Dynamics of the *Toxoplasma gondii* inner membrane complex

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**ABSTRACT**

Unlike most cells, protozoa in the phylum Apicomplexa divide by a distinctive process in which multiple daughters are assembled within the mother (schizogony or endodyogeny), using scaffolding known as the inner membrane complex (IMC). The IMC underlies the plasma membrane during interphase, but new daughters develop in the cytoplasm, as cytoskeletal filaments associate with flattened membrane cisternae (alveolae), which elongate rapidly to encapsulate subcellular organelles. Newly assembled daughters acquire their plasma membrane as they emerge from the mother, leaving behind vestiges of the maternal cell. Although the maternal plasma membrane remains intact throughout this process, the maternal IMC disappears— is it degraded, or recycled to form the daughter IMC? Exploiting fluorescently tagged IMC markers, we have used live-cell imaging, fluorescence recovery after photobleaching (FRAP) and mEos2 photoactivation to monitor the dynamics of IMC biogenesis and turnover during the replication of *Toxoplasma gondii* tachyzoites. These studies reveal that the formation of the *T. gondii* IMC involves two distinct steps—de novo assembly during daughter IMC elongation within the mother cell, followed by recycling of maternal IMC membranes after the emergence of daughters from the mother cell.

**KEY WORDS:** Apicomplexan parasites, Endodyogeny, Schizogony, Inner Membrane Complex, *Toxoplasma gondii*, Plasmodium, FRAP, Photoactivation

**INTRODUCTION**

The phylum Apicomplexa comprises thousands of obligate protozoan parasites (Levine, 1970), including clinically significant pathogens, such as the *Plasmodium* parasites responsible for malaria (Snow et al., 2005) and *Toxoplasma gondii*—a ubiquitous human pathogen affecting ~30% of the population worldwide (Pappas et al., 2009). These parasites replicate rapidly in the tissues of susceptible individuals, and pathogenesis is largely a consequence of uncontrolled proliferation (Tenter et al., 2000; Weatherall et al., 2002).

Unlike most cell biological systems where replication has been studied in detail (including bacteria and archaea, as well as animals, plants and fungi), Apicomplexans do not divide by binary fission. Rather, these parasites replicate using a distinctive mechanism in which multiple progeny are assembled within the mother (Hepler et al., 1966; Senaud and Cerná, 1969; Sheffield and Melton, 1968). This unusual process is termed schizogony when daughter nuclei are formed before membrane assembly or endopolygeny when daughter nuclei and membranes develop in parallel (Ferguson et al., 2008). *T. gondii* tachyzoites exhibit a minimal form of endopolygeny, assembling only two daughters within each mother (endodyogeny). These parasites are also readily cultivated *in vitro*, making *Toxoplasma* a useful model system for exploring the biology and mechanism of Apicomplexan parasite replication.

Central to the process of Apicomplexan replication is a membrane–cytoskeletal scaffolding known as the inner membrane complex (IMC) (Hu et al., 2002a; Sheffield and Melton, 1968). Flattened vesicles [cortical alveoli—the major morphological feature unifying the superphylum Alveolata (Adl et al., 2005; Moore et al., 2008)] are positioned immediately beneath the plasma membrane, giving the appearance of a triple membrane (Foussard et al., 1990; Vivier and Petitprez, 1969), which is sometimes called the parasite ‘pellicle’. The outer leaflet of the IMC anchors the actin–myosin motor complex that is required for motility and invasion (Dobrowolski et al., 1997; Frien et al., 2010; Menard, 2001), whereas the cytoplasmic side is intimately associated with the subpellicular microtubules and alveolins (intermediate-filament-like proteins) that give the parasite its shape (Mann and Beckers, 2001; Morrisette et al., 1997; Nichols and Chiappinio, 1987). Disrupting IMC organization dramatically alters pellicle integrity, cell shape and invasion competence (Khater et al., 2004; Stokkermans et al., 1996; Tremp et al., 2008).

The IMC is also highly dynamic, and its spatial and temporal organization is thought to be crucial for parasite development and replication. At the onset of daughter cell formation, new IMC complexes assemble within the cytoplasm and elongate rapidly, coordinating the segregation of subcellular organelles according to a strict schedule (Nishi et al., 2008). Newly assembled daughters, delimited by the IMC, ultimately emerge from the mother cell, picking up the maternal plasma membrane and sloughing off any residual maternal material (Sheffield and Melton, 1968). Many studies have focused on the cytoskeletal components of the IMC, and several Apicomplexan-specific IMC membrane proteins have been identified (Beck et al., 2010; Bullen et al., 2009; Fung et al., 2012), but our knowledge of alveolar membrane function remains incomplete (Harding and Meissner, 2014). Where does the IMC come from, and how is its assembly and turnover regulated? How does the IMC interact with other organelles during daughter parasite assembly? Exploiting a fluorescently tagged integral membrane protein as a reporter, we have used live-cell imaging and fluorescence recovery after photobleaching (FRAP) to monitor the dynamics of IMC biogenesis and turnover during *Toxoplasma gondii* tachyzoite replication.

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RESULTS

GAP40 permits the visualization of IMC membrane dynamics during parasite replication

Previous studies on the replication of Apicomplexan parasites have defined the IMC as a valuable morphological marker for tracking the cell cycle, including the assembly of daughter parasites (Hu et al., 2002a; Kono et al., 2012; Nishi et al., 2008). These studies focused on alveolins, such as the IMC1 protein – intermediate-filament-like molecules associated with the inner face of the IMC. In order to understand IMC membrane dynamics, we have employed GAP40, an integral IMC protein with nine predicted transmembrane domains, which is also a component of the glideosome protein complex responsible for parasite motility (Frénal et al., 2010). The Ku80 system (Fox et al., 2009; Huynh and Carruthers, 2009) was used to engineer allelic replacements expressing GAP40–YFP at the endogenous locus in RH strain of T. gondii parasites.

