A Member of the Ferlin Calcium Sensor Family Is Essential for Toxoplasma gondii Rhoptry Secretion

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ABSTRACT Invasion of host cells by apicomplexan parasites such as Toxoplasma gondii is critical for their infectivity and pathogenesis. In Toxoplasma, secretion of essential egress, motility, and invasion-related proteins from microneme organelles is regulated by oscillations of intracellular Ca2+. Later stages of invasion are considered Ca2+ independent, including the secretion of proteins required for host cell entry and remodeling from the parasite’s rhoptries. We identified a family of three Toxoplasma proteins with homology to the ferlin family of double C2 domain-containing Ca2+ sensors. In humans and model organisms, such Ca2+ sensors orchestrate Ca2+-dependent exocytic membrane fusion with the plasma membrane. Here we focus on one ferlin that is conserved across the Apicomplexa, T. gondii FER2 (TgFER2). Unexpectedly, conditionally TgFER2-depleted parasites secreted their micronemes normally and were completely motile. However, these parasites were unable to invade host cells and were therefore not viable. Knockdown of TgFER2 prevented rhoptry secretion, and these parasites failed to form the moving junction at the parasite-host interface necessary for host cell invasion. Collectively, these data demonstrate the requirement of TgFER2 for rhoptry secretion in Toxoplasma tachyzoites and suggest a possible Ca2+ dependence of rhoptry secretion. These findings provide the first mechanistic insights into this critical yet poorly understood aspect of apicomplexan host cell invasion.

IMPORTANCE Apicomplexan protozoan parasites, such as those causing malaria and toxoplasmosis, must invade the cells of their hosts in order to establish a pathogenic infection. Timely release of proteins from a series of apical organelles is required for invasion. Neither the vesicular fusion events that underlie secretion nor the observed reliance of the various processes on changes in intracellular calcium concentrations is completely understood. We identified a group of three proteins with strong homology to the calcium-sensing ferlin family, which are known to be involved in protein secretion in other organisms. Surprisingly, decreasing the amounts of one of these proteins (TgFER2) did not have any effect on the typically calcium-dependent steps in invasion. Instead, TgFER2 was essential for the release of proteins from organelles called rhoptries. These data provide a tantalizing first look at the mechanisms controlling the very poorly understood process of rhoptry secretion, which is essential for the parasite’s infection cycle.

KEYWORDS Toxoplasma gondii, calcium, ferlin, micronemes, protein secretion, rhoptries
The apicomplexan parasite *Toxoplasma gondii* infects one in every three humans. Clinical symptoms of toxoplasmosis derive from the tissue destruction and inflammation caused by repeated rounds of host cell invasion, intracellular replication, and lytic egress of the tachyzoite life stage. Egress is mediated by intracellular Ca\(^{2+}\) ([Ca\(^{2+}\)]\(i\)) fluctuations that trigger release of proteins from the microneme organelles (1). Following egress, parasites move via gliding motility to a new host cell, triggered by additional parasite [Ca\(^{2+}\)], oscillations that facilitate further release of micronemes (2). Subsequent host cell invasion also relies on micronemal proteins (3). The micronemes are localized at the apical end of the parasite and are released from the apical tip (4). Following initial recognition of a host cell, the parasite engages in a tighter interaction with the target cell, mediated by proteins secreted from the rhoptries. The club-shaped rhoptries are anchored at the parasite’s apical end, from where they secrete their contents (5, 6). Proteins in the apical rhoptry neck (RONs) are secreted into the host cell before the rhoptry bulb proteins (ROPs) are released (7). RONs function in tightening the parasite-host attachment by forming a moving junction (MJ), whereas ROPs modulate a variety of host cell pathways to accommodate intracellular replication (6). Rhoptry secretion must be preceded by microneme secretion and requires recognition of a host cell. Although the molecular details of the underlying signal transduction pathways and the mechanism of rhoptry exocytosis remain obscure (8), rhoptry release is generally assumed to be Ca\(^{2+}\)-independent. The last step in establishing infection of a new cell is secretion of host cell-remodeling proteins from the parasite’s dense granules, which again is believed to be Ca\(^{2+}\)-independent.

