Identification and Molecular Dissection of IMC32, a Conserved Toxoplasma Inner Membrane Complex Protein That Is Essential for Parasite Replication

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ABSTRACT The inner membrane complex (IMC) is a unique organelle of apicomplexan parasites that plays critical roles in parasite motility, host cell invasion, and replication. Despite the common functions of the organelle, relatively few IMC proteins are conserved across the phylum and the precise roles of many IMC components remain to be characterized. Here, we identify a novel component of the Toxoplasma gondii IMC (IMC32) that localizes to the body portion of the IMC and is recruited to developing daughter buds early during endodyogeny. IMC32 is essential for parasite survival, as its conditional depletion results in a complete collapse of the IMC that is lethal to the parasite. We demonstrate that localization of IMC32 is dependent on both an N-terminal palmitoylation site and a series of C-terminal coiled-coil domains. Using deletion analyses and functional complementation, we show that two conserved regions within the C-terminal coiled-coil domains play critical roles in protein function during replication. Together, this work reveals an essential component of parasite replication that provides a novel target for therapeutic intervention of T. gondii and related apicomplexan parasites.

IMPORTANCE The IMC is an important organelle that apicomplexan parasites use to maintain their intracellular lifestyle. While many IMC proteins have been identified, only a few central players that are essential for internal budding have been described and even fewer are conserved across the phylum. Here, we identify IMC32, a novel component of the Toxoplasma gondii IMC that localizes to very early daughter buds, indicating a role in the early stages of parasite replication. We then demonstrate that IMC32 is essential for parasite survival and pinpoint conserved regions within the protein that are important for membrane association and daughter cell formation. As IMC32 is unique to these parasites and not present in their mammalian hosts, it serves as a new target for the development of drugs that exclusively affect these important intracellular pathogens.

KEYWORDS inner membrane complex, Toxoplasma gondii, coiled-coil domain, endodyogeny, palmitoylation, parasitology

Toxoplasma gondii is a member of the phylum Apicomplexa, which includes many human pathogens, such as Plasmodium falciparum, the causative agent of malaria, and Cryptosporidium spp., a leading cause of diarrheal diseases (1–3). T. gondii is the most widespread member of the phylum, as it infects all mammals, including approximately one-third of the world’s human population (4). Most human infections are asymptomatic, but the parasite causes serious disease in immunocompromised individuals and congenitally infected neonates. Apicomplexan parasites share a number of unique organelles that are necessary for maintaining their intracellular lifestyles and causing disease (5). One of these organelles is the inner membrane complex (IMC), a
peripheral membrane and cytoskeletal system that underlies the parasite’s plasma membrane and plays central roles in motility, invasion, and replication (6).

The IMC is composed of a series of flattened membrane vesicles, known as alveoli, which are sutured together like a quilt, as well as a supporting cytoskeletal network of intermediate filament-like proteins called alveolins (7–9). In T. gondii, the alveoli are organized into three rows of rectangular plates along with a single cone-shaped plate called the apical cap at the apical end of the parasites (10). Detergent fractionation demonstrates that most IMC proteins localize either to the alveolar membrane or the cytoskeletal network, though many network-associated proteins are likely also tethered to the membranes (9, 11, 12). While some IMC proteins, such as the GAPM proteins, IMC25, and GAP40, are embedded in the alveoli via transmembrane domains (13–15), the majority are tethered to the IMC membrane by myristoylation and/or palmitoylation, which are carried out via a cytoplasmic N-myristoyl transferase (NMT) or a target membrane palmitoyl acyltransferase (PAT), respectively (16, 17). Two essential IMC-localized PATs that are responsible for modifying a growing number of known palmitoylated IMC proteins were recently characterized (18–22). While many of the known acylated IMC membrane proteins are dispensable for parasite survival (e.g., ISP1-4 and HSP20), the precise identity and functions of essential palmitoylated proteins are still not well understood.

The IMC plays several crucial roles in the lytic cycle of T. gondii. The IMC houses the glideosome, the actin-myosin motor that provides the force necessary for gliding motility and host cell invasion (23–25). The apical cap portion of the IMC has recently been shown to play an important role in organizing the conoid, a microtubule basket-shaped structure that is critical for the secretion of the micronemes for host cell adhesion, and for the rhoptries, which are necessary for host cell penetration (26–28). Importantly, the IMC also serves as the scaffold for forming daughter buds during endodyogeny, the internal budding process of replication in T. gondii tachyzoites (6). Endodyogeny begins with the duplication and segregation of the centrosomes and kinetochores, followed by subpellicular microtubule assembly, IMC formation, and daughter cell budding. Newly synthesized or replicated organelles are packaged into the developing IMC scaffold until budding is complete, at which point the maternal IMC is disassembled and the daughter cells emerge, adopting the maternal plasma membrane (29). Although all apicomplexan parasites are dependent on the IMC for their various internal budding processes, many components of the IMC machinery are not conserved across the phylum (11, 12, 30, 31).