GAP40–YFP localizes uniformly throughout the parasite pellicle, including the apical and basal ends, as illustrated in Fig. 1 (see supplementary material Fig. S1 and Movie 1 for time-lapse imaging of living parasites). This contrasts with the localization of IMC1 in several significant ways. First, GAP40 labels the full length of the IMC, whereas IMC1 is excluded from the apical and basal ends (cf. Fig. 1A; supplementary material Fig. S1, merged). Second, GAP40 associates with developing daughter parasites earlier during the replicative cycle than IMC1 (cf. Fig. 1A–ii; supplementary material Fig. S1). Third, maternal GAP40 remains clearly visible throughout the process of daughter parasite emergence from the mother, in contrast to IMC1, which can be difficult to visualize, particularly at late stages (cf. Fig. 1Av; supplementary material Fig. S1).

We have used the pattern of GAP40 immunofluorescent signal to define several distinct stages in IMC development, as shown by colored text in Figs 1 and 2, and in cartoon form in Fig. 1C. The precise timing of these stages can be determined by live-cell imaging, as shown in Fig. 2A (and the corresponding time-lapse supplementary material Movie 1). Only maternal IMC is visible during interphase (i.e. prior to the initiation of daughter cell assembly, centrosome duplication, Golgi and apicoplast elongation and partitioning).

The first stage of IMC development, initiation of daughter IMC assembly (defined as t=0 throughout this report, and labeled in blue) occurs just apical to the maternal nucleus, adjacent to the microtubule organizing center (MTOC) and Golgi apparatus (Nishi et al., 2008; see also below). As noted above, GAP40 associates with the developing daughter scaffold before the appearance of IMC1. The second stage, daughter elongation (shown in aqua) proceeds basally over the subsequent 3 h, and is defined by growth of the IMC membrane (GAP40) and cytoskeleton (e.g. IMC1). During elongation, maternal organelles are partitioned between daughter parasites, progressively incorporating the centrosome, Golgi, apicoplast and nucleus (with the associated endoplasmic reticulum) (Nishi et al., 2008). The third stage, the emergence of daughter parasites (magenta) occurs by budding from the mother cell ~180–210 min after initiation, providing daughters with their plasma membrane, as indicated by colocalization with the parasite surface antigen SAG1 (Fig. 1B,C). During the final stage, daughter parasites continue to grow and mature for ~2 h after emergence from the mother (maturation, red). Daughters remain connected by a narrow cytoplasmic bridge, which is ultimately lost as maternal material is degraded, recycled or left behind in the residual body.

![Fig. 1. Stages of the IMC cycle in T. gondii. Colocalization of the integral membrane protein GAP40 (green, A and B) with the cytoskeletal alveolin IMC1 (red, A) and the plasma membrane protein SAG1 (red, B) at various stages throughout the T. gondii replicative cycle. Shown are selected images from fixed specimens, approximately corresponding to times t=0 (i), 45 min (ii), 120 min (iii), 180 min (iv), 200 min (v), 225 min (vi) and 360 min (vii). Arrows indicate the initiation of GAP40 assembly prior to that of IMC1; arrowheads show that maternal GAP40 disappears from the residual body during maturation. (Note that fixed samples provide better resolution and enable colocalization with SAG1 to be performed; see supplementary material Fig. S1 for time-lapse imaging of GAP40 and IMC1 in living cells.) (C) Cartoon illustrating the relative location of the IMC (green) and the parasite plasma membrane (black), along with the various stages defined in the text – initiation, elongation, emergence and maturation (the color coding of the cartoons is maintained throughout all figures). Scale bars: 5 μm.](image-url)
Quantitative analysis of GAP40–YFP fluorescence from multiple parasites over time (Fig. 2B; see also supplementary material Fig. S2 and Table S1 for raw data) demonstrates that this process is highly regular in parasites dividing with normal kinetics. The total fluorescence of developing daughters increases linearly over time, from the earliest stages of initiation (defined as $t=0$, fluorescence $=0$), through elongation and even after emergence from the mother cell, during the final stages of maturation. At $\sim 5$ h after initiation (2 h after emergence), fluorescence intensity reaches a plateau, which is maintained until the beginning of a new replicative cycle.

**Daughter IMCs are assembled de novo during elongation**

Morphological studies provide useful markers for the parasite cell cycle, but reveal little about the origin of the IMC. As a patchwork of flattened vesicles (Dubremetz and Torpier, 1978; Morrissette et al., 1997; Torpier et al., 1991), it has long been assumed that the IMC must be derived from the Golgi (Agop-Nersesian et al., 2010). Treatment with brefeldin A (BFA) arrests daughter IMC development (Fig. 2C; DeRocher et al., 2005), but GAP40 does not colocalize precisely with the Golgi apparatus (Fig. 2D, especially insets at $t=0$ and 60 min). GAP40 is more closely associated with a VP1+ (vacuolar protein 1) proM2AP+ Rab5+ organelle known as the endosome-like compartment (ELC) (Tomavo et al., 2013). This organelle is probably comparable to the recycling endosome, or to the pre-vacuolar compartment in plants (Jackson et al., 2013; Robibaro et al., 2002); see below for further discussion.

