The *Toxoplasma* Dense Granule Proteins GRA17 and GRA23 Mediate the Movement of Small Molecules between the Host and the Parasitophorous Vacuole

**Graphical Abstract**

**Highlights**
- GRA17 and GRA23 are secreted proteins unique to PV-residing apicomplexans
- GRA17 affects the transfer of small molecules through the PV and in vivo virulence
- The GRA17 loss-of-function phenotypes can be rescued by *Plasmodium* EXP2
- GRA17 or GRA23 expression alters the membrane conductance of *Xenopus* oocytes

**Authors**
Daniel A. Gold, Aaron D. Kaplan, ..., Randall L. Rasmusson, Jeroen P.J. Saeij

**Correspondence**
jsaeij@mit.edu

**In Brief**
*Toxoplasma gondii* resides within an intracellular parasitophorous vacuole (PV) selectively permeable to small molecules. Gold et al. identify GRA17 and GRA23 as secreted *Toxoplasma* proteins that mediate the passage of small molecules between the host cytoplasm and PV, providing a molecular explanation for how these vacuole-residing parasites access host nutrients.
The Toxoplasma Dense Granule Proteins GRA17 and GRA23 Mediate the Movement of Small Molecules between the Host and the Parasitophorous Vacuole

Daniel A. Gold,1 Aaron D. Kaplan,2,3 Agnieszka Lis,2,3 Glenna C.L. Bett,2,3,4 Emily E. Rosowski,1 Kimberly M. Cirelli,1 Alexandre Bougdour,5,6 Saima M. Sidik,7 Josh R. Beck,8 Sebastian Lourido,6 Pascal F. Egea,9 Peter J. Bradley,8 Mohamed-Ali Hakimi,5,6 Randall L. Rasmusson,2,3 and Jeroen P.J. Saeij1,4

1Department of Biology, Massachusetts Institute of Technology, Cambridge, MA 02139, USA
2Physiology and Biophysics, The State University of New York, University at Buffalo, Buffalo, NY 14214, USA
3Center for Cellular and Systems Electrophysiology
4Department of Obstetrics and Gynecology
5School of Medicine & Biomedical Sciences, The State University of New York, University at Buffalo, Buffalo, NY 14214, USA
6Universite´ Joseph Fourier, 38000 Grenoble, France
7Whitehead Institute for Biomedical Research, Cambridge, MA 02142, USA
8Department of Microbiology, Immunology and Molecular Genetics, University of California, Los Angeles, Los Angeles, CA 90095, USA
9Department of Biological Chemistry, UCLA David Geffen School of Medicine, Los Angeles, CA 90095, USA
*Correspondence: jsaeij@mit.edu
http://dx.doi.org/10.1016/j.chom.2015.04.003

SUMMARY

Toxoplasma gondii is a protozoan pathogen in the phylum Apicomplexa that resides within an intracellular parasitophorous vacuole (PV) that is selectively permeable to small molecules through unidentified mechanisms. We have identified GRA17 as a Toxoplasma-secreted protein that localizes to the parasitophorous vacuole membrane (PVM) and mediates passive transport of small molecules across the PVM. GRA17 is related to the putative Plasmodium translocon protein EXP2 and conserved across PV-residing Apicomplexa. The PVs of GRA17-deficient parasites have aberrant morphology, reduced permeability to small molecules, and structural instability. GRA17-deficient parasites proliferate slowly and are avirulent in mice. These GRA17-deficient phenotypes are rescued by complementation with Plasmodium EXP2. GRA17 functions synergistically with a related protein, GRA23. Exogenous expression of GRA17 or GRA23 alters the membrane conductance properties of Xenopus oocytes in a manner consistent with a large non-selective pore. Thus, GRA17 and GRA23 provide a molecular basis for PVM permeability and nutrient access.

INTRODUCTION

Protozoan parasites in the phylum Apicomplexa present a global human and veterinary health problem. Toxoplasma gondii, which causes toxoplasmosis, can infect a wide range of warm-blooded animals including humans and is a particular challenge to immune-compromised individuals such as AIDS patients and fetuses of infected mothers. Apicomplexan parasites spend the majority of their life cycle within host cells and have evolved diverse strategies to survive in this niche. Many, like Toxoplasma and Plasmodium, reside within a non-fusogenic parasitophorous vacuole (PV). The PV membrane (PVM) is the physical interface between PV-residing parasites and their host cells during intracellular growth, remaining intact until parasite egress. The PV protects parasites from being sensed by the host cell, but it is also a potential barrier to the scavenging of nutrients from the host and to parasite effectors accessing the host.

Apicomplexan parasites depend on host cells for many essential nutrients. For instance, Toxoplasma is auxotrophic for small molecules including amino acids like tryptophan and arginine, purines, and sterols (reviewed in Coppens, 2014). Some, like sterols, are actively scavenged from the host cell to the PV through sterol transporters on the PV (Ehrenman et al., 2010). Transporters for small metabolites have been identified on the Toxoplasma plasma membrane but not on the PVM (Landfear, 2011). The Toxoplasma, Plasmodium, and Eimeria PVMs are selectively permeable, allowing passive, charge-independent diffusion of small molecules up to 1,300–1,900 Da into the PV, presumably through pore-like molecules on the PVM (Desai and Rosenberg, 1997; Schwab et al., 1994; Werner-Meier and Entzeroth, 1997). This is the putative mechanism by which the parasite within the PV gains access to host metabolites or effluxes toxic metabolic byproducts like lactate into the host cell. The molecular identity of the pores that grant this PVM permeability is unknown.

Toxoplasma partially circumvents the PVM barrier by secreting proteins into the host cell before and during invasion before formation of the PV from secretory organelles that include rhoptries and dense granules. Secreted effectors play a critically important role in Toxoplasma pathogenesis, modulating host signaling and creating a hospitable environment in the PV (reviewed in Hunter and Sibley, 2012; Kemp et al., 2013). Unlike
GRA17 are unique to PV-residing Apicomplexa

<table>
<thead>
<tr>
<th>Organism</th>
<th>GRA17-Like Genes</th>
<th>Gene Names</th>
<th>PV?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Toxoplasma gondii</td>
<td>2</td>
<td>GRA17 (TGGT1_222170), GRA23 (TGGT1_297880)</td>
<td>yes</td>
</tr>
<tr>
<td>Neospora caninum</td>
<td>2</td>
<td>NCLIV_005560, NCLIV_006780</td>
<td>yes</td>
</tr>
<tr>
<td>Hammondia hammondi</td>
<td>2</td>
<td>HHA_222170, HHA_297880</td>
<td>yes</td>
</tr>
<tr>
<td>Sarcocystis spp.</td>
<td>2</td>
<td>S. neurona A, S. neurona B; S. muris DG32</td>
<td>yes</td>
</tr>
<tr>
<td>Eimeria spp.</td>
<td>1</td>
<td>E. acervulina EAH_00026470 (DG32) and E. mitis EMH_0012530 (DG32)</td>
<td>yes</td>
</tr>
<tr>
<td>Plasmodium spp.</td>
<td>1</td>
<td>P. falciparum PFD17_1471100 (EXP2), P. knowlesi PKH_123420 (EXP2), P. reichenowi PRCDC_1470300 (EXP2)</td>
<td>yes</td>
</tr>
</tbody>
</table>

Babesia bovis 0 NA no
Cryptosporidium spp. 0 NA no
Theileria spp. 0 NA no
Gregarina niphandroides 0 NA no

The peptide sequences of GRA17 and GRA23 were used as bait in searches against apicomplexan organisms in EuPathDB.org using the BLASTP 2.0 algorithm or TBLASTN 2.2.26. Subsequent hits receiving a p value less than 0.005 were identified as similar to GRA17 and themselves used as baits in subsequent searches.

"The S. neurona genome is sequenced but unannotated, so the two GRA17-like genes were arbitrarily named “A” and “B.”

"The S. muris genome has not been sequenced, so we cannot determine whether there is a second GRA17-like gene.

"GRA17-like genes were identified for all Eimeria and Plasmodium species whose genomes have been sequenced, but for brevity, only a selection of these were included in this table.

the invasion-secreted rhoptry proteins (ROPs), dense granule proteins (GRAs) are also secreted post-invasion (Karsten et al., 1998).