In addition to trafficking to the correct organellar subdomain, the timing of expression and recruitment of IMC proteins to the daughter buds during endodyogeny also likely plays an important role in IMC protein functions. Some of the earliest expressed proteins include the AC9/AC10/ERK7 complex, which is essential for invasion, and F-box protein TgFBXO1, which plays an important role in replication (26–28, 32). These are followed by ISP1 and ISP3, which are dispensable and recruited independently of the cortical microtubules which drive bud development (21, 33). Some proteins, such as the cytoskeletal alveolins IMC3 and IMC6, are recruited to daughter buds later in the replication process, and other IMC proteins are exclusively recruited to the maternal IMC, after endodyogeny is complete (IMC7, -12, and -14) (9, 34). Although formation of the IMC involves precisely timed recruitment of different IMC components to specific subdomains, the exact contribution of each of these components and the importance of correct timing remain poorly understood.

Here we report the discovery of a novel IMC protein (IMC32), which we identified from our previous in vivo biotinylation (BioID) experiment using the early daughter apical cap protein AC9 as the bait (27, 28). We show that IMC32 is an early daughter IMC protein and characterize its timing of recruitment relative to other IMC components. We also find that IMC32 is a membrane-associated IMC protein that resides in the body portion of developing daughter buds and that localization is dependent on a strongly predicted N-terminal palmitoylation site. Surprisingly, we find that proper protein
trafficking also requires a series of conserved C-terminal coiled-coil (CC) domains. Using a conditional knockdown system, we demonstrate that IMC32 is essential for parasite survival and that its conditional depletion results in the complete collapse of the IMC. Finally, we use a series of truncations and internal deletions to dissect the coiled-coil domains to determine how these domains serve IMC32 function. These deletions show that two of the most conserved regions of the coiled-coils play essential roles in the ability of IMC32 to govern parasite replication. Taken together, we identify and characterize a novel IMC membrane protein that is absolutely essential for parasite replication.

RESULTS

**IMC32 is an early daughter IMC protein.** We previously generated a large data set of candidate IMC proteins from our BioID experiments using AC9 as the bait (28). Our top hits from this data set were filtered for proteins with cyclical expression patterns similar to known IMC proteins, negative genome-wide CRISPR screen (GWCS) scores, homology to known proteins, and potential acylations that may enable IMC membrane association (16, 35, 36). Using these criteria, TgGT1_232150 emerged as an interesting target for study, as it had a cyclical expression profile similar to AC9, as well as to ISP3, a daughter-enriched IMC protein that is palmitoylated and myristoylated (Fig. 1A) (21, 35). TgGT1_232150 also contains a strongly predicted N-terminal palmitoylation site at...
residue 7 and a $-4.32$ GWCS score (see Fig. S1A in the supplemental material) (36, 37). While the protein lacks homology to known proteins, other features include a series of 5 C-terminal coiled-coil domains and 14 phosphorylation sites identified by phosphoproteomics analyses (Fig. 1B; Fig. S1B and C) (38, 39). TgGT1_232150 appears to be conserved in the Apicomplexa, as orthologues are seen in Neospora, Sarcocystis, Eimeria, and Cystoisospora. In addition, BLAST analysis reveals a Plasmodium homologue, PF3D7_0717600, that has limited homology to TgGT1_232150 (2e$^{-14}$) (Fig. S2). While we cannot determine if these are true orthologues or are homologous proteins, PF3D7_0717600 also contains predicted N-terminal palmitoylation sites (residues 7 and 13) and a similar series of C-terminal coiled-coil domains (Fig. S2A to C). Intriguingly, PF3D7_0717600 is also predicted to be very important or essential in Plasmodium by a genome-wide piggyBac transposon screen (40).

To determine the localization of TgGT1_232150, we used CRISPR/Cas9 to tag the endogenous locus with a C-terminal 3×HA epitope tag (41). In nondividing parasites, the protein was difficult to detect and appeared to be spread throughout the cytoplasm (Fig. 1C). In replicating parasites, TgGT1_232150 localizes to the body of parasite daughter buds, as assessed by colocalization with the IMC body alveolin IMC6 (Fig. 1D) (9). Costaining with the apical cap marker ISP1 showed that TgGT1_232150 is excluded from the apical cap portion of the organelle (Fig. 1E) (21). In mid to late daughters, the localization of TgGT1_232150 appeared spotty and seemed to form discontinuous longitudinal stripes along the body of the developing daughter buds. To ensure that this was not due simply to low detection, we used a spaghetti monster OLLAS tag (sm_OLLAS) to increase the strength of the signal (42). We also costained for the IMC suture protein ISC6 to determine if this pattern colocalized with the sutures that tether the alveolar plates of the organelle together (12). Tagging IMC32 with sm_OLLAS substantially increased the brightness of the signal but did not change the localization pattern. We could not observe a definitive colocalization with ISC6, which is predominantly seen as transverse stripes in the developing daughter buds (Fig. S3). Based on these data, we concluded that TgGT1_232150 is a daughter-enriched IMC body protein and named it IMC32.