The Ca\(^{2+}\) signal during egress and invasion is transduced by several molecular mechanisms, including calmodulin (2), calcineurin (9), Ca\(^{2+}\)-dependent protein kinases (CDPK1 [10] and CDPK3 [11–13]), and at the site of microneme exocytotic membrane fusion by the *T. gondii* DOC2.1 protein (14), referred to as TgDOC2 here. Unlike other well-studied Ca\(^{2+}\)-triggered exocytosis models, the only identifiable domain in TgDOC2 is the namesake double C2 domain ("DOC2"), making its organization unconventional. In model organisms, Ca\(^{2+}\)-mediated vesicle fusion with the plasma membrane is organized by at least three DOC2 domain proteins, of which at least one contains a transmembrane domain (15–18). C2 domains bind to other proteins or phospholipids. Ca\(^{2+}\) can make these C2 domain interactions conditional through its association with positionally conserved Asp residues in a select number of C2 domains (15, 19). The ferlins are unique among the DOC2 domain proteins as they contain five to seven C2 domains rather than two and are relatively large (~200 kDa). The ferlins comprise an ancient eukaryotic protein family present in most unicellular organisms (except amoebas and fungi), including the *Apicomplexa* and all multicellular organisms (except higher plants) (20). Although ferlins are relatively understudied due to their absence from neurons and yeast, they typically function in membrane fusion, vesicle trafficking, and membrane repair. Dysfunction of human ferlins can cause deafness and muscular dystrophy (21).

To better understand the machinery underlying *Toxoplasma* Ca\(^{2+}\)-mediated exocytosis, we evaluated the DOC2 domain family in the *Apicomplexa*. Next to the unconventional TgDOC2 (14), we identified three DOC2 proteins of the ferlin family, two of which are widely conserved across *Apicomplexa*. We determined that TgFER2, a conserved representative, is essential for host cell invasion and required for rhoptry secretion. These findings provide critical insight into the poorly understood mechanisms of rhoptry secretion while raising the possibility that, contrary to common assumptions, rhoptry secretion might be a Ca\(^{2+}\)-dependent process.

**RESULTS**

The *T. gondii* genome encodes three ferlin proteins. Next to TgDOC2, a series of BLAST searches of the *Toxoplasma* genome identified four additional proteins containing two or more C2 domains, of which three also contained a transmembrane domain. Two proteins had clear homology to the ferlin family of Ca\(^{2+}\)-sensitive membrane fusion proteins (Fig. 1A). We named these proteins TgFER1
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**FIG 1** (A) Toxoplasma encodes three ferlin proteins, and human otoferlin is shown for comparison. Ferlins are defined by 5 to 7 C2 domains (labeled A to F) and a C-terminal transmembrane (TM) domain, as well as typically a “FerI” domain of unknown function. TgFER3 contains an N-terminal signal peptide, which, in combination with the C-terminal TM domain, could signal glycosylphosphatidylinositol (GPI) anchor addition at the C terminus. Yellow shading in TgFER3 represents coiled-coil domains. Degenerate C2 domains are defined as having a P value below the cutoff in Pfam database searches. (B) Phylogenetic analysis of apicomplexan, chromerid, and human ferlins. Abbreviations are grouped as follows. Human ferlins: L1 to L6, FR1L1 to -6 (FR1L1, dysferlin [Q9HC10.3]; FR1L2, otoferlin [Q9NZM1.1]; FR1L4, A92123.1; FR1L5, A0A812.2; FR1L6, Q2WG9.2). Ox, green algae Ostreococcus tauri (Q01FJ7). Chromerids: Vb, Vitrella brassicaformis (VbFER1, Vbre_12074 plus Vbre_12075; VbFER2, Vbre_9118); and Cv, Chromera velia (CvFER1, Cvel_17519.2; CvFER2, Cvel_9223). Apicomplexa: Tg, Toxoplasma gondii (TgFER1, TGME49_309420; TgFER2, TGME49_260470; TgFER3, TGME49_295472 plus TGME49_295468); Nc, Neospora caninum (NcFER1, NCLIV_053770; NcFER2, NCLIV_026570; NcFER3, NCLIV_002280); Em, Eimeria maxima (EmFER1, EMWEY_00002120; EmFER2, EMWEY_00009280; EmFER3, EMWEY_00017650); Pf, Plasmodium falciparum (PfFER1, PF3D7_0806300; PfFER2, PF3D7_1455600); Pb, Plasmodium berghei (PbFER1, PBANKA_122440; PbFER2, PBANKA_131930); Cp, Cryptosporidium parvum (CpFER1, cgdb_2910; CpFER2, cgdb_2320); and Gn, Gregarina niphonii (GnFER1, Gni_063830; GnFER2, Gni_073830). Alignment and the unrooted Jukes-Cantor phylogenetic tree were generated in Geneious v.6.1.6 (drodes).
does not function in cell division or replication but is essential for completion of Toxoplasma’s lytic cycle.