Plasmodium exports proteins beyond its PVM into the erythrocyte cytosol. EXP2 is the putative protein-conducting pore of the Plasmodium translocon for exported proteins (PTEX) complex that mediates the essential export of secreted proteins containing a Plasmodium export element (PEXEL) sequence motif (de Koning-Ward et al., 2009). There are also PEXEL-independent mechanisms of protein export (Grüning et al., 2012; Slau et al., 2014) that utilize the same PTEX machinery (Beck et al., 2014; Elsworth et al., 2014). In Toxoplasma, post-invasion protein export into the host cell has only been reported for GRA16 and GRA24 (Bougdour et al., 2013; Braun et al., 2013). Most characterized Toxoplasma GRA proteins are involved in organizing the structure and environment of the PV or associated with the PVM, but others such as GRA15, GRA16, and GRA24 play a role in modulating host cell signaling, suggesting that there may be more GRA proteins trafficked beyond the PVM after invasion (Bougdour et al., 2013; Mercier et al., 2005; Rosowski et al., 2011).

We sought to investigate the molecular interaction between the Toxoplasma PV and the host cell by studying a PVM-localized GRA protein with homology to Plasmodium EXP2.

RESULTS

GRA17 is a Secreted GRA Protein Required for Normal PV Morphology

Since EXP2 has been implicated in forming a molecular-translocation channel in the P. falciparum PV, we searched for homologs in the Toxoplasma genome and found two genes encoding proteins with predicted signal peptides: the uncharacterized TGGT1_222170 and TgGRA23 (TGGT1_297880), an identified GRA gene of unknown function (Masatani et al., 2013). TGGT1_222170 and TgGRA23 share 30% amino acid sequence similarity with each other and 26% and 22% similarity with PfEXP2, respectively (Figure S1A). Homologs of these proteins, including Sarcocystis muris DG32 (Freyer et al., 1999), were identified only in Apicomplexa that replicate within a PV (Table 1). Except for Plasmodium spp. EXP2, none of these genes have been functionally characterized, but due to its similarity to GRA23, we provisionally named TGGT1_222170, “GRA17.” There is only one gene similar to TgGRA17 or TgGRA23 found in Eimeria spp. and Plasmodium spp., but we found two distinct homologs in the genomes of other sequenced PV-residing Apicomplexa that are grouped in the family Sarcocystidae (Table 1; Figure S1B). We found that GRA17 and GRA23 are conserved proteins with high gene transcription levels across a global range of Toxoplasma strains (Meio et al., 2013; Minot et al., 2012).

Besides strongly predicted α helices, GRA17 and GRA23 lack sequence motifs that suggest a functional role. To characterize them, we individually disrupted the GRA17 and GRA23 genes and we complemented the ΔGRA17 strain with either HA-tagged GRA17 or Toxoplasma-codon-optimized P. falciparum EXP2 transgenes that we will refer to as “ΔGRA17+GRA17-HA” and “ΔGRA17+PfEXP2-HA” (Figures S2A–S2E). The ΔGRA17 strain was also complemented with GRA23-HA-FLAG (HF) expressed from either the putative gra23 promoter “ΔGRA17+GRA23-HFlow” or from the strong gra1 promoter “ΔGRA17+GRA23-HFhigh.” The differing expression levels in these strains were verified by western blot (Figure S2F). We infected human foreskin fibroblasts (HFFs) with tachyzoites (the fast-replicating stage of Toxoplasma), fixed, and analyzed the sample by deconvolution immunofluorescence (IF) microscopy. The ΔGRA17+GRA17-HA strain was used for many co-staining experiments as it does not express GFP, and the GRA17-HA localization was invariant across all the strains (data not shown). We observed that GRA17-HA is intra-vacuolar and is associated with the PVM and possibly the tubulo-vesicular network (TVN) based on its partial co-localization with GRA7 (a PVM and TVN marker) and
GRA4 (a TVN marker) especially around the periphery of the PV (Figure 1A). GRA23 has previously been characterized as a PVM-localized GRA (Masatani et al., 2013), and to observe its relative co-localization with GRA17, we generated a strain that co-expresses GRA17-V5 and GRA23-HF (the tachyzoites express GFP) and observed substantial co-localization around the PVM (Figures 1A, bottom panel, and S2G); however, the localization of GRA17-HA appeared localized to discrete foci relative to the more continuous pattern of GRA23-HF around the PVM.

\[\text{Pf}^{\text{EXP2-HA}} \text{localized to the PVM in the} \text{D}^{\text{GRA17}} \text{strain, and the localization of GRA23-HF was unaffected by the absence of GRA17 in the} \text{D}^{\text{GRA17+GRA23-HF}} \text{low strain (Figure S2H). We confirmed that GRA17 is a GRA by observing partial co-localization of GRA17 and GRA7 in extracellular tachyzoites (Figure S2I).}

We raised a specific antibody against GRA17 that detected a 19 kDa band that is absent in the \text{D}^{\text{GRA17}} strain (Figure S3A, lanes 1–4).

We tested GRA17 association with the PVM by fractionating HFFs infected with WT+GRA17-HA by ultracentrifugation and extracting a membrane-enriched fraction with either PBS or sodium carbonate. Sodium carbonate treatment extracts peripherally attached proteins such as GRA1, but more tightly membrane-associated proteins like GRA7 remain in the insoluble fraction (Neudeck et al., 2002; Sibley et al., 1995). We found that in contrast to GRA1, GRA17 was only minimally extracted by sodium carbonate and that the majority remained in the insoluble fraction (Figure 1B). Thus, the majority of GRA17 is tightly associated with the PVM. A similar fractionation pattern for GRA23 has been reported (Masatani et al., 2013).

We observed that a significant portion of PVs in the \text{D}^{\text{GRA17}} strain were grossly enlarged relative to the number of parasites in the PV, whereas the \text{D}^{\text{GRA23}} PVs are similar to wild-type (WT) (Figure 1C). Furthermore, the \text{D}^{\text{GRA17+GRA17-HA}}-complemented strains, \text{D}^{\text{GRA17+GRA17-HA}} and \text{D}^{\text{GRA17+PfEXP2-HA}}, completely rescued this aberrant morphology. We were unable to generate a \text{D}^{\text{GRA17+D}^{\text{GRA23}}/\text{D}^{\text{GRA23}}} strain, but we were able to delete the GRA23 gene from the \text{D}^{\text{GRA17+GRA17-HA}} strain, which has low GRA17 expression (Figure S3A, lanes 2 and 8). We observed that this \text{D}^{\text{GRA17+D}^{\text{GRA23}}+\text{GRA17-HA}} strain mimics the aberrant PV morphology of the \text{D}^{\text{GRA17}} strain (Figure 1C), providing evidence of functional overlap between GRA17 and GRA23.

Furthermore, the \text{D}^{\text{GRA17}} strain complemented with high (but not lower) expression of GRA23-HF partially suppressed this phenotype, with infrequent enlarged vacuoles observed (Figure 1C).

We examined whether there were other genes involved in a compensatory role with the \text{D}^{\text{GRA17}} strain by sequencing the total mRNA from bone-marrow-derived mouse macrophages 4 hr post-infection with WT, \text{D}^{\text{GRA17}}, \text{D}^{\text{GRA23}}, and \text{D}^{\text{GRA17+GRA17-HA}} strains and mapping the sequences to the Toxoplasma genome. As previously observed, GRA17 expression in the \text{D}^{\text{GRA17+GRA17-HA}} strain was 2.5-fold less compared to WT despite its ability to functionally rescue the \text{D}^{\text{GRA17}} strain (Figure 3B) (GEO: GSE65980). GRA23 expression levels in the \text{D}^{\text{GRA17+GRA17-HA}} strain were 1.8-fold increased over WT levels, and GRA17 levels were not increased in the \text{D}^{\text{GRA17}} strain (Figure S3B). We found five additional, uncharacterized putative signal peptide-containing proteins that were uniquely upregulated in the \text{D}^{\text{GRA17}} strain (Table S2).
GRA17 Is Involved in the Movement of Small Molecules between the Host and PV

Since PfEXP2 can rescue a ΔGRA17 phenotype, we wondered whether GRA17 or GRA23 plays a role in protein export beyond the PVM. The GRA proteins GRA16 and GRA24 localize to the host nucleus with increasing intensity over the time of infection, peaking at 20 hr, indicating that they are exported across the PVM after the PV has been formed (Bougdour et al., 2013; Braun et al., 2013). We tagged the endogenous copy of GRA24 with HA-FLAG (HF) in the WT, ΔGRA17, and ΔGRA23 strains. We also transiently transformed GRA16-HF into WT, ΔGRA17, and ΔGRA23 extracellular tachyzoites with which we immediately infected HFFs. We then fixed the infected cells at early and late time points post-infection. Across all strains we observed a temporal increase from low to high levels of GRA16-HF and GRA24-HF staining in the host nucleus (Figures 2A and S3C). Thus, neither GRA17 nor GRA23 are individually required for GRA16 and GRA24 export beyond the PVM.