At the start of replication, IMC32 appears as five punctae arranged in a pentagon, prior to the appearance of IMC6, suggesting that it is expressed early during endodyogeny (Fig. 1F). To more precisely analyze the timing of IMC32 expression, we colocalized IMC32 with a number of previously described proteins which also appear early during replication. We first colocalized it with ISP1, an early daughter bud marker that is incorporated into the IMC before IMC6, and found that IMC32 is recruited to the forming daughter buds before ISP1 (Fig. 1G) (21). We then colocalized IMC32 with FBXO1 and AC9, which are components of the IMC apical cap that are recruited to daughter buds at the earliest stages of endodyogeny, and found that IMC32 is recruited to early daughter buds similar to these proteins (Fig. 1H and I). Lastly, we colocalized IMC32 with the centrosome marker CEP250 and found that the early buds were adjacent to the centrosome (Fig. 1J), again consistent with IMC32 being a component of the early daughter cell scaffold (43). From this, we concluded that IMC32 is a daughter-enriched IMC protein that appears very early during parasite replication, preceding recruitment of ISP1 but at approximately the same time as AC9 and TgFBXO1.

**IMC32 is essential and plays an important role in endodyogeny.** The negative GWCS score suggests that IMC32 is either important or essential for parasite survival (36). We were unable to disrupt the gene with CRISPR/Cas9 by using several approaches, indicating that the gene is essential. To use a conditional knockdown approach, we replaced the endogenous promoter with an anhydrotetracycline (ATc) transactivator-based regulation system, which allows for the knockdown of IMC32 transcription upon addition of ATc (IMC32cKD) (Fig. S4) (44). The promoter replacement included the introduction of an N-terminal Myc tag, which could not be detected for unknown reasons. This was not problematic, as the promoter replacement was done in a strain containing an
endogenous C-terminal 2×Strep-3×Ty tag. In spite of the presumed addition of the tag and the altered timing of the truncated SAG4 promoter in the regulatable system, IMC32 localization appeared similar to that of the endogenous tagged protein (Fig. 2B). After 24 h of ATc treatment, we found by Western blotting and immunofluorescence assay (IFA) that IMC32 protein levels had decreased to almost undetectable levels (Fig. 2A and B), supporting an efficient conditional knockdown of IMC32.

To assess the effects of the knockdown on parasite morphology, intracellular parasites were treated for 6 h with (+) and without (−) ATc, syringe lysed, and allowed to infect new monolayers for 24 and 40 h with and without ATc. At 24 h, we saw parasites with disrupted morphology, as assessed by staining with the apical cap marker ISP1 and the cytoskeletal alveolin IMC6 (Fig. 2C). Despite these severe morphological defects, many of the vacuoles continued to expand to the 40-h time point, forming masses of misshapen parasites with a large amount of nuclear material and collapsed or missing IMC components (Fig. 2D). This suggests that in the absence of IMC32, parasites mount continued attempts at replication in spite of the gross morphological defects incurred by the knockdown. Similar results were seen by staining for tubulin and the cytoskeletal apical cap marker AC9, both of which detect very early daughter buds as well as maternal parasites (Fig. 2E to H) (28). We also examined the centrosome in these masses using CEP250 and found multiple foci, further indicating continued attempts at replication (Fig. 2I and J). The defects in replication are ultimately lethal, as the knockdown parasites fail to form plaques in a standard plaque assay (Fig. 2K).

These data demonstrate that IMC32 plays a critical role in the development of the IMC and is essential for parasite viability.

**The IMC32 palmitoylation site is required for localization and function.** To better understand how IMC32 functions in replication, we complemented the knockdown using a 3×HA-tagged wild-type cDNA driven by the endogenous promoter, targeted to the uracil phosphoribosyltransferase (UPRT) locus. The wild-type complement (IMC32cKD + IMC32FL) successfully colocalized with the endogenous conditionally regulated copy of IMC32 (Fig. 3A). To assess the importance of the predicted IMC32 palmitoylation site, we also generated an IMC32 expression construct in which the cysteine at residue 7 was mutated to a serine (IMC32C7S). Disruption of the palmitoylation site resulted in mislocalization of the protein into scattered punctae throughout the cytoplasm, showing that this residue is critical for IMC localization (Fig. 3B).