TgFER2 does not localize to the micronemes. Myc-TgFER2 localization by IFA revealed a dispersed pattern not reminiscent of any Toxoplasma feature (Fig. 2D). Since membrane trafficking proteins are notoriously hard to accurately localize (e.g., synaptotagmin IV is observed on different membranes in the secretory pathway, dependent on cell type, condition, and fixation method [24–26]), we tested fixatives like paraformaldehyde, methanol, and acetone. Although the control SAG1 pattern varied per fixative, the Myc pattern remained unchanged (see Fig. S2A in the supplemental material). Because it is conceivable that TgFER2 localization changes upon egress and/or in response to a change in \([\text{Ca}^{2+}]_i\), we also probed extracellular parasites with the Ca\(^{2+}\) ionophore A23187. Under any extracellular condition, TgFER2 appears as defined cytoplasmic puncta suggestive of either inclusion bodies or a membranous structure of unknown identity (see Fig. S2B). This staining pattern differed from the more disperse intracellular signal, which is likely a result of the different physiological states (e.g., \([\text{Ca}^{2+}]_i\)) of the parasites under these conditions.

Since the transmembrane domain is predictive of membrane association, we resolved the localization by immunoelectron microscopy (IEM). In intracellular parasites, a comparatively small number of gold particles were distributed throughout the cytoplasm (Fig. 3A). Gold particles were notably enriched at the cytoplasmic side of the inner membrane complex (IMC) (Fig. 3A and B) and within the internal structures of the conoid at the apical end (Fig. 3A and C to E). In extracellular parasites, we observed a different pattern with gold particles patched on the cytoplasmic face of the parasite.
mic side of the rhoptries (Fig. 3F to H). The lack of consensus across multiple IFA and IEM experiments and the technical limitations of both methods restrain us from drawing any definitive conclusion about FER2 localization in the tachyzoites. Nonetheless, TgFER2 labeling was never observed on the micronemes, which was inconsistent with our initially hypothesized role for FER2 in microneme secretion.

TgFER2 is not required for microneme secretion and conoid extrusion. To unravel the lethality of TgFER2 depletion, we first assayed parasite egress. After 48 or 96 h of ATc treatment, FER2-cKD parasites egressed normally when treated with Ca\(^{2+}\) ionophore A23187 (Fig. S1B). This suggests that the micronemes are secreted normally. The morphology and distribution of micronemes in TgFER2-depleted parasites are also normal by IFA and transmission electron microscopy (TEM) (see Fig. S3 in the supplemental material). Since TgFER2 was detected in the conoid, we also examined conoid extrusion as another Ca\(^{2+}\)-regulated process (27). Figure S1C in the supplemental material shows that conoid extrusion is normal in the TgFER2-depleted mutant.

Next we directly tested microneme protein secretion through Mic2 release (28). Both untriggered, low-level constitutive secretion and Ca\(^{2+}\) ionophore-induced microneme secretion occurred normally in the absence of TgFER2 (Fig. 4A and B). It is now apparent that micronemes are not uniform and that distinct populations within the parasite contain different proteins (29). We therefore reasoned that TgFER2 might act differentially on these populations and that this might explain our observations. Mic2 is secreted from a Rab5a/c-dependent population of micronemes. Another component of

\[ \text{FIG 3} \quad \text{Subcellular localization of TgFER2.} \quad \text{Shown is immunoelectron microscopy of intracellular (A to D) and extracellular (F to H) Toxoplasma tachyzoites expressing an N-terminal Myc epitope-tagged TgFER2 from the endogenous locus under the TetO7sag4 promoter. In intracellular parasites, Myc antibodies direct gold particle clusters to the cytoplasmic side of the IMC, and there is a strong enrichment inside the conoid, but no strong association with micronemes and minor association with the rhoptries. In extracellular parasites, gold particles are predominantly observed in clusters on the cytoplasmic side of the rhoptry membranes next to localization inside the conoid. (Note the experimental variation: Myc antibody binding was not as efficient in extracellular parasites.)} \]

"R" marks the rhoptries, asterisks mark micronemes, and arrowheads mark gold beads in the conoid. Panel E is a magnification of the region marked in panel D. Scale bars are 250 nm.
this population, Mic10, was also secreted normally in TgFER2-depleted parasites (Fig. 4A). We examined secretion from the Rab5a/c-independent population Mic proteins 3, 5, 8, and 11 containing population by assessing Mic8 secretion by Western blotting. This proceeded normally in the absence of TgFER2. Labeling of Mic3, -5, and -8 by IFA on nonpermeabilized parasites (30, 31) also confirmed secretion of these proteins to the surface of both FER2-replete and -depleted parasites (Fig. 4C; see Fig. S4A and B in the supplemental material). Thus, secretion of all micronemes is TgFER2 independent.