While GRA17 and GRA23 do not appear to affect protein transport across the PVM, we asked if they function as a pore for small molecules and metabolites.

We used the properties of the vital dye 5-(and-6)-carboxy-2,7'-dichlorofluorescein diacetate (CDCFDA) to test this hypothesis. CDCFDA is membrane permeable and non-fluorescent until it enters living cells, where intracellular esterases convert it into the fluorescent, membrane-impermeable 5-(and-6)-carboxy-2,7'-dichlorofluorescein (CDCF). The molecular weight of the CDCF fluorophore (445.2 Da) is less than the established size-exclusion limit of the Toxoplasma PVM (Schwab et al., 1994), so we predicted that it should passively enter into the PV through the PVM but be unable to permeate through the tachyzoite plasma membrane (Schwab et al., 1994). We infected HFFs with the WT, ΔGRA17, ΔGRA17+GRA17-HA, ΔGRA23, ΔGRA17+PfEXP2-HA, ΔGRA17+GRA23+GRA17-HA, and ΔGRA17+GRA23-HF_low-high strains for 24 hr and then added a pulse of CDCFDA to the cell culture medium before washing away the dye and subsequently imaging the living cells. As predicted, we observed CDCF-fluorescence intensity equivalent to the host cytosol within nearly all the PVs of the GRA17 nor GRA23 are individually required for GRA16 and GRA24 export beyond the PVM.

While GRA17 and GRA23 do not appear to affect protein transport across the PVM, we asked if they function as a pore for small molecules and metabolites.

We used the properties of the vital dye 5-(and-6)-carboxy-2,7'-dichlorofluorescein diacetate (CDCFDA) to test this hypothesis. CDCFDA is membrane permeable and non-fluorescent until it enters living cells, where intracellular esterases convert it into the fluorescent, membrane-impermeable 5-(and-6)-carboxy-2,7'-dichlorofluorescein (CDCF). The molecular weight of the CDCF fluorophore (445.2 Da) is less than the established size-exclusion limit of the Toxoplasma PVM (Schwab et al., 1994), so we predicted that it should passively enter into the PV through the PVM but be unable to permeate through the tachyzoite plasma membrane (Schwab et al., 1994). We infected HFFs with the WT, ΔGRA17, ΔGRA17+GRA17-HA, ΔGRA23, ΔGRA17+PfEXP2-HA, ΔGRA17+GRA23+GRA17-HA, and ΔGRA17+GRA23-HF_low-high strains for 24 hr and then added a pulse of CDCFDA to the cell culture medium before washing away the dye and subsequently imaging the living cells. As predicted, we observed CDCF-fluorescence intensity equivalent to the host cytosol within nearly all the PVs of the
WT, ΔGRA17+GRA17-HA, ΔGRA17+PfEXP2-HA, and ΔGRA23 strains but excluded from the tachyzoites themselves (Figure S4A). By contrast, 36% of the PVs in the ΔGRA17 strain had equivalent intra-PV CDCF-fluorescence compared to the host cytosol (p = 4 × 10^{-8} to WT, Figure 2B, red arrow and right panel), suggesting a role for GRA17 involving the passage of small molecules into the PV. We observed a similar phenotype for the ΔGRA17/ΔGRA23+GRA17-HA strain, with CDCF-fluorescence in 51% of PVs (p = 0.002 to WT and no statistical difference to ΔGRA17; Figure 2B, right panel). In contrast to the ΔGRA17+GRA23-HF_{low} strain, which was similar to the ΔGRA17 strain (45% intra-PV CDCF-fluorescence; p = 0.002 from WT), we observed a partial rescue of dye transfer with the ΔGRA17+GRA23-HF_{high} strain with CDCF-fluorescence in 73% of PVs that was statistically distinct from the ΔGRA17 strain (p = 0.01; Figure 2B), further suggesting a synergistic role for GRA23 in small molecule permeability. Since we observed PVs of varying permeability within the same host cell (Figure 2B, ΔGRA17), the GRA17-mediated effect on PVM permeability to the small molecule fluorophore is PV autonomous and independent of host intracellular fluorophore concentration.

We confirmed this phenotype by introducing the membrane-impermeable dyes lucifer yellow (522 Da) or Dextran (3,000 Da)-Alexa-488 into infected cells by the technique of scrape-loading (McNeil et al., 1984). Since cellular dye uptake varies around the scrape, there was a large degree of heterogeneity of both intracellular and intra-PV dye intensities. We scored the intra-PV dye intensity relative to its host cell dye intensity as “present,” “absent,” or “intermediate” (Figure S3D). We found that the scrape-loaded lucifer yellow results were similar to the CDCFDA results: in contrast to >90% of WT, ΔGRA17+GRA17-HA, and ΔGRA23 PVs, only 22% of ΔGRA17 PVs were scored as having fluorescence fully equivalent to their host cells (Figure 2C). When the larger dextran (3,000 Da) dye was scrape-loaded into infected cells, we found that most of the PVs of both WT and ΔGRA17 failed to uptake the dye, consistent with the established Toxoplasma PVM size-exclusion limit (Schwab et al., 1994). These results provide further evidence that PfEXP2 can rescue ΔGRA17 phenotypes and for functional overlap between GRA17 and GRA23.

The Absence of GRA17 Results in Highly Dynamic, Unstable PV

We found that fixatives altered the appearance and numbers of the enlarged, aberrant ΔGRA17 PVs (data not shown), so we imaged the ΔGRA17 strain live over an extended time period post-infection to observe the dynamics of these PVs. Nearly all of the PVs we observed by live imaging appeared enlarged at some point during the experimental time course. We also observed many enlarged PVs “collapsing,” where the tachyzoites within them turn opaque and cease to divide. As an example, we present a multiple infected host cell where all but one PV collapses over a 5:30 hr time course (Figure 3A; Movie S1). The remaining intact PV within this cell remained enlarged and continued to divide, ultimately egressing from the cell at 62:00 hr (Movie S2). We have also observed PV collapse in singly infected cells (Figure S4A). Furthermore, we observed dynamic alteration of the size and shape of the vacuoles as exemplified by a PV that becomes grossly enlarged to the size of the entire host cell before ultimately egressing (Figure S4B; Movies S3 and S4). Since the tachyzoites within these enlarged PVs divide normally and we regularly observe egress, it indicates that they are viable.

In concordance with the PV collapse, we observed that ΔGRA17 tachyzoites grew more slowly than WT tachyzoites. We measured this effect by quantifying the mean area of plaques formed in HFF monolayers by infection with the ΔGRA17 and ΔGRA23 strains relative to WT. These assays measure the overall growth of the parasites in vitro. We found that ΔGRA17 formed plaques that were 53% the size of the WT strain (p < 0.0001; Figures 3B and S5A), and the ΔGRA17+GRA17-HA strain fully rescued this growth defect. The ΔGRA23 strain formed plaques that were slightly smaller than the WT strain, but this effect was not significant (p = 0.21; Figures 3B and S5A). In contrast to the smaller plaques made by the ΔGRA17 strain, we observed that a strain that overexpresses GRA17-HA (WT+GRA17-HA; Figure S3A, lanes 9 and 10) formed plaques that were 67% larger than WT (p < 0.0001; Figures 3B and S5A), implying that excess GRA17 can accelerate growth of the parasite. The collapsed PVs with non-dividing tachyzoites suggest a basis for the slow growth of the ΔGRA17 strain in vitro.