In order to examine the dynamics of IMC assembly, we tracked parasite replication and GAP40 fluorescence after selectively bleaching either the maternal or daughter IMC. As shown in Fig. 3A (and supplementary material Movie 3), daughter parasite fluorescence appears and elongates with normal kinetics even after the maternal IMC is completely bleached, indicating that the IMC must be synthesized de novo (Fig. 3B). No fluorescence recovery was observed in the maternal IMC, indicating that newly synthesized GAP40 is added to daughter IMC scaffolds only. GAP40 is unable to diffuse in mature parasites, as no recovery was observed after bleaching only a portion of the maternal IMC (Fig. 3C). In order to assess the movement of membrane proteins throughout the process of IMC assembly, we bleached the daughter IMC at various stages during elongation and monitored fluorescence recovery, as shown in Fig. 4. Daughter IMCs that are bleached early during elongation (Fig. 4A) recover rapidly, becoming uniformly fluorescent over their entire length, suggesting that newly synthesized IMC is added throughout the entire daughter IMC (or is able to diffuse within the plane of the membrane, or both). Bleaching a portion of the daughter IMC slightly later during the process of elongation results in only partial fluorescence recovery (Fig. 4B). Comparing images from 170 and
160 min shows a slight decline in apical fluorescence and partial fluorescence recovery, but the apical end remains more brightly stained, indicating that GAP40 movement is limited. These data imply that GAP40 is highly mobile at early stages of elongation, but that its mobility declines over time, concomitant with assembly of the membrane–cytoskeletal scaffold.

Parasites were also transiently transfected with GAP40 fused to mEos2 (Baker et al., 2010; McKinney et al., 2009), a photoconvertible fluorescent reporter that undergoes a conformational change when excited at 405 nm, resulting in a shift from green to red fluorescence. Photoactivation causes a portion of GAP40–mEos2 to fluoresce red (Fig. 5A,B), allowing it to be followed by time-lapse microscopy, analogous to a pulse-chase experiment. During the process of elongation, both red and residual green fluorescence remained essentially constant in maternal parasites, as no additional GAP40 is added to the maternal IMC and maternal GAP40 does not redistribute to daughter parasites (Fig. 3). However, in the developing daughters, whereas green fluorescence increased due to the addition of new material (see above, e.g. Fig. 2), red fluorescence declined, as photoactivated GAP40 disseminated throughout the developing daughter IMC membrane (180 min time-points and Fig. 5C). Total red fluorescence in daughter parasites remained constant but fluorescence intensity per unit of IMC membrane decreased proportionate to the growth of developing daughters.

Maternal IMC is not left behind in the residual body, but is internalized and incorporated into daughter parasites during maturation

As daughter parasites emerge, unsegregated maternal material is sloughed off in the ‘residual body’ (Sheffield and Melton, 1968). In contrast to the plasma membrane, maternal GAP40 did not remain in the residual body (Fig. 1C; Fig. 6A, column vii). The disappearance of maternal IMC from the residual body coincided with an accumulation of GAP40 in the cytoplasm of emerging and maturing daughter cells [Fig. 6A, column vi; also visible in Fig. 1Avi,Bvi; Fig. 2; supplementary material Movie 1 (213–288 min)]. This cytoplasmic accumulation of GAP40 was associated with – but clearly distinct from – the Golgi apparatus (Fig. 2D, 300 min), as previously noted during the early stages of IMC assembly (Fig. 2D, 0–60 min). Colocalization with VP1 (Miranda et al., 2010) and proM2AP (Rabenau et al., 2001) indicated that GAP40 accumulates in the ELC (Robibaro et al., 2001; Tomavo et al., 2013), as shown in Fig. 6C.

To determine whether this late-appearing cytoplasmic pool of GAP40 was actually derived from maternal GAP40, as shown in the cartoon in Fig. 6B, we bleached the maternal IMC in dividing cells and monitored cytoplasmic accumulation of GAP40 after the emergence of daughter parasites (Fig. 7; supplementary material Movie 4). As daughter cells progressed through maturation, a cytoplasmic pool of GAP40 briefly accumulated in the cytoplasm of daughters from control (unbleached) mothers but not in daughters of the bleached mothers (cf. 383 min images in Fig. 7A; supplementary material Movie 4). Moreover, whereas the intensity of daughter parasites was independent of maternal IMC bleaching during elongation (cf. 255 min; Fig. 3), daughter parasites emerging from bleached mothers became progressively dimmer during maturation (cf. 488 min).

Quantification confirmed that the fluorescence of daughter parasites from bleached mothers increased throughout elongation, at an approximately linear rate, comparable to that of daughters from control (unbleached) mothers (aqua lines in Fig. 7B; supplementary material Fig. S3), confirming de novo synthesis of the IMC. However, during maturation (red), the fluorescence
Fig. 4. GAP40 redistribution declines during daughter elongation. GAP40 fluorescence in daughter parasites recovers rapidly at early stages of elongation (closed arrowheads), owing to a combination of de novo synthesis and diffusion or trafficking (A), but fluorescence is unable to fully redistribute when bleaching is applied after ~150 min post-initiation (B, open arrowheads). Bleached areas are outlined in yellow. Scale bars: 5 μm.

Fig. 5. Photoactivation of GAP40–mEos2 indicates protein movement within the elongating IMC. Exposure of transiently expressed GAP40–mEos2 to violet light at various times after initiation (A,B) converts a portion of this reporter to a red fluorescent protein. Red fluorescence remains uniform even during elongation (arrowheads), suggesting diffusion within the IMC. (C) Cartoon illustrating the diffusion of red fluorescence (shows red channel only after photoactivation). Scale bars: 5 μm.
of daughters from unbleached controls continued to rise (solid symbols), whereas that of daughter cells from bleached mothers slowed (open symbols).