Surface antigen SAG1 and Mic8 were deposited in trails behind parasites ± ATc, implying that TgFER2-depleted parasites are still motile (Fig. S4A and B). This was confirmed by scoring the total number of motile parasites and the type of motility displayed by individual FER2-cKD parasites by video microscopy (Fig. S4C). However, invading parasites, which are clearly identifiable by the stripping of Mic proteins off the apical invading parasite surface, were only observed in the presence of TgFER2 (Fig. 4C; Fig. S4A), suggesting a microneme-independent invasion defect.

**TgFER2 is required for host cell invasion.** We further examined host cell invasion through a series of invasion and attachment assays. As controls, we used mutants with defects at different points of host cell attachment and/or invasion. These include the TgDOC2 temperature-sensitive mutant (ts-DOC2) devoid of all microneme secretion (14), the calcineurin (CnA) mutant, which secretes micronemes normally but does not attach properly (9), the AMA1 mutant, which secretes micronemes but shows an increase in aborted invasion events due to failures in functional MJ formation (32, 33),
and the DHHC7 mutant, which lacks the palmitoyltransferase responsible for anchoring the rhoptries at the apical end and as a result is defective in rhoptry secretion (5).

Early events in parasite attachment are mediated by the binding of SAG proteins to glycans on the host cell surface. We assayed this by attachment of parasites to fixed host cells (30). Only the ts-DOC2 mutant, which completely fails to secrete micronemes, demonstrated reduced attachment relative to the wild-type and uninduced controls. Thus, all other mutants have normal microneme secretion (Fig. 5A). These data in general indicated that micronemal proteins support initial attachment, in the absence of MJ formation as that is not possible on fixed host cells.

Next we tested attachment and invasion by the “red-green invasion assay” (34). TgFER2-depleted parasites invaded at a much lower frequency (Fig. 5B). The defect intensified 3-fold upon prolonged TgFER2 depletion (96 h), which importantly did not affect their viability as these parasites are still fully capable of egress (Fig. S1B). As expected, all other mutants showed severe invasion defects. In this assay, quantification of the total number of
parasites per field allows for an estimation of parasite attachment. By this metric, a defect in the attachment of TgFER2-depleted parasites to host cells was observed. By 96 h of knockdown, the numbers of parasites attached to host cells dropped nearly 4-fold (Fig. 5B). This approaches the levels observed in the ts-DOC2 mutant, where attachment and invasion are both severely defective. The similarity between the ts-DOC2 mutant, where the primary defect is in attachment, and the DHHC7 mutant, with a penetration defect, highlights that this assay is unable to distinguish between these two interconnected phenotypes.

It has been observed that the motility and attachment dynamics of Toxoplasma are different under conditions of shear stress (35). To investigate whether these conditions might better clarify the phenotype of the TgFER2 knockdown, we measured the ability of FER2-cKD parasites ± ATc to adhere to human vascular endothelial cells (HUVECs) under flow. Depletion of TgFER2 led to a significant decrease in the number of parasites retained in the chamber (Fig. 5C; see Fig. S5 in the supplemental material), although attachment of TgFER2-depleted parasites was less compromised under flow relative to static conditions. This further underscores that TgFER2 is essential for invasion but does not pinpoint the nature of the defect. FER2-cKD parasites were then scrutinized for their interactions with host cells by video microscopy. Both wild-type and FER2-cKD parasites were able to glide across the host cells. In contrast to control parasites, TgFER2-depleted parasites were not able to invade host cells (Fig. 6; see Movie S1 in the supplemental material). Surprisingly, FER2-cKD parasites still exhibited “impulse motility” characteristic of invading parasites (36). This typical burst of forward motion immediately preceding invasion is followed by a momentary pause when parasites secrete the RONs and create the MJ before proceeding with invasion and subsequent parasitophorous vacuole formation. Both control and induced FER2-cKD parasites displayed bursts of impulse motility followed by a pause. However, only in control parasites was this pause followed by forward motion (invasion) at approximately half the original speed. In contrast, the velocity of induced parasites dropped essentially to 0 μm/s, and they failed to invade. This observation suggests that TgFER2 functions in the very late stages of invasion and indicates that the MJ either is not formed or is not of sufficient strength to support the force required for parasite penetration into cells.

TgFER2 is required for rhoptry secretion. As shown in Fig. 3 and Fig. S3 and S6 in the supplemental material, the localization, morphology, and apical anchoring of the rhoptries were not affected by TgFER2 depletion. Thus, FER2 is not required to tether the rhoptries at the apical end of the parasite. To test rhoptry function, we first monitored the release of rhoptry neck proteins by tracking RON4 distribution and assayng MJ formation (Fig. 7A). In the noninduced control, we readily observed MJ
formation, but no MJ formation was detected in the absence of TgFER2. These data suggest that the RON proteins are either not secreted, or if they are, they fail to assemble into the MJ.