GRA17 and GRA23 Affect the In Vivo Growth and Virulence of Toxoplasma

We examined whether the effects of GRA17 on overall Toxoplasma proliferation in vitro would manifest in vivo. We infected CD-1 outbred mice intraperitoneally (i.p.) with 100 tachyzoites of the WT, ΔGRA17, ΔGRA17+GRA17-HA, and ΔGRA23 strains. We also infected an additional cohort with 1,000 tachyzoites of the ΔGRA17 strain. None of the mice infected with the WT, ΔGRA23, and ΔGRA17+GRA17-HA strains survived beyond 12 days post-infection (Figure 4B, right panel). In contrast to the smaller plaques made by the ΔGRA17 strain, we observed that a strain that overexpresses GRA17-HA (WT+GRA17-HA; Figure S3A, lanes 9 and 10) formed plaques that were 67% larger than WT (p < 0.0001; Figures 3B and S5A), implying that excess GRA17 can accelerate growth of the parasite. The collapsed PVs with non-dividing tachyzoites suggest a basis for the slow growth of the ΔGRA17 strain in vitro.

We infected CD-1 outbred mice intraperitoneally (i.p.) with 100 tachyzoites of the WT, ΔGRA17, ΔGRA17+GRA17-HA, and ΔGRA23 strains. We also infected an additional cohort with 1,000 tachyzoites of the ΔGRA17 strain. None of the mice infected with the WT, ΔGRA23, and ΔGRA17+GRA17-HA strains survived beyond 12 days post-infection (Figure 4A), consistent with the reported hyper-virulence of the type I genetic background of our strains (LD_{100} = ~1 tachyzoite). By contrast, all of the mice infected with either the low or high dose of the ΔGRA17 tachyzoites survived. All of these mice were seropositive and were resistant to subsequent challenge with 10,000 WT (type I) tachyzoites (Figure 4A, arrow). Subsequently, we infected cohorts of CD-1 mice with 100 tachyzoites each of the ΔGRA17, ΔGRA17+GRA17-HA, ΔGRA17+PfEXP2-HA, and ΔGRA17+GRA23+GRA17-HA strains and monitored both body weight and survival. We found that none of the mice infected with ΔGRA17+GRA17-HA or ΔGRA17+PfEXP2-HA survived past 12 days post-infection, whereas all of the mice infected with the ΔGRA17 or ΔGRA17+GRA23+GRA17-HA strains survived the duration of the experiment (Figure 4B, top). In contrast to the mice infected with ΔGRA17 parasites, those infected with ΔGRA17+GRA23+GRA17-HA experienced a significant drop in body weight and deterioration in body condition (e.g., hunched stance and ruffled fur) during the course of infection before recovering fully by 30 days post-infection (Figure 4B, bottom). Thus, PfEXP2 can rescue the loss of GRA17 in vivo, and the deletion of GRA23 with low GRA17 expression renders Toxoplasma avirulent. Since the type I strain is hyper-virulent in mice, we tested the accelerated growth in vivo by overexpressing GRA17 in the far less virulent type III strain (LD_{100} = 10^{5} tachyzoites), which also
accelerated its in vitro growth (Figure S5B). We infected BALB/c mice with this strain and a WT (type III) strain. We found that all the infected mice survived, but the mice infected with type III+GRA17-HA had a higher parasite burden (Figure S5C), lost more weight (Figure S5D), had a worse body condition (e.g., hunched posture and ruffled coat), and had a higher brain tissue cyst count in some but not all of this cohort (Figure S5E).

GRA17 and GRA23 Are Synergistically Required for Growth

We were able to detect ΔGRA23 phenotypes only in the background of significantly reduced GRA17 expression, and the overexpression of GRA23-HF in the ΔGRA17 background was able to partially rescue ΔGRA17 phenotypes, suggesting GRA17 and GRA23 operate synergistically. While we were able to generate multiple, independent clonal tachyzoite isolates of ΔGRA17/ΔGRA23+GRA17-HA, we could not generate a ΔGRA17/ΔGRA23 strain in parallel by multiple methods. In order to determine whether a ΔGRA17/ΔGRA23 strain is viable, we used the CRISPR gene disruption technology developed for Toxoplasma (Shen et al., 2014; Sidik et al., 2014). We chose target strains where GRA17-HA is the only species of GRA17 present so we could follow its fate by IF microscopy. As a test, we co-transformed the ΔGRA17/ΔGRA23+GRA17-HA strain with Cas9-3xFLAG, a guide RNA (gRNA) to target protospacer sequences early within the GRA17 coding sequence as well as a double-stranded oligonucleotide donor template containing a nonsense mutation near the predicted Cas9-cleavage site. We co-transformed donor templates with our gRNAs, since our target strain background is ΔKu80, which eliminates non-homologous end-joining in favor of homologous recombination (Fox et al., 2009). We used three different gRNA/donor template pairings, fixed at 24 hr and 48 hr, and observed these gRNAs led to similar GRA17-HA loss relative to no loss with a non-specific gRNA (Figures 5A, top panel, and S6).

After transformation of both the ΔGRA17/ΔGRA23+GRA17-HA and ΔGRA17+GRA17-HA strains with Cas9-3xFLAG, a GRA17-specific gRNA/donor [residues 29–36]/donor template [Leu32stop] and subsequent infection of HFFs, we fixed infected cells early to compare the initial loss of GRA17-HA signal in the population (24 hr) to late, when the parasites will form plaques in the host cell monolayer (120 hr). Both target strains were transformed at an equivalent rate as measured by Cas9-3xFLAG detection (~27%–28%; Figure 5B, top panel). We also found that ΔGRA17+GRA17-HA and
GRA17 and GRA23 Alter the Membrane Conductance of Xenopus Oocytes

We expressed GRA17 and GRA23 in Xenopus oocytes to see if they can alter membrane permeability consistent with the insertion of large, relatively non-selective pores into the cell membranes. The expression of such a non-selective pore makes several experimentally testable predictions. The first is that expression will be detrimental to cells, resulting in significant cell death due to loss of cell contents and inability to maintain volume homeostasis. When varying levels of GRA17 and GRA23 cRNAs were injected into Xenopus oocytes, a dose-dependent increase in cell lysis was observed relative to water-injected cells or to GRA15 cRNA, a GRA protein that localizes to the PVM (Rosowski et al., 2011) which is not predicted to affect membrane permeability (Figure 6A). HA-tagged versions of GRA17, GRA23, and GRA15 were detected in total membrane fractions isolated from cRNA-injected oocytes (Figures S7A and S7B), and GRA17 and GRA23 were also observed localized to the oocyte plasma membrane in fixed histology sections (Figure S7C). The second prediction is that increased membrane permeability through expression of putative non-selective pores would cause a depolarizing shift in membrane resting potential prior to cell lysis, and GRA17 and GRA23 both demonstrated this shift relative to water- or GRA15-injected oocytes (Figure 6B). Finally, the current to voltage relationship of the net membrane current for a large non-selective, constitutively open pore should result in an increased linear current-voltage relationship (Figure 6C), a depolarization of the reversal potential (Vrev; Figure 6D, top panel), and an increased conductance (G; Figure 6D, bottom panel) for GRA17 and GRA23 relative to water- or GRA15-injected oocytes, which we observed. The results of these exogenous expression studies are strongly consistent with the hypothesis that GRA17 and GRA23 can form non-selective high-conductance pores when exogenously expressed in cell membranes. We injected a separate batch of Xenopus oocytes with PfEXP2-HA cRNA and, as with the Toxoplasma GRAs, found that it was present in total membrane and cytosolic fractions (Figure S7D). Relative to GRA17 and GRA23, PfEXP2 cRNA-injected oocytes had a more modest decrease in survival (73% compared to 93% of water-injected oocytes; Figure S7E). There was no statistically significant difference between PfEXP2 cRNA-injected oocytes and water-injected control resisting membrane potential (RMP) (Figure S7F); however, the control oocytes had a significantly depolarized RMP compared to the previous batches of oocytes used for the Toxoplasma GRAs (Figure 6B), indicating that PfEXP2 forming a large membrane pore. Further work is needed to characterize the role of PfEXP2 as a membrane pore.