Internal fluorescence arose only in daughters emerging from unbleached mothers (gray versus yellow lines in Fig. 7B). The appearance and disappearance of this material coincided precisely with the disappearance of maternal IMC from the residual body and the cessation of GAP40 addition due to de novo synthesis. These observations strongly support the hypothesis that maternal GAP40 is internalized during maturation and recycled into daughter IMCs as illustrated in Fig. 7C.

The replicative cycle of RH strain \( T. gondii \) is typically \(~7\) h, as reported previously (Behnke et al., 2010; Fichera et al., 1995; Nishi et al., 2008), but we observed some variability from specimen to specimen, due to subtle differences in environmental conditions and/or mild phototoxicity during laser photobleaching (parasites that ceased to divide owing to more severe phototoxicity were excluded from analysis). Although atypical, more-slowly replicating parasites provided increased time resolution for the analysis of IMC elongation, maturation and recycling (Fig. 7B, left; supplementary material Movie 4; Fig. S3) and the results obtained from slowly replicating parasites were quantitatively indistinguishable from those of parasites replicating with normal kinetics (Fig. 7B, right; supplementary material Movie 3).

The GAP40–mEos2 fusion protein was also exploited to investigate the apparent recycling of maternal IMC during maturation (Fig. 8A,B). As noted previously (Fig. 5), while elongating daughters became increasingly green due to the addition of newly synthesized material, red fluorescence became increasingly difficult to discern because a fixed amount of photoactivated GAP40 dispersed throughout the growing daughter IMC (cf. Fig. 8A, 135 min). By \(~2\) h post-activation, green fluorescence was evident in daughter parasites, but the red fluorescence of daughter IMCs had declined to near background levels (arrows in 170 min images). Daughter parasites beginning to emerge from the mother cell at 200 min were clearly green but not red (arrows). At this stage, maternal morphology was lost, as the material that was not incorporated into the developing daughters – including the maternal IMC (red) – was sloughed off in the form of the residual body (solid arrowheads). Over the subsequent 1–2 h (cf. 260 min), maternal IMC was gradually lost from the residual body (solid arrowheads), in parallel with transient internalization of GAP40 into the maturing daughter cytoplasm (open arrowheads) and the appearance of (red, maternal) GAP40 in the mature daughter IMC (arrows). Recycled maternal GAP40 became incorporated throughout the mature daughter parasite IMC (390 min).

Further support for the recycling of maternal IMC into daughter parasites came from transient expression studies, as parasites lost the transfected plasmid over time. When the synthesis of fluorescent GAP40 was fortuitously lost during interphase, the fate of the maternal IMC could be followed in the absence of labeled daughter IMC (Fig. 8C). The maternal IMC was visible in the residual body at the time of emergence (180 min), but later appeared in the daughter IMCs (270 min), trafficking through the Golgi and ELC region (210 min).
DISCUSSION
Apicomplexan parasites replicate by a highly unusual process, assembling daughter cells within the mother using a membrane–cytoskeletal structure known as the IMC. In addition to providing a scaffold for daughter cell assembly and the maintenance of parasite shape (Khater et al., 2004; Sheffield and Melton, 1968) (Tremp et al., 2008), the mature IMC also serves to anchor molecular complexes that are associated with parasite motility (the glideosome), host cell attachment and invasion (Gaskins et al., 2004; Keeley and Soldati, 2004). By analogy with the alveolae of ciliates and dinoflagellates, the IMC might also play (as yet undefined) roles in storage and/or homeostasis. Most previous studies on Apicomplexan replication have focused on the IMC cytoskeleton (Anderson-White et al., 2011; Anderson-White et al., 2012; Dearnley et al., 2012; Hu et al., 2002a; Hu et al., 2006), but we know little of how the patchwork of flattened vesicles that comprises the IMC membrane is assembled (Torpier et al., 1991).

In order to directly examine IMC membrane assembly, we used the integral IMC membrane protein GAP40 (Frénal et al., 2010) to define four distinct developmental stages – initiation, elongation, emergence and maturation (Figs 1, 2, 6; supplementary material Figs S1–S4). Careful analysis by time-lapse video microscopy, in combination with photobleaching (Figs 3, 4, 7; supplementary material Fig. S3) and photoactivation (Figs 5, 8), revealed that the IMC of T. gondii tachyzoites is assembled in two distinct phases – elongation involves de novo synthesis (Fig. 3), whereas post-emergence maturation involves salvage and recycling of the maternal IMC (Figs 7, 8). Using modern imaging and analysis techniques, it is possible to quantitatively track the timing of these processes in great detail (Fig. 2B; Fig. 7B; supplementary material Fig. 7. Maternal IMC is recycled into developing daughters after their emergence. (A) The entire IMC of one parasite was photobleached (yellow), whereas that of its sister was not (providing an internal control). Time-lapse imaging reveals comparable fluorescence of the daughter IMCs elongating within bleached versus unbleached mothers but, after emergence (308 min), daughters maturing from the unbleached mother are brighter than those from the bleached mother. During the maturation phase, cytoplasmic fluorescence is evident in daughters from the unbleached control parent (filled arrowhead) but not the bleached parent (open arrowhead). (B) Quantification of IMC fluorescence in daughters from unbleached control parasites (solid symbols) versus bleached parasites (open symbols). Panels are based on independent experiments, corresponding to images in Fig. 7A (left; see also supplementary material Movie 4) and Fig. 3A (right; supplementary material Movie 3). Color coding indicates IMC initiation (blue), elongation (aqua), emergence (magenta) and maturation (red). Note that fluorescence is comparable in daughters emerging from bleached versus unbleached mothers through the elongation stage but diverges after emergence, as daughters maturing from unbleached controls remain bright, whereas those from bleached mothers continue to grow without accumulating additional fluorescence. Transient cytoplasmic fluorescence is observed in daughters maturing from unbleded controls (gray) but not bleached mothers (yellow). Filled and open diamonds indicate the fluorescence of parental parasites before bleaching. (C) Cartoon showing de novo synthesis of the IMC during elongation and salvage of the maternal IMC during maturation. Scale bar: 5 μm.
Figs S2, S3). The resulting timecourse for tachyzoite development (supplementary material Fig. S4) is remarkably consistent with previous studies (Fichera et al., 1995; Nishi et al., 2008; Behnke et al., 2010), but highlights the entire IMC developmental cycle, including daughter parasite initiation, elongation and emergence (~3.5 h), followed by maturation (~2 h), including recycling of the maternal IMC.