We then conducted plaque assays with the three strains (IMC32cKD, IMC32cKD + IMC32FL, and IMC32cKD + IMC32C7S) and measured the mean plaque size with and without ATc. As expected, IMC32cKD parasites made large plaques without ATc and no plaques were formed upon ATc treatment (Fig. 3C and D). The IMC32cKD + IMC32FL line rescued the knockdown, as it was able to form plaques that were the same size with and without ATc, noting that ATc treatment alone has a minor effect on parasite growth (Fig. S5) (44). Surprisingly, IMC32cKD + IMC32C7S parasites in the presence of ATc were able to form very small plaques, with a mean plaque size only 1.7% the size of the plaques formed by IMC32cKD + IMC32FL plus ATc. Extending the plaque assays to 14 days showed that the IMC32cKD + IMC32C7S plaques do not get significantly larger, suggesting that these parasites fail to replicate further after forming small plaques. In agreement with this, IMC32cKD + IMC32C7S parasites were unable to be cultured in the presence of ATc, and multiple attempts to knock out the endogenous wild-type copy of the gene in this background failed. Therefore, despite making small plaques initially, the IMC32C7S parasites essentially cannot rescue the function of the IMC32 knockdown, demonstrating that the palmitoylation site is necessary for IMC32 localization and function.

**IMC32 localization and function are not dependent on phosphorylation.** Protein phosphorylation has been shown to play an important role in regulating the function of critical IMC components; thus, we assessed the 14 phosphorylation sites in IMC32 (Fig. 4A; Fig. S1B) (39, 45–48). To this end, we generated a construct with all 14 sites mutated to alanines and expressed it in the knockdown strain (IMC32cKD + IMC32MutPhos). We examined the localization of the phosphorylation mutant by IFA and
IMC32 is essential and plays an important role in endodyogeny. (A) Western blot showing ~97% depletion of IMC32-2×Strep3×Ty after 24 h of ATc treatment. Parasites were allowed to infect for 8 h before being treated with ATc. Mouse anti-Ty was used to detect IMC32, and mouse anti-ROP7 was used (Continued on next page)
found that localization was not affected (Fig. 4B). Because ATc treatment affects growth rates (Fig. S5), we compared the mutant to a strain expressing the wild-type copy of IMC32 in the presence of ATc and found no significant difference in plaque size (Fig. 4C).

We therefore conclude that these phosphorylation sites are not important for the function of IMC32.

**Coiled-coil 5 is not essential for IMC32 localization or function.** Given that the phosphorylation sites were dispensable, we suspected that the five coiled-coil domains at the C terminus of the protein might play an important role in IMC32 localization or function. Coiled-coil domains are supercoils of two or more alpha-helices that are involved in protein binding-dependent functions, such as vesicle transport and structural scaffolding, which may agree with a role of IMC32 in organelle formation (49). We were particularly interested in coiled-coils 1, 4, and 5, as these regions contain

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**FIG 3** The IMC32 palmitoylation site is required for localization and function. (A) Complementation with full-length IMC32:HA (HA) driven from its own promoter shows colocalization with endogenous 2×Strep3×Ty-tagged IMC32 (wild type [WT]). Red, rabbit anti-HA; green, mouse anti-Ty. (B) Mutagenesis of the predicted palmitoylation site (IMC32C7S) results in mistargeting to punctate spots in the cytoplasm. WT, endogenous 2×Strep3×Ty-tagged IMC32. Red, rabbit anti-HA; green, mouse anti-Ty. (C) Plaque assays showing that the IMC32 knockdown cannot make plaques at 8 days postinfection. Complementation with full-length IMC32 (IMC32cKD + IMC32FL) rescues plaque formation, while complementation with IMC32C7S results in very small plaques which do not appear larger from 8 to 14 days. Scale bar, 0.5 mm. (D) Quantification of plaque assays at days 8 and 14 showing that the small plaques formed at day 8 by IMC32C7S do not grow significantly larger by extending the plaque assay (ns [not significant], \( P > 0.05 \); **, \( P < 0.01 \)).
substantial amino acid identity to its malarial counterpart, suggesting that they may be important for IMC32 function (Fig. S2A).

To begin exploring these coiled-coil domains, we generated a truncated copy of IMC32 that removed just CC5 (residues 905 to 976), including a strongly conserved region in the *Plasmodium* homologue (residues 952 to 974) (Fig. 5A; Fig. S2) and asked whether the truncated protein could localize to daughter buds and effectively rescue the IMC32cKD knockdown. When expressed in the IMC32cKD line, the truncated protein (IMC32cc5) localized correctly to the IMC (Fig. 5B). Upon knockdown of the endogenous copy, plaque assays showed that growth was not significantly reduced compared to the strain complemented with IMC32FL (Fig. 5C). Thus, although part of CC5 is conserved, it is dispensable for IMC32 localization and function.