DISCUSSION

Intracellular pathogens that reside within vacuoles face a challenge of scavenging nutrients and modulating the host cell beyond their vacuole membrane. The existence of putative pores providing selective permeability to small molecules in the Toxoplasma and Plasmodium PVMs has been described (Desai and Rosenberg, 1997; Schwab et al., 1994), and now we have provided a molecular basis for this phenomenon in any Apicomplexa. The flow of small molecules between the host cytosol and PV plays a critical role in tachyzoite growth and survival, and its reduction may result in either restriction to nutrients or the inability of toxic metabolic byproducts to efflux.
out of the PV. Overexpressing GRA17 leads to more rapid growth, suggesting that nutrients scavenged by Toxoplasma via GRA17 may be limiting factors for normal growth of the parasite.

An important question is how the ∆GRA17 strain remains viable in vitro when it is starved from access to small molecules. A hint as to why some parasites retain viability lies in the aberrant morphology of the ∆GRA17 PVs. The tachyzoites within the enlarged vacuoles are viable as they replicate and egress. The aberrant PVs may then be the byproduct or consequence of a compensatory mechanism for GRA17 absence. Compensation may include a change in the osmolarity or other structural forces, resulting in PV enlargement. The collapsed PVs may be due to ineffective compensation to reduced access to small molecules or to overcompensation that leads to PV damage. PV collapse, as well as the sensitivity of the enlarged PVs to fixatives, hints at the dynamism and fragility of the ∆GRA17 PVs and implies a precarious balance of compensatory forces.

A prime candidate for a compensatory molecule is GRA23 based on the body of evidence we have presented. One model is that GRA17 and GRA23 may form a variety of homo- or hetero-multimers whose biophysical properties may differ depending on the composition of the complex. Alternatively, GRA17 and GRA23 may function on the PVM separately, as monomers or otherwise, having partially overlapping but ultimately separable functions. GRA17 and GRA23 may function similarly to α-helical pore-forming molecules like colicins or the pro-apoptotic protein Bax that can exist in either a water-soluble state or as an integral membrane pore that alters permeability (reviewed in Parker and Feil, 2005). Particularly intriguing are Bax-derived small α-helical peptides that can homo-multimerize within the membrane via a toroidal mechanism that does not require direct protein-protein interactions but instead contains proteins interspersed with lipids (García-Sáez et al., 2006; Qian et al., 2008). We have immunoprecipitated both GRA17 and GRA23 from PVM-containing fractions in the cell (data not shown), but we have so far not been able to identify specific interacting proteins. Our mRNA sequencing experiment did identify several other genes encoding putatively secreted protein products that are uniquely upregulated in the ∆GRA17 strain, but none of these have been functionally characterized to date. Thus, future work is required to define the exact molecular functions of GRA17 and GRA23 as well as identify associated molecular factors.

The observation that Plasmodium EXP2 can rescue ∆GRA17 phenotypes hints at a possible role of GRA17 or GRA23 in protein export in Toxoplasma. However, GRA16 and GRA24 accumulation in the host nucleus via post-invasion in both the ∆GRA17 and ∆GRA23 strains argues against such a role, although GRA17 and GRA23 may have redundant functions in protein export. It will be difficult to distinguish between a potential defect in export in the absence of both proteins versus
parasites that are dead or dying and thus unable to actively export protein. Ultimately, very little is known about Toxoplasma protein export, and there are currently only two reported exported proteins to test as candidates. Even though Plasmodium EXP2 has not been formally shown to be the translocation pore for protein export, it begs the question of whether it serves a dual role of nutrient pore and protein channel, since EXP2 is the only GRA17-like protein in Plasmodium spp. The aforementioned a-helical pore-forming proteins can form pores big enough for peptides in many cases (Parker and Feil, 2005), suggesting this dual functionality is feasible. It is possible that the other proteins in the PTEX complex may direct the function of EXP2 from a potential nutrient pore toward a role in protein export.

GRA17 and GRA23 may help elucidate another longstanding question of Toxoplasma biology concerning the sporozoite, the sexually derived stage found in the oocyst. After excystation from the oocyst and invasion of the host cell, the sporozoite forms a unique vacuole (PV1) in which the sporozoite undergoes no replication and ultimately egresses from the PV1 to form a second vacuole (PV2) in the same cell where it completes its conversion to the tachyzoite stage and undergoes replication. Unlike tachyzoite PVs and PV2, the PV1 is impermeable to small molecules and devoid of a subset of GRAs typically associated with PV ultrastructure, suggesting that their absence plays a role in the unique properties of the sporozoite PV1.

We believe GRA17 and GRA23 present exciting tools to understand the challenge of how pathogens proliferate within an intracellular vacuole, which is a phenomenon widely conserved in pathogens across evolution.

EXPERIMENTAL PROCEDURES

Reagents
5-(and-6)-carboxy-2',7'-dichlorofluorescein diacetate (C-369) and Dextran 3000-Alexa488 (D34682) were obtained from Life Technologies and lucifer yellow CH dipotassium salt (L0144) from Sigma-Aldrich.

Parasite and Cell Culture and Plaque Size Assays
Toxoplasma tachyzoites and HFFs maintenance and plaque size assays were performed as described (Rosowski et al., 2011).

Transient Transformation and Cas9-Mediated Gene Disruption
DNA sequences targeting GRA17 were cloned into the pSS013 Cas9 vector (Sidik et al., 2014) as described in Supplemental Experimental Procedures. Transient transformations of the pSS013 plasmid were performed by co-electroporating 50–100 μg of circular DNA per strain and 25 μl of pre-annealed donor template oligonucleotides (100 μM) into freshly scrape-syringe-lysed tachyzoites, as described in Supplemental Experimental Procedures, and used to infect HFFs immediately. Confluent HFFs on coverslips were infected for fixation at 24 hr, 48 hr, and 120 hr at MOIs of 2, 0.5, and 0.04, respectively; fixed; and processed for IFA. For cloning purposes, transformed parasites were serially diluted into HFFs grown on 96-well plates without selection. Single-plaque wells were replica-passaged for propagation or for IFA with anti-HA antibodies to determine whether GRA17-HA was disrupted.

Microscopy of Fixed Samples
Microscopy was performed with NIS-Elements software (Nikon) and a digital camera (CoolSNAP EZ; Roper Scientific) connected to an inverted fluorescence microscope (eclipse Ti-S; Nikon) and either phase contrast or DIC.
imaging. Deconvolution imaging was performed using the OptiGrid Structured Light Imaging System (QIOPTIQ Imaging Solutions).

**Cell Fractionation**

Membrane enriched fractions of infected HFFs were obtained by mechanical lysis and differential centrifugation and treated with 0.1 M sodium carbonate (pH 11) or PBS as described (Sibley et al., 1995). The resulting pellet and supernatant were subjected to SDS-PAGE and western blotting.

**Live Microscopy**

HFFs were grown on glass-bottom 24-well plates (Greiner Bio-One) and infected with tachyzoites for 24 hr in growth media. The cells were washed with PBS and replaced with growth media minus phenol red (GMPR) supplemented with 10 μM CCFDA for 10 min at 37°C. CCFDA was sequentially diluted into GMPR from a 10 mM DMSO solution. The dye-containing media was removed, and the cells were washed three times with PBS, replaced with GMPR, and were immediately imaged. scrape-loading experiments were modified from et-Fouly et al. (1987). Briefly, infected cells were prepared as above on glass bottom 24-well plates. Either 0.5 μM lucifer yellow CH or 0.5 μg/ml Dextran3000-Alexa488 were diluted in GMPR and incubated for 5 min at 37°C. The disposable tip of a 10 μl pipette was used to scrape a cross- crossed series of straight lines on the bottom of the well. The cells were washed three times with PBS, replaced with GMPR, and were immediately imaged. For the extended time course experiments, HFFs were infected with the jGRA17 strain. Cells were imaged every 30 min over a 72 hr period using a 40 x objective (NA = 0.95) on a Nikon TE2000 inverted microscope equipped with an environmental chamber, Hamamatsu ORCA-ER digital camera, and NIS Elements Imaging Software.