Bleaching of the maternal IMC prior to daughter assembly demonstrates that initiation and elongation occur almost entirely through de novo synthesis. Although bleached maternal IMC fails to recover, daughter parasites become brightly fluorescent (Fig. 3A) – comparable to the fluorescence of daughters from unbleached mothers (Fig. 7B). This observation supports previous reports that the maternal IMC in dividing *T. gondii* tachyzoites remains intact from the very earliest stages of daughter assembly to the emergence of daughters from the mother cell (Dubey et al., 1998; Hu et al., 2002a; Sheffield and Melton, 1968). Early ultrastructural studies suggested that the daughter IMC might originate as an outgrowth of the maternal IMC (Vivier and Petitprez, 1969), but see below for further discussion.

It is interesting that GAP40 appears to be added to the entire daughter IMC uniformly throughout the process of assembly (Figs 3, 4, 8), suggesting that the IMC grows in contrast to the processive assembly of subpellicular microtubules (Hu et al., 2002b). FRAP also reveals that GAP40 is able to disperse within the daughter IMC during the first ~2 h of assembly (Fig. 4A; Fig. 5), but becomes increasingly immobile during elongation (Fig. 4B) and is essentially fixed upon emergence and in mature parasites (Fig. 3C). This contrasts with the cytoskeletal protein IMC1, which is continuously remodeled during IMC assembly (Hu et al., 2002a; Mann et al., 2002). Further studies have failed to distinguish between diffusion versus motor- or vesicle-based trafficking, as the motility of intracellular parasites precludes the analysis of small photobleached spots and protein synthesis inhibitors are toxic to intracellular parasites due to indirect effects on the host cell (data not shown). Nevertheless, it is clear that GAP40 mobility declines with time, presumably as a consequence of association with the microtubule and IMC cytoskeleton, lipid rafts and assembly of the glideosome complex (Johnson et al., 2007; Mann et al., 2002).

New parasites ultimately emerge by budding out of the mother, a process that clothes developing daughter IMCs in the maternal plasma membrane and leaves any remaining maternal material behind in the residual body (Fig. 6). In *Plasmodium* parasites, the residual body includes the polymerized heme residue of hemoglobin digestion (hemozoin). Previous studies on *Toxoplasma* have revealed that the mitochondrion is incorporated into developing daughters very late during development, and that rhoptries and micronemes continue to form during and after emergence. We now report that the daughter IMC membrane also continues to expand after emergence, through salvage of the maternal IMC. Photobleaching clearly demonstrates that daughter
IMC growth prior to emergence is independent of the mother, but growth after emergence is almost entirely attributable to salvage of the maternal IMC (Fig. 7). We have termed this newly appreciated phase of the parasite developmental cycle ‘maturaiton’.

Previous ultrastructural studies (Vivier and Petiprez, 1969) suggested that daughter IMCs might derive from that of the mother, but argued for an early maternal contribution. On the basis of live-cell imaging, we demonstrate that the daughter IMC is indeed derived from salvaged maternal components, but that this recycling occurs only late during the parasite replicative cycle. As noted above, all early IMC synthesis occurs de novo, leaving all maternal IMC in the residual body at the point of fusion with the ER.

During maturation, however, maternal IMC disappears from the residual body and becomes incorporated throughout both daughter IMCs (Figs 8A, C, 390 min). The disruption of both de novo IMC assembly (Fig. 2C, left) and IMC salvage (Fig. 2C, right) by BFA treatment, and the association of GAP40 with the ELC (Tomavo et al., 2013) argues that trafficking to the IMC proceeds by the ER–Golgi–ELC pathway. As no vesicles were observed trafficking GAP40 from the residual body to the daughter IMCs, the removal of maternal IMC from the residual body is likely to proceed by fusion with the ER en route to the Golgi and ELC (Fig. 6C, Fig. 2D; supplementary material Movie 5).

It remains uncertain what machinery is involved in IMC formation and recycling, in part due to the lack of adequate molecular markers. Dominant-negative Rab1A and Rab1B proteins disrupt the assembly of the glideosome complex and IMC biogenesis, and perturb vesicular trafficking, organellar segregation and cytokinesis, but the causal relationships between these effects is unclear (Agop-Nersesian et al., 2010; Agop-Nersesian et al., 2009). Overexpression of syntaxin 6 (Stx6, which regulates retrograde transport from the ELC to the Golgi) also disrupts vesicular trafficking to the IMC during the late stages of daughter cell assembly (Jackson et al., 2013), suggesting the possible involvement of this protein in maternal IMC recycling. As additional markers are defined, it will be interesting to further explore IMC biogenesis and recycling, and the relationship of these processes to other membrane trafficking pathways (including dense-granule secretion and endocytosis).