We next performed “evacuole” assays to determine the quantity and quality of rhoptry protein secretion in TgFER2 mutants. Evacuoles, which are rhoptry protein clusters injected in the host cell cytosol, were visualized with ROP1 antiserum and classified as illustrated in Fig. 7B (33). We used the AMA1-cKD as control for partial rhoptry secretion and weak MJ strength (32, 33) and DHHC7-cKD parasites as a control for defective rhoptry secretion (5). To examine the strength of the overall parasite-host cell interaction, the number of parasites per field was counted and differentiated by whether they were associated with evacuoles (Fig. 7C). More specifically, the strength of the MJ itself can be measured by assaying the number of evacuoles per field and whether they are associated with parasites (Fig. 7D). Levels of rhoptry secretion were differentiated by the relative size of evacuole patterns (Fig. 7E): small punctate ROP1 staining indicates less secretion than long trails or clusters. For FER2-cKD parasites, the number of parasites per field was consistent with the observations from the attachment and invasion assays: depletion of TgFER2 decreased parasite attachment (Fig. 7C). Among the parasites that were attached, very few were associated with evacuoles, indicating that they have not secreted the rhoptries’ contents into the host cell. The TgFER2 data are comparable with the results for the DHHC7 mutant. However, they differed from parasites that lack AMA1, where few parasites attach but the majority of the parasites have secreted rhoptries and generated evacuoles (32, 33). TgFER2-

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**FIG 7** Rhoptry secretion of TgFER2-depleted parasites is impaired. (A) Formation of the MJ. A constant number of parasites across the various mutants and controls were incubated with host cells for 10 min. MJ formation was visualized with RON4 antiserum under semipermeabilizing conditions by 0.02% saponin. SAG1 stains the extracellular portion of the parasites. Arrowheads mark successfully invaded parasites that are not accessible to the SAG1 antibodies. Brightness and contrast adjustments were made identical for both conditions, and, thus, signals are directly comparable. (B) Representative examples of parasite and evacuole features scored in the evacuole assay represented in panels C to E. na, not applicable. (C to E) Evacuole assay to monitor rhoptry bulb secretion and assess stability of the MJ attachment. Parasites as indicated were grown under ATc for 48 h (+) or 96 h (++) and incubated with host cells for 10 min. Evacuole formation was visualized using ROP1 antiserum following paraformaldehyde fixation. n = 3 ± SD. For across-samples statistics for ±ATc (one-way ANOVA), correction is marked above the bar. Pairwise statistics for ±ATc (Student’s t test) are marked above the connector line. *, P < 0.01; **, P = 0.001; ***, P < 0.0001. Asterisk color corresponds with the variable compared across samples; black asterisks correspond with analysis of the total number of parasites. Indicated below the graphs in panels C, D, and E are the percentages of the total number of events recorded that represent association with the evacuole (evac.) or parasite (par.) or display trails/groups of evacuoles, respectively. The data presented in these panels are derived from the same experiments.

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depleted parasites, like the DHHC7 mutant, appear to secrete very few, if any, rhoptries. When we assess how many of the observed evacuoles are associated with parasites, it becomes clear that TgFER2 depletion is much more similar to DHHC7 depletion than to parasites lacking AMA1 (Fig. 7D). Finally, we observed relatively few extensive evacuole patterns for both TgFER2- and DHHC7-depleted parasites (Fig. 7E) and conclude that in the rare events of rhoptry secretion, very little material was released. Overall, we conclude that TgFER2 is required for the secretion of the rhoptries, which is necessary to invade host cells.

**DISCUSSION**

Micronemes and rhoptries are essential to the invasion of apicomplexan parasites. These fascinating cellular structures are likely derived from ancestral organelles that persist in modern predatory protozoa and were adapted during the evolution of the *Apicomplexa*’s intracellular, parasitic lifestyle (37). While the molecular details of the initiation of microneme secretion are incompletely understood, the critical role of intracellular Ca²⁺ fluxes has been known for decades. Far less is known about either the mechanisms of rhoptry secretion or the trafficking of their contents.

It has long been hypothesized that secretion from both micronemes and/or rhoptries requires a membrane fusion event, but evidence for a canonical secretion machinery has been elusive. Using C2 domains as the anchor for a bioinformatic search for potential components of this pathway, we were unable to find homologs for either synaptotagmins or the canonical DOC2 family of Ca²⁺ sensors that function in mammalian neurotransmitter release (18, 38). We did find orthologs of the ferlin family of Ca²⁺-sensing membrane fusion proteins. TgFER1 and TgFER2 are widely conserved across the *Apicomplexa*, whereas the degenerate TgFER3 is found only in the *Coccidia* and a single *Chromerid* species, illustrative of the ancient history of these processes.

