with distilled water. Grids were then incubated with 2% methylcellulose: 4% uranyl acetate 9:1 on ice in the dark for 15 min, picked-up on a wire loop and air-dried. All chemicals were from Electron Microscopy Sciences (USA), solvents were from Sigma. Observations and image acquisition were performed on a Jeol 1200 EXII transmission electron microscope at the Electron Microscopy Platform of the University of Montpellier (MEA; http://mea.edu.umontpellier.fr). Transmission electron microscopy images were processed with Fiji for contrast optimization and the Image J plugin was used to make the EM panels.

**Cryo-electron-tomography (Cryo-ET)**

Freshly isolated *Toxoplasma gondii* cells were suspended in HBSS along with fiducials (10 nm colloidal gold from Ted Pella for alignment of tilt series). A 4 µl drop of the suspension was applied onto EM grids, excess liquid blotted away, and plunge frozen in a liquid ethane/propane mixture (pre-cooled using liquid nitrogen) using a EM GP2 automatic plunger (Leica Microsystems, Wetzlar, Germany). The blotting chamber was set to 95-100% relative humidity at 37°C and blotting was done either from the sample side of the grid or from the back using Whatman filter paper #1. Plunge-frozen grids were subsequently loaded into autogrid cartridges (ThermoFisher). EM cartridges containing frozen grids were stored in liquid nitrogen and maintained at ≤−170°C throughout storage, transfer and cryo-ET imaging.
Cryo-ET data collection was performed on a ThermoFisher Krios G3i 300 keV field emission gun cryo-TEM equipped with a 6k x 4k K3 direct electron detector (Gatan, Inc.) at the Beckman Center for Cryo-Electron Microscopy in the Singh Center for Nanotechnology, University of Pennsylvania. The camera was operated in electron counting mode to enable motion correction. An energy filter (Gatan Imaging Filter, Gatan, Inc.) with a slit width of 20 eV was used to increase the contrast of the projection images. Additionally, a Volta phase plate was used to boost the image contrast at defoci of negative 2-3 µm. A magnification of 33,000X with corresponding pixel sizes of 2.65 Å was used for imaging. SerialEM software was used for all imaging. Cells were first assessed at lower magnifications for suitability of ice thickness and outer membrane integrity (the parasites sometimes tend to vesiculate their outer membrane possibly in response to blotting). Once the target cells were identified and marked, anchor maps were used to revisit these locations and collect tilt-series in an automated fashion. Each tilt-series was collected from negative 60° to positive 60° with an increment of 2° in an automated fashion using the low dose functions of tracking and focusing. The cumulative dose of each tilt-series ranged between 100 and 150 e/Å². Once acquired, tilt-series were binned twice or four times before aligned using the 10 nm colloidal gold as fiducials and reconstructed into tomograms by our in-house automated computation pipeline utilizing the IMOD software package. For presentation purposes, tomograms were favorably oriented in three dimensions before averaging a few slices around the slice of interest (to enhance contrast) using the slicer window in IMOD. In a few cases (Figs. 4b-e, contrast was further enhanced using the non-linear anisotropic diffusion filter in IMOD.

**Immunopurification and mass spectrometry analysis**

Immunopurification was performed using anti HA magnetic beads (Pierce R88836) as previously described. Purified proteins were loaded on a SDS-PAGE and digested in gel (2
bands per sample) as previously described\textsuperscript{58}. Samples were loaded onto a 25 cm reversed
phase column (75 mm inner diameter, Acclaim Pepmap 100\textsuperscript{®} C18, Thermo Fisher Scientific)
and separated with an Ultimate 3000 RSLC system (Thermo Fisher Scientific) coupled to a Q
Exact HFX (Thermo Fisher Scientific). MS/MS analyses were performed in a data-
dependent mode. Full scans (375 – 1,500 m/z) were acquired in the Orbitrap mass analyzer
with a resolution of 60,000 at 200 m/z. For the full scans, 3e6 ions were accumulated within a
maximum injection time of 60 ms. The twelve most intense ions with charge states ≥ 2 were
sequentially isolated (1e5) with a maximum injection time of 45 ms and fragmented by HCD
(Higher-energy collisional dissociation) in the collision cell (normalized collision energy of
28\%) and detected in the Orbitrap analyzer at a resolution of 30,000.
Raw spectra were processed using the MaxQuant\textsuperscript{59} using standard parameters with label-free
quantification (LFQ) and match between runs\textsuperscript{60}. MS/MS spectra were matched against the
UniProt Reference proteomes of \textit{T. gondii} and Human (respectively Proteome ID
UP000001529 and UP000005640) and 250 frequently observed contaminants as well as
reversed sequences of all entries (MaxQuant contaminant database). Statistical analysis were
done using Perseus on LFQ data\textsuperscript{61}.