**Mouse Survival and Physiology**

For mouse infection, infected HFFs were mechanically lysed with 30G needles, washed with PBS, and centrifuged at 582 x g for 7 min to collect tachyzoites and then diluted in PBS and counted. For the survival assays, 5 female CD-1 mice (age 6-8 weeks, Charles River Laboratories) were infected i.p. with 100 tachyzoites per each strain (plus a cohort of mice was infected with 1,000 tachyzoites of jGRA17). Surviving mice were re-infected i.p. at day 30 post-infection with 10,000 tachyzoites of WT and monitored for survival for an additional 21 days. For all infections, relative parasite MOI was assessed by plaque assays, and all surviving mice were verified to be seropositive for Toxoplasma antigen. Mouse below a body condition score of 2 were euthanized and counted as dead in accordance with institutional and federal regulations. All mice were maintained in specific pathogen-free conditions. The MIT Committee on Animal Care approved all protocols (assurance number A-3125-01).

**Electrophysiology**

Xenopus laevis were cared for according to standards approved by the Institutional Animal Care and Use Committee of the University at Buffalo. The frogs were anesthetized in tricaine (1 g/l; Sigma). Oocytes were digested in 82.5 mM NaCl, 2 mM KCl, 1 mM MgCl₂, 5 HEPES (pH 7.4), 1 mg/ml collagenase, and 1 mg/ml protease. The supernatant was subsequently centrifuged at 582 x g for 7 min to collect tachyzoites and then diluted in PBS and counted. For the survival assays, 5 female CD-1 mice (age 6-8 weeks, Charles River Laboratories) were infected i.p. with 100 tachyzoites per each strain (plus a cohort of mice was infected with 1,000 tachyzoites of jGRA17). Surviving mice were re-infected i.p. at day 30 post-infection with 10,000 tachyzoites of WT and monitored for survival for an additional 21 days. For all infections, relative parasite MOI was assessed by plaque assays, and all surviving mice were verified to be seropositive for Toxoplasma antigen. Mouse below a body condition score of 2 were euthanized and counted as dead in accordance with institutional and federal regulations. All mice were maintained in specific pathogen-free conditions. The MIT Committee on Animal Care approved all protocols (assurance number A-3125-01).

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes seven figures, two tables, four movies, and Supplemental Experimental Procedures and can be found with this article online at http://dx.doi.org/10.1016/j.chom.2015.04.003.

**AUTHOR CONTRIBUTIONS**


**ACKNOWLEDGMENTS**


Received: September 8, 2014
Accepted: March 20, 2015
Published: May 13, 2015

**REFERENCES**


Supplemental Information

The *Toxoplasma* Dense Granule Proteins GRA17 and GRA23 Mediate the Movement of Small Molecules between the Host and the Parasitophorous Vacuole

Fig. S3

A

<table>
<thead>
<tr>
<th>Lane</th>
<th>WT</th>
<th>ΔGRA17</th>
<th>ΔGRA23</th>
<th>ΔGRA17-HA</th>
<th>WT</th>
<th>ΔGRA17-HA</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>extra</td>
<td>1.4</td>
<td>1.4</td>
<td>1.8</td>
<td>2</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>intra</td>
<td>1.1</td>
<td>0.7</td>
<td>1.2</td>
<td>0.8</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

---

B

Fold expression over WT

- ΔGRA17
- ΔGRA17 + GRA17-HA
- ΔGRA23

---

C

GRA16-HA-FLAG transient transformation

WT

ΔGRA17

ΔGRA23

post-infection: 2 h

20 h

---

D

ΔGRA17 scrape loading

Present

Intermediate

Absent

Fluorescence intensity (au)
Figure S1. Sequence alignment and analysis of GRA17, related to Table 1.

(A) Alignment of the primary peptide sequences of *Toxoplasma gondii* GRA17 (TGGT1_222170)\(^1\) and GRA23 (TGGT1_297880), *Neospora caninum* NCLIV_005560 and NCLIV_006780, *Sarcocystis muris* DG32 (O43946_SARMU), *Sarcocystis neuron* A and B (unannotated), *Eimeria acervulina* EAH_00026470, *E. mitis* EMH_0012530 and the EXP2 genes from *Plasmodium falciparum* (PF3D7_1471100), *Plasmodium knowlesi* (PKH_123420), and *Plasmodium reichenowi* (PRCDC_1470300) using Profile ALiGnMEnt (PRALINE) (Bawono and Heringa, 2014) which scores amino acid conservation. The scoring scheme works from 0 for the least conserved alignment position, up to 10 for the most conserved alignment position. The color assignments are depicted in the legend (*above*). (B) The phylogenetic tree represents full-length peptide sequences from the previous alignment, using the Neighbor-Joining method with the program MEGA5 (Tamura et al., 2011). The percentage of replicate trees in which the associated taxa clustered together in the bootstrap test (500 replicates) is shown next to the branches. The tree is drawn to scale (*below*), with branch lengths in the same units as those of the evolutionary distances used to infer the phylogenetic tree. The evolutionary distances were computed using the JTT matrix-based method and are in the units of the number of amino acid substitutions per site. All positions containing gaps and missing data were eliminated.

\(^1\) A *Toxoplasma* Kozak sequence is present immediately upstream of the second methionine (residue 58) from the predicted GRA17 coding sequence on ToxoDB.org (v13.0) and the observed size of the GRA17 band (19 kDa; Fig. S3A) matches the signal peptidase cleavage product predicted from the GRA17 coding sequence starting from that second methionine in contrast to the size discrepancy that exists with the current gene model on ToxoDB.org (v13.0). This suggests that Met58 (from the current gene model) is the actual GRA17 start codon.
Figure S2. Confirmation of gene knockouts and transgenic expression, related to Figure 1.

(A) Schematic diagram depicting the genomic loci of the genes of interest (GOI) (top) and the gene knockout-targeting construct (pTKO2-HXGPRT or pTKO2-DHFR-TS) used to disrupt GOI loci (bottom). The construct consists of the HXGPRT or DHFR-TS selection cassettes flanked by UTRs derived from TgDHFR, which in turn is flanked by 5’ and 3’ targeting sequences (described in Experimental Procedures) corresponding to regions flanking the GOI coding sequences. P1 and P2 refer to primers annealing within the coding region of the GOI (either GRA17 or GRA23). P3 and P5 are sense and antisense primers annealing within the HPT or DHFR-TS selection cassettes. P6 and P4 refer to sense and antisense primers annealing outside the 5’ and 3’ flanking sequences surrounding either the GRA17 or GRA23 loci. (B) Schematic diagram depicting the genomic locus of TgUPRT (top) and the knock-in-targeting construct (pTKO2-UPRT) containing the transgenes used to disrupt the UPRT loci (bottom). The construct encodes the transgenes and accompanying UTRs, which in turn are flanked by 5’ and 3’ targeting sequences (described in Experimental Procedures) corresponding to the UPRT loci. P1, P2, and HA are sense and antisense primers annealing within the coding region of the transgenes. P7 and P8 refer to sense and antisense primers annealing outside the 5’ and 3’ targeting sequences surrounding the UPRT locus. (C) The absence of PCR products amplified by P1/P2 from genomic DNA derived from the ΔGRA17 and ΔGRA23 strains (in contrast to WT) verify of the disruption of the GRA17 and GRA23 loci (top two panels, respectively). The PCRs products amplified by the P6/P5 and P3/P4 primers verify the orientation of the knockout construct within the GRA17 or GRA23 loci (bottom four panels). (D) The PCRs products amplified by the P6/P5 and P3/P4 primers from genomic DNA derived from the ΔGRA17/ΔGRA23+GRA17-HA strain (in contrast to WT and ΔGRA17+GRA17-HA strains) verify the orientation of the DHFR-TS knockout
cassette within the GRA23 locus (top two panels). The PCRs products amplified by the P7/HA, P1/P8, and P1/HA primers from genomic DNA derived from the ΔGRA17+GRA17-HA and ΔGRA17/ΔGRA23+GRA17-HA strains (in contrast to WT) verify the presence and orientation of the GRA17-HA transgene within the UPRT locus (third, fourth, and fifth panels from top). The absence of a PCR product amplified by P1/P2 from genomic DNA derived from the ΔGRA17/ΔGRA23+GRA17-HA strain (in contrast to ΔGRA17+GRA17-HA) verifies the disruption of the GRA17 locus (bottom panel). (E) The PCRs products amplified by the P7/HA and P1/P8 primers from genomic DNA derived from the ΔGRA17+EXP2-HA and ΔGRA17+(gra1-5'UTR)-GRA23-HA-FLAG (HF) strains (in contrast to WT) verify the presence and orientation of the (gra1-5'UTR)-EXP2-HA (top two panels) and (gra1-5'UTR)-GRA23-HF_{high} (bottom two panels) transgenes within the UPRT locus. (F) Wild-type (WT), ΔGRA17+(gra23-5'UTR)-GRA23-HF_{low}, and ΔGRA17+(gra1-5'UTR)-GRA23-HA-FLAG_{high} extracellular parasites were lysed with sample buffer, subjected to SDS-PAGE and Western blotted with anti-HA and anti-SAG1 antibodies. (G) HFFs were infected with WT+GRA17-V5/GRA23-HF as in Fig. 1A except that the samples were fixed with methanol to quench GFP fluorescence and stained with anti-HA and anti-V5 antibodies as indicated (scale bar = 10 µm). (H) HFFs were infected for 24 h with ΔGRA17+PfEXP2-HA, ΔGRA17+(gra23-5'UTR)-GRA23-HA-FLAG_{low}, and ΔGRA17+(gra1-5'UTR)-GRA23-HA-FLAG_{high} parasites, fixed and stained with anti-HA and anti-SAG1 antibodies and imaged by IF and DIC microscopy; scale bar = 5 µm. (I) Extracellular tachyzoites were fixed and stained with anti-HA (red) and anti-GRA7 (green) antibodies and imaged with deconvolution as above and merged with a DIC brightfield image; scale bar = 5 µm.