Detailed studies of *Toxoplasma* FER2 demonstrated its requirement for secretion from the rhoptries. This finding provides one of the first mechanistic insights into rhoptry secretion, firmly linking it to the activity of this C2 domain-containing protein. It is generally accepted that rhoptry secretion must be preceded by microneme secretion and requires contact with an appropriate host cell (4, 39). However, neither the transduction of this attachment signal nor the process by which it leads to secretion of the organelle’s contents has been clarified. Although it is known that Mic8 is required for rhoptry release and has been postulated to be key in a signal transduction pathway (40), there are no experimental data supporting this model. Furthermore, AMA1 (33) and RON5 (41) also appear to be involved in rhoptry secretion, but these mechanisms are equally unknown. Our finding that the Ca²⁺ sensor TgFER2 is required for rhoptry secretion provides a tantalizing hint at the molecular mechanism. The presence of an Asp residues constellation consistent with Ca²⁺-binding capacity in TgFER2’s C2F domain supports this model (see Fig. S7 in the supplemental material). This is consistent with Ca²⁺ binding being restricted to the C2E and C2F domains in mammalian ferlins (21). We have as yet been unable to definitively demonstrate a role for Ca²⁺ in TgFER2 function specifically. However, ferlins are Ca²⁺-sensing proteins, and it is well established that a rise in [Ca²⁺], accompanies host cell invasion (2). Thus, while the conventional belief has been that this fluctuation acted only on activation of motility, conoid extrusion, and microneme secretion, our collective evidence provides a hint that rhoptry secretion may be similarly dependent on variations in [Ca²⁺]. The relative importance of the protein’s individual C2 domains in this process and their relative Ca²⁺-binding abilities are exciting open questions to be experimentally determined.

Of the mammalian ferlins, otoferlin is currently the best studied, yet its mechanism of action remains poorly understood (42). Otoferlin is expressed in many tissues, but in cochlear hair cells (CHCs) it controls the release of neurotransmitter upon an increase in [Ca²⁺]; (43, 44). A rise in [Ca²⁺]; leads otoferlin to interact with phospholipids (42) and SNARE proteins *in vitro* (45), although SNAREs have been debated to be absent from the site of secretion in CHCs (46). This highlights the potential for ferlin proteins to facilitate...
membrane fusion in the absence of SNAREs, an important parallel to *Toxoplasma*, in which there is currently no evidence for either rhoptry- or microneme-resident SNAREs.

As part of this study, we compared different invasion and egress mutants across several commonly used assays, which allowed for several important observations. First, the fixed host cell attachment data (MJ absent) demonstrate that microneme proteins contribute a large portion to the attachment strength. Somewhat unexpectedly, the red-green invasion assay did not differentiate the various mutants very well, with the exception of the partial attachment defect previously reported for the CnA mutant (9). Thus, this assay is not capable of specifically attributing individual phenotypes to defects in attachment versus invasion. In contrast, the vacuole assay was very powerful in differentiating different aspects of MJ formation and rhoptry secretion.

Overall, our findings support two interesting hypotheses. First, if ferlins act as Ca\(^{2+}\) sensors during Ca\(^{2+}\)-dependent secretion in the *Apicomplexa*, TgFER2 may represent the link between the previously observed Ca\(^{2+}\) fluctuations during invasion and the well-described mechanics of MJ formation. If on the other hand, the essential role of TgFER2 during rhoptry secretion is calcium independent, this would signify a fascinating evolutionary divergence from the canonical function of ferlins as Ca\(^{2+}\) sensors. While additional work will be required to distinguish between these models, the work presented here is a critical step in our understanding of these critical virulence processes.