\textbf{Amino acid sequence alignments and phylogenetic analyses, cell and parasite culture,}
parasite cloning strategies, parasite transfections, parasite immunoblots, \textit{T. gondii}
plaque assays, \textit{T. gondii} intracellular growth assay, \textit{T. gondii} immunofluorescence-based
induced egress assay, \textit{T. gondii} conoid extrusion assay, \textit{T. gondii} gliding assays, \textit{T. gondii}
attachment assay, Ciliata culture conditions, Ciliata biolistic transformation, generation
of \textit{Tetrahymena} knockout strain, Ciliata immunoblots, Ciliata immunofluorescence
microscopy, statistical analysis, reagents and antibodies are found in Supplementary
Methods.
Data availability

The datasets generated during and/or analysed during the current study are available from the corresponding author upon request.

References


38. Zhang, M. et al. Uncovering the essential genes of the human malaria parasite

39. Briguglio, J. S., Kumar, S. & Turkewitz, A. P. Lysosomal sorting receptors are essential

40. Mueller, C. et al. The *Toxoplasma* protein ARO mediates the apical positioning of
   rhoptry organelles, a prerequisite for host cell invasion. *Cell Host Microbe* **13**, 289–301
   (2013).

41. Beisson, J., Cohen, J., Lefort-Tran, M., Poupille, M. & Rossignol, M. Control of
   membrane fusion in exocytosis. Physiological studies on a *Paramecium* mutant blocked

   genes involved in exocytotic membrane fusion in *Paramecium*. *Genetics* **130**, 461–70

43. Varghese, T. Fine structure of the endogenous stages of *Eimeria labbeana*. I. The first

44. Sheffield, H. G. Electron microscope study of the proliferative form of *Besnoitia

   (1976).

46. El Hajj, H. et al. Molecular signals in the trafficking of *Toxoplasma gondii* protein MIC3 to

   antibody blocks interactions between components of the malarial invasion machinery.


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**Author contributions**

E.A. performed most of the work on *Toxoplasma*. M.M.C generated the *Plasmodium* data and completed the work on *Toxoplasma* during revision. A.G. performed the phylogenetic analyses. E.A. D.S and R. N. generated the *T. thermophila* data. C.S. generated the donor plasmid for *Pfnd9-iKO*. A.N.G., N.D.S.P., R.N. and M.M. contributed to *Toxoplasma* phenotypic analyses. D.M.P.V. performed the IP analyses. S. U. performed the mass spectrometry analyses. E.A., L.B-S and J.F.D. performed electron-microscopy. E. A., P.R.F and J.F.D. performed the freeze fracture analyses. S.K.M. and Y.-W.C. performed the cryo-ET. A.N.G. and S.K.M. prepared the samples for cryo-ET. M.L. designed the study with the support of E.A. M.L. supervised the research. E.A. and M.L. wrote the paper with editorial support from M.M.C., A.P.T., D.S., A.N.G., S.K.M., Y.-W.C. and B. S. The data are included in the main manuscript and in the supplementary materials.
Competing interests
The authors declare no competing financial interests.

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Supplementary Information is available for this paper.