Figure S3. GRA17 physical and functional characterization, related to Figure 2. (A) HFFs were infected with the indicated Toxoplasma strains for 24 h and lysed to obtain intracellular (intra) lysates or tachyzoites were allowed to completely egress to
obtain extracellular (extra) lysates, which were then subjected to SDS-PAGE and Western blotted with either serum raised against a C-terminal epitope of GRA17 or antibodies against *Toxoplasma* Surface Antigen (SAG)-1, used here as a loading control. The *arrows* distinguish between the epitope-tagged recombinant and endogenous forms of GRA17. The *asterisk* denotes a non-specific band found in the intracellular lysates. The designations “r” and “e” refer to the recombinant and endogenous species of GRA17, respectively. SAG1 bands were quantified by optical density and normalized to either intracellular or extracellular WT lanes as indicated. (B) Total mRNA was isolated and sequenced from bone marrow-derived mouse macrophages that were infected with equivalent MOI-matched wild-type, ΔGRA17, ΔGRA17+GRA17-HA, and ΔGRA23, and tachyzoites for 4 hours. The expression data is expressed as fold change of the transcript counts of the *GRA17* (*red*), *GRA23* (*blue*), *GRA7* (*charcoal*) and *T. gondii* GAPDH (*orange*) genes compared to wild-type levels (*dotted line*). This experiment was performed once. (C) Wild-type, ΔGRA17, and ΔGRA23 tachyzoites were transiently transformed with GRA16-HA-FLAG and immediately used to infect HFFs and fixed at 2 h and 20 h post-infection and subjected to IF with anti-HA antibodies (*green*) or stained with Hoechst dye (*red*) and overlaid on a DIC image. This experiment was performed twice; scale bar = 10 µm. (D) Sample images of ΔGRA17-infected HFFs that have been scrape-loaded with Lucifer Yellow dye (*green*) alone or overlaid onto a DIC image that exemplify the intra-PV fluorescence scores as described in Fig. 2C (*left panel*). Using the NIS Elements software, fluorescence intensity (arbitrary units) was plotted across a line of defined length within the vacuole or as a reference, outside the vacuole for each example (*right panel*; scale bar = 10 µm).

**Figure S4. GRA17 dynamically affects PV morphology, related to Figure 3.**
Sub-confluent HFFs on were infected with ΔGRA17 tachyzoites and imaged live under identical conditions as in Fig. 3A. Images obtained from two different cells; phase-contrast microscopy from indicated time-points. (B) As in (A), time-points of images from one cell range from 20 h to 51:30 h post-infection when the tachyzoites egress. The red asterisks indicate the same PV followed over time, which along with its host cell moved out of the fixed field of view times over the experimental time course. Scale bar = 10 µm for both subfigures.

**Figure S5. GRA17 affects in vitro and in vivo growth, related to Figure 4.**

(A) Representative plaque images from the analysis in Fig. 3C from each strain are presented. Plaques were imaged at 4× magnification by phase-contrast microscopy taken out of phase to increase contrast; scale bar = 0.5 mm. (B) Confluent monolayers of HFFs in individual wells of 24-well plates were infected with 300 tachyzoites of each indicated type III strain for 7 days. The area of at least 30 plaques from each strain was measured and the mean area was normalized to wild-type (WT). This experiment was performed once. (C) Five female BALB/c mice were each infected with 10⁴ tachyzoites intraperitoneally with the indicated strains expressing firefly luciferase and were imaged in vivo at 4 days post-infection. One representative of the 5 mice infected is shown (left). Each bar of the graph (right) represents a mean of the total flux (photons/sec/cm²/sr) per mouse of each strain as an assessment of parasite burden. Error bars represent ±SD and a Student’s t test was used to measure statistical significance (P=0.05). (D) The average weight of the five mice infected with each strain in (B) was plotted over time as an assessment of overall health. Error bars represent ±SD and a two-tailed Student’s t test was used to measure statistical significance. (E) At 45 days post-infection, *Toxoplasma* tissue cysts were counted by staining homogenates of brains derived from each mouse with dolichos biflorus-FITC, a marker for the cyst wall. Cysts from each mouse brain homogenate were diluted and blindly counted in quadruplicate and back-
calculated to give the total number of cysts per brain (circle or square plots). The horizontal red lines represent the mean number of total brain cysts per cage of mice infected and error bars represent the standard deviation across diluted replicates for each brain (P = 0.13, student’s t test).

**Figure S6. Cas9-mediated disruption of GRA17, related to Figure 5.**

Confluent HFFs were infected with ΔGRA17/ΔGRA23+GRA17-HA parasites that were transiently transformed with Cas9-3xFLAG and either non-specific or GRA17-specific gRNAs along with GRA17 donor oligonucleotide templates containing a nonsense mutation (described in Supplemental Experimental Procedures). The infected cells were fixed at 24 h and subjected to IF with anti-HA and anti-FLAG antibodies. The percentage of Cas9-3xFLAG-positive vacuoles and GRA17-HA-positive vacuoles out of the total were scored. This experiment was performed only once.

**Figure S7. GRA17, GRA23, GRA15, and EXP2 expression in Xenopus oocytes, related to Figure 6.**

(A) *Xenopus laevis* oocytes injected with 29 ng of GRA17-HA or 33 ng GRA23-HA cRNAs or mock injected for 3 days and snap-frozen. The oocytes were lysed and fractionated into total membrane and cytosol fractions as described in Supplementary Experimental Procedures. The lysates were subjected to SDS-PAGE and Western blotted with anti-HA and anti-GAPDH antibodies as indicated. (B) *Xenopus* oocytes were injected with 32 ng of GRA15-HA cRNA or mock injected for 3 days and fractionated into total membrane (M) or cytosol (C) fractions and Western blotted as in Fig. S7A. (C) *Xenopus* oocytes were injected with 29 ng of GRA17-HA, 33 ng of GRA23-HA cRNA, or equivalent volume water for 3 days and were snap-frozen, cryo-sectioned, fixed and processed by immunohistochemistry with anti-HA antibodies. Images were taken at 20× magnification using either phase-contrast brightfield (BF) or fluorescent deconvolution microscopy (scale bar = 50 µm). The oocyte membrane is indicated (white arrows) in
contrast to the background fluorescence of the pigment granules. (D) *Xenopus* oocytes were injected with 54 ng of *EXP2-HA* cRNA or mock injected for 3 days and fractionated into total membrane (M) or cytosol (C) fractions and Western blotted as in Fig. S7A. (E) Percent per condition survival of *Xenopus laevis* oocytes injected with 30 ng of *PIEXP2-HA* mRNA or equal volume H$_2$O, 3 days post-injection. Survival was based on a morphological appearance of the oocytes. Fractions on top of each bar graph represent the number of oocytes that survived out of the total injected for each condition. (F) Resting membrane potentials (RMP) of surviving oocytes injected with *PIEXP2-HA* cRNA or equal volume H$_2$O. (G) Average conductance (G) was computed by fitting the I-V relationship as a straight line. (H) Average membrane currents measured from a holding potential of -90 mV potentials of -120 mV to +50 mV were applied for 2 seconds. Peak currents from each applied voltage were used to generate an I-V relationship. For all above sub-figures, data are shown as the mean ± SE. Confidence levels were calculated using Student’s paired t-test, where * signifies P<0.05 when compared to H$_2$O-injected oocytes.