**MATERIALS AND METHODS**

**Parasites and mammalian cell lines.** Transgenic derivatives of the RH strain were maintained in human foreskin fibroblasts (HFFs) as previously described (47). For the attachment assay under fluidic shear stress, HUVECs were cultured in EGM-2 medium containing EGM-2 SingleQuot supplements and growth factors (Lonza, Allendale, NJ). TgFER2 CDS was amplified using primers YFP-FER2-F/R and Nhel/EcoRV cloned into tub-YFPYPF(MCS)/sagCAT (48) to generate ptub-YFP-FER2/sagCAT, which was used for Sanger sequencing validation of the gene model. FER2-cKD parasites were generated by BglII/NotI cloning of PCR-amplified FER2 sequence (primers BamHI-FER2-F/NotI-FER2-R) into N-terminal Myc epitope-tagged plasmid derived from dihydrofolate reductase (DHFR)-TetO7/sag4-Nt-GOI (Wassim Daher, Université de Montpellier) and linearized by XbaI prior to transfection. ts-DOC2 parasites were generated by first 5xTY tagging the DOC2 locus using the PCR ampiclon (primers 5xTy_upstream_F/5xTy_PlusLink_R) from plasmid pLIC-5xTY-DD24/HX (Chris Tonkin, Walter and Eliza Hall Institute) and BglII/EcoRV cloning into tub-YFPYPF(MCS)/sagCAT. The tub promoter was Pmel/BglII replaced with the 3′ DOC2 homologous region PCR amplified from genomic DNA (gDNA [primers DOC2_3-target_F/R]). The CAT cassette was Pmel/NotI replaced with a DHFR minigene cassette and plasmid Nhel linearized prior to transfection. A CRISPR/Cas9 (clustered regularly interspaced short palindromic repeats with Cas9) plasmid was generated to mutate DOC2 F124 to S124 using primers DOC2_proto_F/R (49) and cotransfected with hybridized oligonucleotides DOC2_FM–>SV_F/R in RHΔKu80ΔHX-DOC2-5xTY parasites. All primer sequences are provided in Table S1 in the supplemental material.

**Imaging.** The following antisera were used: anti-Myc monoclonal antibody (MAb) 9E10, anti-SAG1 MAb DG52 (50), anti-Mic2 MAb 6D10 (51), mouse anti-AMA1 (33), rabbit anti-Mic3 (52), rabbit anti-Mic8 (53), and mouse anti-ROP1 (54). Alexa 488- or 594-conjugated secondary antibodies were used (Invitrogen). Images were collected on a Zeiss Axiosvert 200 M wide-field fluorescence microscope, and images were deconvolved and adjusted for phase contrast using Velocity software (Perkin Elmer).

**Egress assay.** The egress assay was performed as described previously (9, 14). Freshly lysed parasites, pretreated ± ATc for 24 h, were inoculated into HFF cells and incubated ± ATc for an additional 24 h. For 96 h, parasites treated ± ATc for 68 h were inoculated and incubated ± ATc for an additional 30 h. Egress was triggered by treatment with 2 μM A23187 or dimethyl sulfoxide (DMSO) at 37°C for 5 min, followed by IFA with rat anti-IMC3 (48). Intact vacuoles were counted for each sample in at least 10 fields, and the percentage of egress was calculated relative to the DMSO control.

**Attachment and invasion.** The combined attachment/invasion assay was performed as previously published (14, 34) with modifications described in reference 9. Tachyzoites treated ± ATc for the hour indicated (ts-DOC2 parasites incubated at 35 and 40°C) were added to host cells in a 96-well plate, centrifuged (28 × g, 3 min, room temperature), and allowed to invade for 1 h at 37°C. Noninvasive extracellular parasites were detected using Alexa 594-conjugated anti-SAG1 T41E5 (55). Following fixation and permeabilization, all parasites were visualized with Alexa 488-conjugated anti-SAG1 T41E5. At least 300 parasites were counted per sample.

**Attachment to fixed host cells.** Assay was performed as previously described (33). HFF confluent 96-well optical bottom plates were fixed with 3% formaldehyde (PFA) plus 0.06% glutaraldehyde for 5 min at 4°C, followed by overnight 0.16 M ethanolamine quenching at 4°C. Wells were prewashed with 0.2% bovine serum albumin (BSA) in Dulbecco’s modified Eagle’s medium (DMEM). Cytoplasmic yellow fluorescent protein (YFP)-expressing TATiΔKu80 parasites mixed in a 1:1 ratio were used as an internal control (9), centrifuged (28 × g, 5 min, 20°C) on the monolayer, and incubated for 30 min at 37°C. Wells...
were rinsed 3 times with phosphate-buffered saline (PBS), fixed with 4% PFA for 30 min at 4°C, and permeabilized with 0.25% Triton X-100 for 10 min. After blocking with 1% BSA in PBS, the parasites were probed with rabbit anti-green fluorescent protein (anti-GFP [Torrey Pines Biolabs]), and mouse anti-SAG1 DG52. Three random fields in 3 independent wells were counted.

**Attachment under fluidic shear stress.** Attachment under fluidic shear stress was performed as described previously (35, 56). Microfluidic channels containing fibronectin were coated overnight with HUVECs. Freshly lysed parasites treated ± ATc for 48 or 96 h were either stained with CMTPX CellTracker red or carboxyfluorescein succinimidyl ester (CFSE [Life Technologies]), counted, and combined 1:1. In each replicate experiment, the dyes were switched on the parental and knockdown parasite lines. Parasites were flowed at a shear force of 0.5 dyne/cm² for 10 min at 37°C and were fixed under flow conditions with 4% PFA for 30 min, followed by imaging on a Nikon Eclipse Ti microscope.