Toxoplasma tachyzoites divide by endodyogeny, making recycling highly efficient. Other life-cycle stages (and many other Apicomplexan parasites) produce multiple daughters by endopolygeny or schizogony, in which nuclei divide prior to the IMC (Ferguson et al., 2008). Whether the IMC is recycled during the emergence and maturation of these parasites is unknown, but the production of multiple daughters certainly diminishes the potential savings from recycling, as the production of 16 daughters (as is typical during the intraerythrocytic replication of P. falciparum, for example) necessitates the production of at least 15 new IMC equivalents. The rapid degradation of the maternal IMC during maturation in Toxoplasma might provide useful insights into the rapid IMC degradation observed as Plasmodium merozoites differentiate into ring-stage parasites within the infected erythrocyte (Bannister et al., 2000; Hepler et al., 1966). Given the importance of the IMC in parasite structural organization and motility, and the fact that Apicomplexan parasite pathogenesis is a direct consequence of rapid proliferation, better understanding of the IMC might also yield new therapeutic strategies. The availability of good markers for both the cytoskeletal and membrane components of the IMC (Fig. 1), and the high degree of time resolution with which the parasite cell cycle can be defined (Fig. 2B, supplementary material Figs S2, S3) provides useful tools for small-molecule screening.

**MATERIALS AND METHODS**

**Cells and parasites**

Human foreskin fibroblasts (HFFs) were cultivated at 37°C under a humidified atmosphere containing 5% CO₂, as previously described (Roos et al., 1995), in a 5:1 mixture of high glucose Dulbecco’s Modified Eagle’s Medium (DMEM, Life Technologies, Grand Island, NY) to Medium 199 (Life Technologies), supplemented with 10% newborn calf serum (NBS, Thermo Scientific, Waltham, MA), 50 μM penicillin, 50 μg/ml streptomycin and 25 μg/ml gentamicin (Life Technologies). Immediately prior to inoculation of the cells with T. gondii tachyzoites, this growth medium was replaced with Minimal Essential Medium (MEM, Life Technologies) supplemented with 2 μM Glutamax (Life Technologies), 1% heat-inactivated fetal bovine serum (FBS, Thermo Scientific) and antibiotics (as above).

**Plasmids**

The allelic replacement plasmid pLicGAP40YFP-dhfrTHXGpRT was engineered by PCR amplification of 1538-bp-spanning TgGAP40 (TgME49_249850; ToxoDB), using the primers shown in supplementary material Table S2, and integration into the Lic sequences in pYFP.Lic.HXG [kindly provided by Vern Carruthers, University of Michigan (Huynh and Carruthers, 2009)]. Plasmid pLicIMC1mCherry-dhfrDHFR was engineered similarly, using a 1950-bp fragment from the 3’ end of TgIMC1 (TgME49_231640) integrated into pmCherry.Lic.DHFR. All plasmids were confirmed by restriction digestion and sequencing. After linearization with Ksal (for GAP40) or BsiWI (for IMC1), 15×10⁶ freshly harvested RHΔku80ΔhxgpRT strain T. gondii tachyzoites (Huynh and Carruthers, 2009) were electrotoporated with 50 μg of plasmid and were selected in 25 μg/ml mycophenolic acid with 50 μg/ml xanthine (GAP40) or 1 μM pyrimethamine (IMC1) (Roos et al., 1995). Clonal plaques were isolated by limiting dilution and screened by fluorescence microscopy for transgene expression. Plasmid p trabACP-YFP-ΔavrIL was engineered by replacing the ACP sequences in p trabACP-YFP-LICΔavr (Nishi et al., 2008) with GAP40 (BglII–AvrII). Plasmid p trabGAP40mEos2-sqCAT was engineered by replacing the YFP-PA in p trabGAP40YFP-HA-sqCAT (Roos et al., 2008) with mEos2 amplified as an AvrII–AvrII fragment from the construct mEos2/AvrII. mEos2 was kindly provided by Michael Davidson, Florida State University (Kauchanawong et al., 2010). Parasites were transfected with 50 μg of plasmid as above and transient transfectants were examined at ~18 h post-transfection. All transgenes (YFP, mCherry, mEos2) utilized standard (non-optimized) coding sequences.

**Immunofluorescence microscopy**

HFF cells were grown to confluence on 22-mm glass coverslips, infected with T. gondii tachyzoites and incubated at 37°C for a further 18–24 h. Coverslips were then fixed for 15–20 min (4% formaldehyde and 0.05% glutaraldehyde in PBS), permeabilized for 15 min (0.25% Triton X-100 in PBS) and blocked for 1 h at room temperature in 3% bovine serum albumin (BSA) fraction V plus 0.25% Triton X-100. After incubation for 1 h with murine monoclonal anti-SAG1 [1:400 in blocking solution; kindly provided by Lloyd Kasper, Dartmouth College (Mineo et al., 1993)] or anti-IMC1 [1:2000, kindly provided by Gary Ward, University of Vermont (Mann and Beckers, 2001; Wichroski et al., 2002)] coverslips were washed three times with 0.25% Triton X-100 in PBS and stained for 1 h in Alexa-Fluor-594-conjugated goat anti-mouse-IgG antibody (1:5000; Life Technologies). For DNA labeling, samples were then incubated for 10 min with 4,6-diamidino-2-phenylindole dihydrochloride (DAPI, EMD Millipore, Billerica, MA) at a final concentration of 0.5 μg/ml in PBS, washed twice with 0.25% Triton X-100 and once with PBS and mounted on glass slides in Fluoromount-G (Southern Biotech, Birmingham, AL).