The conserved region of coiled-coil 4 is essential for IMC32 function. To assess the role of CC4, we first removed CC5 and most of CC4 by truncating residues 764 to 976 (IMC32cc4-5) and expressed this construct in the knockdown strain (Fig. 6A). IFA showed that IMC32cc4-5 still localized to daughter buds (Fig. 6A, ATc). However, we found that IMC32cc4-5 could not complement the knockdown, as seen by morphologically disrupted parasites that phenocopy the IMC32cKD line (Fig. 6A, +ATc). This result suggests that a region within CC4 is important for IMC32 function.

To further delimit important regions of CC4, we dissected the domain in three parts, designated A to C (Fig. 6B). We then generated a series of internal deletions within CC4 and analyzed whether the deletion constructs could rescue the IMC32cKD phenotype. We first deleted region C (residues 880 to 912), which lacks homology with PF3D7_0717600. The deletion construct (IMC32cc4c) successfully colocalized with endogenous IMC32 in daughter buds (Fig. 6C). Upon knockdown of the endogenous copy, we observed a significant reduction in plaque size compared to IMC32FL with ATc, indicating that region C of CC4 is an important but not essential region of CC4 (Fig. 6D). We next deleted region B (residues 816 to 880), which also lacks homology to PF3D7_0717600. This construct (IMC32cc4b) also correctly localized to the daughter buds in the IMC32cKD line and fully rescued the knockdown (Fig. 6E and F), revealing that region B is not important for IMC32 function. Finally, we assessed region A (residues

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**FIG 4** IMC32 localization and function are not dependent on phosphorylation. (A) Gene model for IMC32 with the 14 predicted phosphorylation sites mutated to alanine, plus a C-terminal 3×HA epitope tag. (B) IFA showing colocalization of endogenous IMC32 (WT) and the mutant phosphorylation copy of IMC32 expressed in the IMC32cKD + IMC32mutphos line. Red, rabbit anti-HA; green, mouse anti-Ty. Scale bar, 4 μm. (C) Quantification of plaque assays with IMC32cKD, IMC32cKD + IMC32cc5, and IMC32cKD + IMC32mutphos parasites under conditions with and without ATc reveals no significant (*P* > 0.05) difference in plaque sizes between IMC32cc5 and IMC32mutphos with ATc.

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764 to 816), which contains the highest similarity between T. gondii and P. falciparum (Fig. 6G; Fig. S2). We found that IMC32 ΔD localizes to daughter buds but completely fails to rescue the IMC32 knockdown (Fig. 6H and I), demonstrating that this conserved region in CC4 is essential for IMC32 function.

Coiled-coil 1 is strongly conserved and essential for protein function. CC1 also contains a region of substantial conservation in P. falciparum (Fig. 7A; Fig. S2). Thus, we generated two additional IMC32 truncations, one which removes all of the coiled-coils except CC1 (IMC32ΔCC1-5) and another which truncates all of the C-terminal coiled-coil domains (IMC32ΔCC2-5). IMC32ΔCC1-5 still localized correctly with the endogenous copy of IMC32, demonstrating that the N-terminal region of the protein plus CC1 is sufficient for trafficking (Fig. 7B). Surprisingly, in spite of containing the palmitoylation site that is sufficient for localization of other IMC proteins, IMC32ΔCC1-5 did not localize to the daughter buds and instead localized as scattered puncta throughout the parasite (Fig. 7C). To ensure that this was not a peculiarity of this single construct, we made two other constructs to test whether the first 144 or 239 amino acids of IMC32 were sufficient for IMC targeting, and both of these showed similar spotted cytoplasmic staining (Fig. S6).

We next generated an internal deletion which removed just CC1 and determined if this construct could localize to the daughter IMC and rescue the IMC32cKD phenotype. When analyzed by IFA, we saw that IMC32ΔCC1 could still target to the daughter IMC, suggesting that the remaining coils may also aid in trafficking (Fig. 7D). To assess whether CC1 was important specifically for the function of IMC32, we performed a plaque assay and found that IMC32ΔCC1 could not rescue the IMC32cKD phenotype (Fig. 7E). This confirmed that CC1 is essential for protein function, as suggested by its conservation with P. falciparum. Together, these data show that the palmitoylation site and at least one of the coiled-coil domains are required for trafficking to developing daughters but that the conserved CC1 is specifically essential for the function of IMC32.
The application of BioID in T. gondii has enabled a better understanding of the protein composition of the IMC (11, 12, 28, 46). These experiments have revealed a growing number of novel proteins with unique localizations, characteristics, and functions, demonstrating the complex nature of this apicomplexan-specific organelle. We originally identified IMC32 from BioID fused to AC9, which is cytoskeletal and restricted to the apical cap, rather than membrane associated and IMC body localized like IMC32.