**Conoid extrusion assay.** The conoid extrusion assay was performed as published (27). Freshly lysed parasites grown ± ATc for 48 h were resuspended in 10% fetal bovine serum (FBS) in HS buffer. Conoid extrusion was induced using 0.5 M ethanol or 5 µM A23187 for 30 s. Parasites were fixed and scored for conoid extrusion by phase-contrast microscopy. Samples were counted blindly, scoring more than 350 parasites per sample.

**Microneme Mic2, Mic8, and Mic10 secretion by Western blotting.** Microneme Mic2, Mic8, and Mic10 secretion by Western blot was performed as published (31). Freshly lysed parasites treated ± ATc for 48 h were resuspended in DMEM/FBS and added to a 96-well round-bottom plate, and secretion was induced by 1 µM A23187 or DMSO for 5 min at 37°C. For constitutive microneme secretion, there was no stimulation at 37°C for 60 min. Supernatants were probed by Western blotting using MAB 6D10 anti-Mic2 (51), rabbit anti-Mic2 (53), rabbit anti-Mic8 (57), and MAB anti-Gra1 (58). Signals were quantified using a densitometer.

**Motility assessments.** Motility was analyzed by video microscopy essentially as described previously (14). Intracellular tachyzoites grown for 96 h ± ATc were physically harvested, resuspended in modified Ringer’s medium, and added to HFF confluent glass-bottom culture dishes (MatTek). The dish was imaged using a 63× objective at 37°C. Videos were recorded with 1-s intervals. Velocities of individual invasion events were analyzed using the ImageJ/FIJI Cell Counter plug-in.

**Moving junction formation.** Moving junction formation was determined as published (40) with previously described modifications (9). Parasites grown ± ATc for 48 h were inoculated into a HFF confluent 24-well plate by centrifugation (28 × g, 5 min, 20°C) and incubated at 37°C for 10 min. Wells were rinsed twice with PBS, fixed with 4% PFA at 4°C, and partly permeabilized with 0.02% saponin. MJ was detected using rabbit anti-ROP4 (7), and all parasites were detected following full permeabilization with MAb anti-SAG1 DG52.

**Evacuole assay.** Evacuoles were determined as described (59) with modifications. A total of 1 × 10⁷ parasites grown ± ATc for 48 h were inoculated into HFF confluent 24-well plates. The plate was centrifuged (28 × g, 15 min, 23°C) and incubated at 37°C for 10 min. Wells were rinsed twice with PBS, fixed with 4% PFA at 4°C, and 0.25% Triton X-100 permeabilized. Evacuoles were detected by MAb Tg49 anti-ROP1 (54). More than 100 events per sample per experiment were counted.

**Immunoelectron microscopy.** Following washing with PBS, overnight-infected HFF cells were fixed in 4% PFA in 0.25 M HEPES (pH 7.4) for 1 h at room temperature and then in 8% PFA in the same buffer overnight at 4°C. They were infiltrated, frozen, and sectioned as previously described (60). Sections were immunolabeled with anti-Myc 9E10 in 1% fish skin gelatin and then with goat anti-IgG antibodies, followed by 10-nm protein A-gold particles before examination with a Phillips CM120 electron microscope under 80 kV.

**Transmission electron microscopy.** Parasites were fixed in 4% glutaraldehyde in 0.1 M phosphate buffer (pH 7.4) and processed for routine electron microscopy (61). Briefly, cells were postfixed in osmium tetroxide and treated with uranyl acetate prior to dehydration in ethanol, treatment with propylene oxide, and embedding in Spurr’s epoxy resin. Thin sections were stained with uranyl acetate and lead citrate prior to examination with a JEOL 1200EX electron microscope.

**Statistics.** Student’s paired t test and one-way analysis of variance (ANOVA) using post hoc Bonferroni correction were used where indicated against the TaNΔKu80 line.

**SUPPLEMENTAL MATERIAL**

Supplemental material for this article may be found at [https://doi.org/10.1128/mBio.01510-18](https://doi.org/10.1128/mBio.01510-18).

FIG S1, TIF file, 0.1 MB.

FIG S2, TIF file, 3.3 MB.

FIG S3, TIF file, 2.8 MB.

FIG S4, TIF file, 2 MB.

FIG S5, TIF file, 0.1 MB.

FIG S6, TIF file, 5.1 MB.
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