Imaging was performed on an Olympus IX70 inverted microscope equipped with a UPlanSApo 100× oil-immersion objective (NA1.4).
300 W xenon arc lamp and a CooISNAP HQ monochrome cooled-CCD camera. The excitation and emission filters used for DAPI were 360/40 nm and 455/50 nm, respectively; for GFP were 470/40 nm and 520/40 nm, and for mCherry or RFP were 572/35 nm and 632/60 nm. Image stacks were captured using DeltaVision SoftWorx software (Applied Precision, Issaquah, WA) and deconvoluted (Figs 1, 6 only) using the constrained iterative algorithm, to minimize the effects of out-of-focus fluorescence. Step size was 0.1 or 0.2 μm and acquisition depth was ~2–3 μm, satisfying Nyquist sampling. Images were further analyzed using open-source Fiji software (Schindelin et al., 2012) and were imported into PowerPoint for figure preparation.

**Time-lapse microscopy**

Confluent HFF cell monolayers were cultivated in 35-mm glass-bottomed dishes (Ibidi, Verona, WI), infected with *T. gondii* tachyzoites at a multiplicity of infection (MOI) of ~2:1 in DMEM lacking Phenol Red (Life Technologies), supplemented with 1% FBS, 1 mM sodium pyruvate, 2 mM glutamine, 100 U/ml penicillin, 100 μg/ml streptomycin and 50 μg/ml gentamicin, and incubated for 12–16 h at 37°C. Prior to imaging, cultures were rinsed with warm PBS lacking divalent cations (to remove extracellular parasites) and incubated in fresh Phenol-Red-free DMEM (as above) supplemented with 10% FBS (to minimize laser phototoxicity) and 25 mM HEPES pH 7. Samples were then transferred to a Chamlide TC stage-top environmental chamber (Live Cell Instruments, Guelph, ON, Canada) that was equipped with a digital temperature, CO₂ and humidity control unit and equilibrated ~2 h before data acquisition.

Time-lapse imaging was performed on an Olympus IX-71 spinning-disc confocal microscope equipped with a UPlanSapo 100× oil-immersion objective (NA1.4), CSU-10 scanner (Yokogawa, Newnan, GA), and C9100-13 EMCCD camera (Hamamatsu, Bridgewater, NJ). 5 μm image stacks (26 planes×0.2 μm steps) were acquired by excitation at 488 nm and 561 nm (1% laser power) every 15 or 30 min, using the emission filters ET525/50 for GFP and ET630/75 for mCherry or RFP (Spectral Applied Research). Data was collected using MetaMorph 7.7.4 (Molecular Devices, Downingtown, PA), and processed using MetaMorph and Fiji software. Some images were contrast enhanced for figure presentation.

For quantification of IMC development (Fig. 2), 9–12 time-lapse image stacks (26 planes, as above) were collected over a continuous 9 h session for each of 15 vacuoles in eight fields. At each time-point (for each parasitophorous vacuole) the central image plane was selected from the stack and Fiji software was used to collect perimeter length and total fluorescence (after background subtraction) for all maternal, daughter and/or grand-daughter IMCs, based on manually drawn lines representing all distinctly resolvable IMC structures (supplementary material Table S1a for images and S1b for quantification).

Individual time-lapse series were aligned based on the estimated time of daughter parasite initiation (‘Offset’ column in supplementary material Table S1 and Fig. S2A). Offset values can be reliably estimated to <15 min resolution, as confirmed by the closely coincident timing of daughter parasite emergence at ~195–210 min, grand-daughter IMC initiation at 600–615 min etc. Multiple vacuoles representing all stages of replication were analyzed, from interphase ‘mother’ parasites shortly after invasion through to the formation of daughter parasites (2 per vacuole) and grand-daughters (4 per vacuole). IMC fluorescence was calculated by sliding-window analysis, pooling samples within hourly windows (e.g. initiation t=±30 min), and is presented as the mean ± s.d. (n=6–24; see supplementary material Table S1 for sample sizes).

**Fluorescence recovery after photobleaching**

Photobleaching was performed on the same Olympus IX-71 spinning-disc confocal microscope described above, using a MicroPoint Galvo ablation system (Andor Technology, South Windsor, CT) consisting of a nitrogen-confocal microscope described above, using a MicroPoint Galvo ablation laser (CrystaLaser model DL405-050-O, wavelength of 405 nm).

Owing to the spontaneous photobleaching properties of mEos2, single images were acquired (no stacks) at selected time-points only using MetaMorph 7.7.4 (Molecular Devices). Data was collected at 488 and 561 nm (5–25% laser power; emission filters 525/50 nm for GFP and 617/73 nm for TRITC). Some images were contrast enhanced for figure presentation but all quantitative measurements were performed using unprocessed data.

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**Competing interests**

The authors declare no competing interests.

**Author contributions**

All experiments were conducted by D.T.O.; both authors contributed equally to all aspects of experimental design, data analysis and manuscript preparation.