Fig. 1 | Rhoptry secretion is dependent on rosette formation.

a, Immunofluorescence assay (IFA) of control untagged line and endogenously HA₃-tagged TgNd6 and HA₃-tagged TgNd9 tachyzoites. The green arrow points to TgNd6 apical puncta. ARO is a marker for rhoptries. Shown are maximum intensity projections of confocal z-stacks of fixed parasites. b, Super-resolution microscopy of the apical dot of TgNd6. Schematic of apical end of tachyzoite. Maximum intensity projections of z-stacks of TgNd6-HA₃ parasites transiently expressing RNG1_GFP (top) and of TgNd6-HA₃ parasites expressing centrin 2 (CEN2)-Ty₂ (bottom). The protein RNG1_GFP marks the apical polar ring (APR) and CEN2_Ty marks pre-conoidal rings (PCR). Higher magnifications show that TgNd6-HA₃ (red arrow) localizes above the apical polar ring (green arrow) and co-localizes partially with CEN2. DIC: differential interference contrast. c, Immunogold labelling of TgNd6-HA₃. Right panel shows TgNd6-HA₃ on the AV. Insert panel: higher magnification of the AV. Micronemes (m) and rhoptries (Rh) are visible in transverse section of the conoid (Co). d, Quantification of invasion after depletion of TgNd6 and TgNd9. Mean ± SD of n=3 independent experiments. e, Immunoblot showing microneme secretion assessed by the release of proteolytically cleaved AMA1 (arrow=processed/secreted TgAMA1) in Tgnd6-
iKD ± IAA 24 h (left) and Tgnd9-iKD ± ATC 72 h (right). P = pellet, Sup = supernatant, Sup ind = propanolol-induced supernatant. GRA3, loading control. f, Rhooptry secretion assay by Secreted Cre, epitope-tagged (SeCrEt). Rhooptry secretion quantification of Tgnd6-iKD (left) ± IAA and Tgnd9-iKD (right) ± ATC. Mean ± SD of n=3 independent experiments. g, Left: Freeze-fracture electron microscopy of a T. gondii tachyzoite (P face) showing a rosette of intramembranous particles (white arrow) at the middle of the apex. Middle: Higher magnification of the left panel. The white arrowheads point to the eight IMPs of the rosette. Right: Quantification of rosettes of IMPs in Tgnd9-iKD ± ATc 72 h and Tgnd6-iKD ± IAA 24 h using freeze fracture. (d, f) Unpaired two tail student’s t test: **** p-value < 0.0001, *** p-value < 0.001, ** p-value < 0.01, * p-value < 0.05.
**Fig. 2** | **PfNd9** is essential for rhoptry secretion in *P. falciparum*.

**a**, Freeze-fracture electron microscopy of a *P. falciparum* merozoite (P face) showing a rosette of intramembranous particles (white arrow). Higher magnification at the bottom. Bar is 100 nm. **b**, Growth curves (parasitaemias) of *p230p DiCre* (Ctrl) and *Pfnd9-iKO* mutant ± rapamycin shows that *PfNd9*-depleted parasites have a growth defect. On the right: Giemsa staining of the growth experiment illustrating development and reinvasion of *p230p DiCre* (Ctrl) and *Pfnd9-iKO* asexual parasites (along 2 cycles) ± rapamycin treatment. **c**, Quantification of egress of *Pfnd9-iKO* ± rapamycin schizonts. Data collected from 8 movies of *Pfnd9-iKO* ± rapamycin. **d**, Left: IFA illustrating AMA1 protein stored in micronemes (top) or secreted and translocated at the surface of the parasite prior to egress (bottom). **e**, Quantification of rhoptry secretion events in *Pfnd9-iKO* ± rapamycin-treated schizonts using anti-*PfRAP2* antibodies to visualise rhoptry secretion events (‘spits’ of RAP2 exported into the RBC). (e) Unpaired two tail student’s *t* test: **** *p*-value < 0.0001.
**Fig. 3** Nd6 and Nd9 are part of an Alveolata complex essential for organelle secretion in *T. gondii* (Apicomplexa) and *T. thermophila* (Ciliata)

a. Mass spectrometry analysis of immuno-isolated TgNd9-HA3. Left: Volcano Plot of proteins differentially enriched in TgNd9 vs control IP. This plot presents the fold change (Difference) and significance (-Log p) obtained from a t-test of three independent IPs using LFQ intensity values. Right: Schematic representation of TgNd proteins using SUPERFAMILY62. RCC1: regulator of chromosome condensation 1-like domains (RLDs), a versatile domain that performs many different functions, including guanine nucleotide exchange on small GTP-binding proteins63. LRR: Leucine Rich Repeat domain. ARM: Armadillo Repeat domain. C2: lipid-calcium binding domain. 