**DISCUSSION**

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Although IMC32 and AC9 reside in distinct subcompartments of the IMC, both appear very early in emerging daughter buds, suggesting roles in the initial formation of the IMC. While it is unclear whether these proteins directly interact, biotinylation of IMC32 by AC9 may occur while the proteins reside in their final destinations (which are abutting one another) or labeling may occur during the trafficking of IMC components during endodyogeny.

The tethering of IMC32 appears to be dynamic: it localizes to the IMC of daughter buds but also shows diffuse staining throughout the cytoplasm after division, rather than localizing peripherally to the maternal IMC. A similar localization is seen in the palmitoylation site mutant IMC32C7S, suggesting that reversible palmitoylation may allow the protein to recycle into emerging daughters at each round of replication (50).

There are two palmitoyl acyltransferases in the IMC which have recently been characterized in detail, TgDHHC2 and TgDHHC14 (18). TgDHHC14 is restricted to daughter buds and is not present in the apical cap, similar to that seen for IMC32, which suggests that IMC32 may be a substrate of TgDHHC14. In addition, conditional knockdown of DHHC14 results in a similar disruption of IMC biogenesis. N-terminal acylation of other IMC proteins such as HSP20 (palmitoylation) and ISP1-3 (myristoylation and palmitoylation) is not only necessary but also sufficient for IMC membrane localization (19–22). However, we show that deletion of coiled-coils eliminates IMC32 trafficking, indicating that the palmitoylation site alone is not sufficient for localization. Although we cannot exclude the possibility that our deletion constructs (Fig. 7C; Fig. S6) caused gross misfolding of the protein, IMC32 localization appears to be dependent on both palmitoylation and its C-terminal coiled-coil domains.

While IMC32 plays a critical role in parasite replication, the precise mechanism by which it functions and how it is regulated remain to be determined. The continued expansion of the parasites upon IMC32 knockdown indicates that the parasites are not arresting in the cell cycle. Thus, the resulting phenotype is likely independent of the essential TgCyclins and TgCrk factors that serve to regulate the progression of cell division and whose disruptions result in growth arrest (6, 51). Whether or not IMC32 is

![FIG 7 Coiled-coil 1 is strongly conserved and essential for protein function. (A) Alignment showing similarity of IMC32 residues 522 to 549 with its Plasmodium homologue, PF3D7_0717600. Black highlights indicate identity; gray highlights indicate similarity. (B) Gene model for IMC32ΔCC2-5 and localization of IMC32ΔCC2-5 analyzed via IFA. IMC32ΔCC2-5 colocalizes with the endogenous copy of IMC32 (WT) in daughter buds. Red, rabbit anti-HA; green, mouse anti-Ty. (C) Gene model for IMC32ΔCC1-5. IMC32ΔCC1-5 fails to localize to the IMC of daughter buds and instead is localized to punctae throughout the cytoplasm. Red, rabbit anti-HA; green, mouse anti-Ty. (D) Gene model of IMC32ΔCC1 and IFA showing IMC32ΔCC1 localization to daughter buds. Red, mouse anti-HA antibodies; green, Rabbit anti-Ty. (E) Quantification of plaque assays with IMC32cKD, IMC32cKD + IMC32ΔCC1-5, and IMC32cKD + IMC32ΔCC2-5 parasites under conditions with and without ATc. IMC32ΔCC1-5 parasites were unable to form plaques (P < 0.0001). Scale bars for all images, 4 μm.

(27, 28). Although IMC32 and AC9 reside in distinct subcompartments of the IMC, both appear very early in emerging daughter buds, suggesting roles in the initial formation of the IMC. While it is unclear whether these proteins directly interact, biotinylation of IMC32 by AC9 may occur while the proteins reside in their final destinations (which are abutting one another) or labeling may occur during the trafficking of IMC components during endodyogeny.
regulated posttranslationally is also unknown. The simultaneous mutagenesis of all 14 predicted phosphorylation sites did not affect IMC32 function. This result does not exclude the possibility that phosphorylation of other sites might be important in regulating IMC32 or that a phosphomimetic mutant could affect its function (46, 48).

The C-terminal group of five coiled-coil domains in IMC32 suggests that this region is likely to mediate protein-protein interactions (49). Coiled-coils have been shown to be essential for GAP45, likely enabling interaction with other components of the glideosome (14). Coiled-coil domains also appear to be enriched in the apical annulus proteins that are embedded in the IMC sutures (52), as well as the conoid, which is cytoskeletal and IMC associated (53). Our dissection of the coiled-coil domains of IMC32 demonstrated that CC1 and CC4, which are most strongly conserved within P. falciparum, are essential for protein function. It is likely that these regions mediate interactions with other IMC proteins or are responsible for forming dimers or trimers of the protein. Interestingly, while CCS is relatively well conserved and dispensable, the region just upstream of this (region C of CC4) is not conserved and necessary for optimal parasite growth. Ultimately, the identification of protein partners or structural information of the protein will provide a better understanding of how IMC32 mediates its critical role in parasite replication.