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**Supplementary material**


**References**


**Figure S1. Staging the IMC cycle in living *Toxoplasma gondii* tachyzoites.** Time-lapse imaging of *T. gondii* parasites coexpressing GAP40-YFP (transient transfectant) and IMC1-mCherry (endogenously-tagged locus). See Fig 1 legend for further description. Scale bars = 5 μm.
**Figure S2. Time-lapse imaging and quantitative dynamics of GAP40.** Panel A, staging of parasite replication in 9-12 time-lapse images (dots) for 15 parasitophorous vacuoles, aligned according to the estimated time of daughter parasite initiation; color coding indicates interphase (gray), IMC initiation (blue), elongation (aqua), emergence (magenta), and maturation (red). Black circles indicate time points where mother and daughter parasites are readily distinguished, permitting accurate quantification (Supplementary Table S1 & Fig 2B). Panel B (same as Fig 2B), quantification of GAP40 fluorescence in maternal, daughter, and grand-daughter parasites (dashed, solid and dotted lines, respectively), determined from 14 sets of time-lapse images (above); sliding window analysis of 1 hr bins, presented as mean ± s.d. (n = 6-24 samples; Supplementary Figure S2 and Table S1). Color coding indicates IMC initiation (blue), elongation (aqua), emergence (magenta), and maturation (red); gray indicates interphase parasites, and purple IMC disappearance.
Figure S3. Bleaching of the maternal IMC affects daughter IMC maturation but not elongation. Accumulation of GAP40 in the IMC during the initiation (blue) and elongation (aqua) of daughter parasites is unaffected by bleaching of the maternal IMC (open symbols), but diverges from controls (closed symbols) during parasite emergence (magenta), indicating that the maternal IMC is recycled into daughter parasites during emergence and maturation (red). Curves represent mean ± s.d. for 12 daughter parasites developing within unbleached (control) mothers (closed symbols), and 8 daughter parasites developing within mothers bleached at various points within the yellow bar (open symbols). Data includes parasites shown in the left-hand panel of Fig 7B. Filled and open diamonds indicate the fluorescence of parental parasites before bleaching.
Figure S4. Summary time course of *T. gondii* tachyzoite development. Following the initiation of daughter parasite assembly within the maternal cytoplasm (*blue*), elongation (*aqua*) involves *de novo* synthesis of the IMC membrane (*GAP40, green*) and cytoskeleton (*IMC1, red*) over the subsequent 3.5 hr. Daughter parasites acquire their plasma membrane as they emerge from the mother (*magenta*), but continue to mature (*red*) over ~2 hr, through the salvage of maternal material (dark shading). Previous studies have focused on organellar segregation during the process of endodyogeny (shaded bar and yellow; *Nishi et al.*, 2008), *i.e.* the 7 hr period between emergence of daughter parasites from the maternal cell, and the emergence of granddaughters. Such analysis fails to fully recognize that complete development of the IMC is an ~11.5 hr process, initiating within the maternal cell, and continuing through dismantling of the daughter cell and salvage of the IMC membrane.
Movie 1. GAP40 Dynamics throughout *T. gondii* replicative cycle. Time-lapse video showing IMC membrane dynamics in GAP40-YFP transgenic parasites (C-terminally tagged at the endogenous genomic locus). Corresponds to Fig 2.

Movie 2. IMC assembly initiates in close proximity to the Golgi apparatus. Time-lapse video of GRASP-mRFP transgenic parasites transiently expressing the IMC marker GAP40-YFP (green). Corresponds to Fig S3.

Movie 3. The daughter IMC is assembled *de novo*. Time lapse video showing the *de novo* assembly of daughter IMCs in GAP40-YFP transgenic parasites after photobleaching of the maternal IMC (top; bottom = unbleached control parasites). Corresponds to Fig 3.
**Movie 4. The maternal IMC is recycled into daughter parasites during maturation.** Time-lapse series of replicating GAP40-YFP transgenic parasites after photobleaching of the maternal IMC (right; left = unbleached control). Video clearly reveals cytoplasmic accumulation of maternal IMC shortly after emergence from unbleached mothers, and recycling into the maturing IMC of daughters. Cytoplasmic accumulation was not observed in daughters emerging from bleached mothers. Corresponds to Fig 7.

**Movie 5. Internalized maternal IMC accumulates transiently in close proximity to the Golgi apparatus during maturation.** Time-lapse video of maturing GRASP-mRFP transgenic parasites transiently expressing the IMC marker GAP40-YFP (green). Movie shows cytoplasmic accumulation of maternal IMC near the Golgi apparatus during maturation.

**Table S1. Quantification of Gap40 fluorescence in maternal, daughter and grand-daughter parasites.** The first page of the Excel spreadsheet displays unenhanced sequential time-lapse images of parasites developing within 15 parasitophorous vacuoles from 8 fields (supplementary material Fig. S2). Vacuoles are marked to indicate parasites used for quantification, as well as these parasites’ history (mothers; daughters a or b; granddaughters a1, a2, b1 or b2), and the stage of replication, using the same blue → aqua → magenta → red color scheme for initiation, elongation, emergence and maturation applied throughout this manuscript. Page 2 presents quantitative fluorescence levels, sequentially aligned on the horizontal axis according to the inferred time since initiation (see supplementary material Fig. S2). Results are summed at the bottom of the table, using a sliding window analysis as described above, and presented in graphical form as Fig. 2B.

**Download Table S1**
<table>
<thead>
<tr>
<th>Primer Set</th>
<th>Forward Primer Sequence (5’-3’)</th>
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* Lic sequences and restriction sites shown in italics; *T. gondii* genomic sequence are underlined.