b. Immunofluorescence (IF) of endogenously HA3-tagged TgNdP1 and TgNdP2 tachyzoites. The white arrow points to TgNdP1 apical dots of two adjacent parasites, which are magnified on the right. ARO: rhoptry marker. DAPI: DNA marker. DIC: differential interference contrast. 

c. Quantification of invasion after depletion of TgNdP1 (left; using ATc) and TgNdP2 (right; using IAA) along with negative control strains. Mean ± SD of n=3 independent experiments. 

d. Rhoptry secretion quantification of TgndP1-iKD (left) ± ATc and TgndP2-iKD (right) ± IAA. Mean ± SD of n=3 independent experiments. 

e. Quantification of rosettes of IMPs in TgndP1-iKD ± ATc 72 h and in TgndP2-iKD ± IAA 24 h using freeze-fracture. 

f. Quantification of mucocyst exocytosis by dibucaine assay. Data collected from three experiments. (c-f)

Unpaired two tail student’s t test: **** *p*-value < 0.0001, *** *p*-value < 0.001, ** *p*-value < 0.01, * *p*-value < 0.05.
**Fig. 4** | **The rhoptry secretion machinery includes the apical vesicle and the rosette**

- **a**, A slice through a tomogram showing a side view of the apical complex – conoid (brown), pre-conoidal apical rings (PCR; gray; two in number), micronemes (yellow), plasma membrane (PM; light blue) and the rhoptry secretion system consisting of the rosette (dark blue), apical vesicle (AV; magenta), rhoptry (orange) and rhoptry tip density (cyan). Original image (right) is annotated with color overlays (left).
- **b**, Left: magnified image of the boxed region in (a) showing the connections between the rhoptry, rhoptry tip density, AV, rosette and the PM. The rhoptry tip is 9 nm distant from the AV. Right: a pair of 3-dimensional segmentations from the data on the left. The PM is rendered transparent in one of these segmentations to see the rosette.
- **c**, Magnified image of the boxed region in (b) showing the side view of the rosette. The AV is 14 nm distant from the PM.
- **d**) Top view of the rosette from a horizontal tomogram section perpendicular to the plane in (c), showing an 8-fold rotational symmetry and a diameter of ~67 nm.
- **e** AV connected with the PM via a rosette in the absence of docked rhoptry. All measurements are made in 3D. Images in (b-e) are computationally filtered to boost contrast (see Methods). The images in (c) and (d) are from two different cells oriented differently on the EM grid resulting in better resolved side view and top view, respectively.
- **f**, Quantification of apical rosettes in *TgARO-iKD* mutants ± ATc 72 h.
- **g**, Left: Ultrastructure of wild type (RH strain type I) tachyzoites with the AV positioned beneath the plasma membrane, and above the tip of the rhoptry neck. Right: In *TgARO-iKD* ATc-treated tachyzoites (72 h), the AV is still properly positioned at the apex (presumably under the rosette), while rhoptries are not docked on the AV. Inset shows a magnification of the vesicles. co: conoid; m: micronemes; Rh: rhoptry; APR: apical polar ring; PCR: pre-conoidal rings; AV: apical vesicle; ICM: pair of intraconoidal microtubules.
- **h**, Schematic of similarities and differences of the exocytic machinery between Ciliata and
Apicomplexa. Exocytosis in Alveolata (Ciliata, Dinoflagellata and Apicomplexa) is outlined by the presence of a rosette of particles embedded in the outer membrane, defining the site of exocytosis. In Ciliata, organelles discharge can have a defensive or predatory function.

Rhoptry exocytosis in Apicomplexa is one of the critical steps of host cell invasion and therefore fundamental for parasitism. In Apicomplexa, but not in Ciliata, an apical vesicle (AV) of unknown function is present between the tip of the rhoptry and the plasmalemma, plausibly involved in the injection of rhoptry proteins into the host cell. PM, plasma membrane; PVM, parasitophorous vacuole membrane; AS, alveolar sac [in Ciliata], which is homologous to the IMC (inner membrane complex) of Apicomplexa.