While the central functions of the IMC in motility, invasion, and replication are conserved in apicomplexans, surprisingly few of its constituents are conserved from T. gondii to P. falciparum. Conserved elements include components of the glideosome that are critical for motility and the alveolins, which make up the cytoskeletal scaffold of the organelle (8, 54, 55). In contrast, many of the membrane components appear to be unique in each organism, suggesting that these players have specialized functions within their host cells. Furthermore, most of the proteins that have been characterized are frequently expected to be dispensable based on genome-wide functional studies (11, 12, 21, 32, 36). One early daughter protein that is important for parasite replication is TgFBXO1 (32). However, TgFBXO1 localizes to the apical cap region and does not appear to be essential for parasite survival. In contrast, IMC32 localizes to the IMC body and is absolutely essential for replication, as its conditional depletion results in a lethal collapse of the IMC. Along with its early timing of expression and conservation, this suggests that IMC32 may be a foundational component of the IMC membranes during daughter bud formation. Together, this study broadens our understanding of the T. gondii IMC by revealing a conserved lynchpin of the organelle that presents a novel target for the development of therapeutics across the apicomplexan phylum.

MATERIALS AND METHODS

Toxoplasma and host cell culture. Parental T. gondii RHΔhxgprt and subsequent strains were grown on confluent monolayers of human foreskin fibroblasts (HFFs; ATCC, Manassas, VA) at 37°C and 5% CO2 in Dulbecco’s modified Eagle medium (DMEM) supplemented with 5% fetal bovine serum (Gibco), 5% Cosmic Calf serum (HyClone), and 1× penicillin-streptomycin-glutamine (Gibco). Tetracycline (Tet)-free medium for knockdown lines used 10% tetracycline-free fetal bovine serum (Omega Scientific). Constructs containing selectable markers were selected using 50 μg/ml mycophenolic acid-xanthine (HXGPRT), or 5 μM pyrimethamine (56–58). Homologous recombination to the UPRT locus was negatively selected using 5 μM 5-fluoro-deoxyuridine (59).

Antibodies. The hemagglutinin (HA) epitope was detected with mouse monoclonal antibody (MAb) HA.11 (BioLegend, San Diego, CA) or rabbit polyclonal antibody (pAb) anti-HA (ThermoFisher). The Ty1 epitope was detected with mouse MAb BB2 (60). Toxoplasma-specific antibodies include mouse MAb anti-ISP1 (13), rabbit pAb anti-IMC6 (61), and mouse MAb anti-ROP7 (1B10) (62).

IFA and Western blotting. For immunofluorescence assays (IFA), confluent HFFs grown on coverslips were infected with T. gondii. After the time periods noted in the text above, the coverslips were fixed with 3.7% formaldehyde and processed for indirect immunofluorescence as described previously (63). Primary antibodies were detected by species-specific secondary antibodies conjugated to Alexa Fluor 488/594 (ThermoFisher). Coverslips were mounted in Vectashield (Vector Labs, Burlingame, CA), viewed with an Axio Imager.Z1 fluorescence microscope (Zeiss), and processed with ZEN 2.3 software (Zeiss).

For Western blotting, intracellular parasites were lysed in 1× Laemmli sample buffer with 100 mM dithiothreitol (DTT) and boiled at 100°C for 10 min. Lysates were resolved by SDS-PAGE and transferred to nitrocellulose membranes, and proteins were detected with the appropriate primary antibody and corresponding secondary antibody conjugated to horseradish peroxidase. Chemiluminescence was induced...
using the SuperSignal West Pico substrate (Pierce) and imaged on a ChemiDoc XR+ system (Bio-Rad, Hercules, CA). Western blot signal was quantified with ImageLab 5.2.1 software.

**Endogenous epitope tagging.** For C-terminal endogenous tagging, a pu6-Universal plasmid containing a protospacer against the 3' untranslated region (UTR) of IMC32 approximately 100 bp downstream of the stop codon was generated using primers P1/P2, as described previously (41). A homology-directed repair (HDR) template was PCR amplified using the Δκu80-dependent LIC vectors p3-HA.LIC-DHFR and p2-Strep-3×Ty.CAT that include the epitope tag, 3' UTR, and selection cassette (64). The 60-bp primers (P3/P4) (Table S1) include 40 bp of homology immediately upstream of the stop codon or 40 bp of homology within the 3' UTR downstream of the CRISPR/Cas9 cut site. This template was amplified in a total of 400 μl, purified by phenol-chloroform extraction, precipitated in ethanol, and electroporated into RHΔhxgprt parasites, along with 100 μg of the sequence-verified pu6-Universal plasmid. Transfected cells were allowed to invade a confluent monolayer of HFFs, and appropriate selection was applied. Successful tagging was monitored by IFA, and clonal lines of tagged parasites were obtained through limiting dilution. AC9, TgFBXO1, CEP250, and ISC6 were tagged using CRISPR/Cas9 with primers P32-P47 as described previously (12, 28, 32, 43).

**Endogenous promoter replacement.** For replacing the endogenous promoter with a Tet-regulatable one, a pu6-Universal plasmid containing a protospacer against the 5' UTR of IMC32 approximately 100 bp upstream of the start codon was generated using primers P15/P16 (41). A homology-directed repair (HDR) template was PCR amplified using the vector pT8TAT1-HX-Tet-mycNtCOR (65), including a transactivator (tTA), seven corresponding operators (TetO), an HXGPRT selection cassette, and a 1×Myc tag. The 60-bp primers (P17/P18) include 40 bp of homology 100 bp upstream of the CRISPR/Cas9 cleavage site or 40 bp of homology immediately downstream of the start codon. This template was amplified in a total of 400 μl, purified by phenol-chloroform extraction, precipitated in ethanol, and electroporated into RHΔhxgprt parasites already endogenously tagged with 2×Strep-3×Ty.CAT, along with 100 μg of the sequence-verified pu6-Universal plasmid. Transfected cells were allowed to invade a confluent monolayer of HFFs in Tet-free medium, and appropriate selection was applied the following day. Successful tagging was monitored by IFA, and clonal lines of tagged parasites were obtained through limiting dilution.

**Complementation of IMC32cKD with IMC32HA.** The majority of the IMC32 coding region was PCR amplified from genomic DNA using primers P9/P10 and ligated together with annealed oligonucleotides P7-P8 to produce the entire coding region. This was then cloned into puPRTKO-IS3HA (11) using NotI/BglII (all enzymes purchased from NEB). The endogenous promoter was amplified from genomic DNA and inserted with NsiI/BglII in front of the coding sequence, resulting in puPRTKO-IMC32HA. This complement vector was then linearized with DraIII and transfected into IMC32cKD parasites along with a universal pu6 that targets the UPRT coding region. This was followed by selection with 5 μg/ml 5-fluorodeoxyuridine for replacement of UPRT as described previously (59). IMC32HA-expressing clones were screened by IFA, and an HA-positive clone was designated IMC32HA. For mutation of the predicted palmitoylation site at position 7, puPRTKO-IMC32HA was used as a template for amplification with primers P13/P14. The resulting partial coding sequence was cloned into BglII/Hpal-digested puPRTKO-IMC32HA. The same processes for linearization, transfection, and selection were followed as described for IMC32HA. Clones were screened by IFA, and an HA-positive clone was designated IMC32HA.C75. For mutagenesis of the phosphorylation sites, a synthetic gene which simultaneously changed all of the phosphorylated serines and threonines to alanine was generated.

For IMC32 C-terminal truncations, puPRTKO-IMC32HA was used as a template to amplify truncations of the coding region. Primers P19/P20 were used for deletion of CC5 (ΔCC5), P19/P21 for ΔCC4-5, P19/P22 for ΔCC2-5, and P19/P23 for ΔCC1-5. The inserts were then cloned into a NotI-digested puPRTKO-IMC32HA. For IMC32 internal deletions, puPRTKO-IMC32HA was used as a template for the Q5 site-directed mutagenesis kit (NEB). Using the online NEBaseChanger tool (http://nebasechanger.neb.com/), appropriate primers were designed for each internal deletion. Primers P24/P25 were used for deletion of residues 764 to 816 (Δ764-816), P26/P27 were used for Δ816-880, P28/P29 were used for Δ880-912, and P30/P31 were used for ΔCC1. The same processes for linearization, transfection, and selection were followed for all truncation and internal deletion constructs.

**Plaque assay.** Six-well plates were seeded with HFFs and allowed to reach confluence. Serial dilutions (100, 200, and 300 parasites/well) of individual strains were used to infect the six-well plates and allowed to form plaques for 8 days, except where stated otherwise. For conditional knockdown of IMC32, ATc was added to the Tet-free medium (1 μg/ml) prior to the addition of parasites. Eight days after infection, HFF monolayers were fixed in 100% methanol for 3 min, washed with phosphate-buffered saline (PBS), and stained with crystal violet for visualization. The area of 50 plaques per condition was measured using ZEN software (Zeiss). Each plaque assay was performed a single time, because the IMC32 knockdown was lethal and we were assessing whether any level of rescue could be obtained. Statistical significance was calculated using a two-sample two-tailed t test. Box-and-whisker plots were generated using Prism GraphPad. The borders of the box represent the first quartile, the median, and the third quartile, while the whisks represent minimum and maximum values.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, TIF file, 0.3 MB.

**FIG S2**, TIF file, 0.7 MB.

**FIG S3**, TIF file, 0.3 MB.
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