**Table S1. DNA oligonucleotides used in this study, related to Figure 1.**

The name, sequence and brief description of each DNA oligonucleotide is presented in table form.

**Table S2. Genes uniquely up-regulated in the absence of GRA17, related to Figure 2.**

Analysis of the mRNA sequenced from the *Toxoplasma*-infected bone-marrow-derived macrophages in Fig. S3B. The data was filtered for genes expressed above an FPKM value threshold of 20 and containing strongly predicted signal peptides by SignalP 4.1. Relative expression was calculated by taking the binary logarithms of the ratio of FPKM values of either the ΔGRA17 or ΔGRA23 strains compared to the FPKM value the Wild-
type strain for each gene and filtered to include only those values ≥ 0.8 or ≤ 0.2, respectively. The ToxoDB.org gene identities, gene product names and values of the relative expression calculated above are listed in the table.

**Videos S1 and S2. A subset of ΔGRA17 PVs collapse and their tachyzoites become inviable over time, related to Figure 3.**

**Videos S1 and S2** depict the same infected host cell as described in Figure 3A. Sub-confluent HFFs were infected with ΔGRA17 tachyzoites. **Video S1** spans 19:30 h to 35:30 h post infection. Note the collapse of all but one PV in the host cell over this time course. **Video S2** spans a duration from 41:30 h to 62:00 h post-infection. Note the remaining PV contains dividing tachyzoites that eventually egress from the host cell but the tachyzoites in the collapsed PVs remain inviable. Images in both videos were collected every 30 minutes from a fixed field of view. Scale bars represent 10 μm.

**Supplemental Videos 3 and 4. ΔGRA17 enlarged PVs change shape and size dynamically and contain viable tachyzoites, related to Figure 3.**

**Videos S3 and S4** depict the same infected host cell as described in Figure S5. Sub-confluent HFFs were infected with ΔGRA17 tachyzoites. **Video S3** spans 21:30 h to 37:00 h post-infection. Note the dynamic growth of the PV, doubling of the number of tachyzoites and movement of the infected cell over this time course (imaged from a fixed point). **Video S4** spans a duration from 37:30 h to 51:30 h post-infection. Note the continued growth of the PV and the ultimate egress of the tachyzoites within it. Images in both videos were collected every 30 minutes; scale bars = 10 μm.
Table S2

<table>
<thead>
<tr>
<th>Gene ID</th>
<th>Product Description</th>
<th>$\log_2 (\Delta \text{GRA17/WT})$</th>
<th>$\log_2 (\Delta \text{GRA23/WT})$</th>
</tr>
</thead>
<tbody>
<tr>
<td>TGME49_278400</td>
<td><em>T. gondii</em> family A protein</td>
<td>1.3</td>
<td>0.2</td>
</tr>
<tr>
<td>TGME49_319350</td>
<td>SAG-related sequence SRS17B</td>
<td>1.0</td>
<td>-0.2</td>
</tr>
<tr>
<td>TGME49_297880</td>
<td>GRA23</td>
<td>0.8</td>
<td>-5.9</td>
</tr>
<tr>
<td>TGME49_270170</td>
<td>hypothetical protein</td>
<td>0.8</td>
<td>0.2</td>
</tr>
<tr>
<td>TGME49_310610</td>
<td>hypothetical protein</td>
<td>0.8</td>
<td>-0.5</td>
</tr>
<tr>
<td>TGME49_222270</td>
<td>hypothetical protein</td>
<td>0.8</td>
<td>0.2</td>
</tr>
</tbody>
</table>

Supplemental experimental procedures

Antibodies

Rabbit polyclonal antibodies were raised against a synthetic C-terminal peptide derived from *TgGRA17* (CVPRLEALSAKMAVKQKAMQG) that was conjugated to keyhole limpet hemocyanin (KLH). The peptide synthesis, rabbit immunization and serum collection was conducted by Covance Research Products (Denver, PA). We used serum at a 1:1000 dilution for Western Blotting; this antibody fails to detect GRA17 by immunofluorescence (IF) microscopy. We used antibodies raised against HA (3F10, rat polyclonal, Roche, 1:500 for WB and IF; 1:100 for immunohistochemistry), FLAG (M2 mouse monoclonal, Sigma, 1:1000), GAPDH (6C5, mouse monoclonal, Santa Cruz, 1:750), V5 (mouse monoclonal, Life Technologies, 1:1000), *Toxoplasma* Surface Antigen (SAG)-1 (Burg et al., 1988) (DG52, purified mouse monoclonal, 1:5000 dilution for IF), *TgSAG1* (rabbit polyclonal, gift of J. Boothroyd, 1:5000 for Western Blotting or IF), *TgGRA4* (rabbit polyclonal, gift of L. Knoll, 1:2000), *TgGRA7* (Dunn et al., 2008) (rabbit polyclonal, gift of J. Boothroyd, 1:2000), *TgGRA1* (Charif et al., 1990), and *TgROP1* (Ossorio et al., 1992). Secondary antibodies for IF were conjugated to Alexa-
488 and Alexa-594 (Life Technologies), and HRP-conjugated antibodies (Kirkegaard & Perry Laboratories) were used for Western Blotting.

**Generation of transgenic, knockout, and complemented strains**

We expressed GRA17 (TGGT1_222170) as a transgene by amplifying its coding region with the primers TGGT1_222170_-1500_F and TGGT1_222170_HA_R (or TGGT1_222170_V5_R) containing the putative endogenous GRA17 promoter from genomic DNA isolated from *Toxoplasma GT1* to clone into the pTKO-att vector that encodes GFP and the hypoxanthine-xanthine-guanine ribosyl transferase (HXGPRT) selectable marker using a Gateway (Life Technologies) cloning strategy described in (Rosowski et al., 2011). Electroporation, selection by mycophenolic acid/xanthine and cloning by limiting dilution were performed as previously described (Donald and Roos, 1998; Rosowski et al., 2011) into RHΔHXGPRT and CEPΔHXGPRT C22 (Boyle et al., 2007) (types I, II, and III respectively). To construct the vector pLIC-P_{GRA23}-GRA23-HA-FLAG, the promoter region of GRA23 (TGGT1_297880) and GRA23 coding sequence was amplified using the primers LIC_ TGGT1_297880_F and LIC_ TGGT1_297880_R from genomic DNA of the RH wild-type strain. The resulting PCR product was cloned into the pLIC-HF-dhfr vector using the LIC cloning method as described (Huynh and Carruthers, 2009) to generate the pLIC-GRA23-HF-dhfr plasmid.

We generated GRA17 and GRA23 knockouts with targeting constructs derived from the pTKO2 vector and Multisite Gateway Pro 3-Fragment Recombination (Life Technologies) strategy previously described (Rosowski et al., 2011) (Fig. S2A). Briefly, the 5’ and 3’ UTRs of GRA17 or GRA23 were engineered to flank the HXGPRT selectable marker in the pTKO2 plasmid. The fragment containing the GRA17 5’UTR was amplified with TGGT1_222170_5’UTR_attB1 and TGGT1_222170_5’UTR_attB4. The GRA17 3’UTR was amplified with TGGT1_222170_3’UTR_attB2 and TGGT1_222170_3’UTR_attB3. The fragment containing the GRA23 5’UTR was
amplified with primers TGGT1\_297880\_5'UTR\_attB1 and TGGT1\_297880\_5'UTR\_attB4. The GRA23 3'UTR was amplified with primers TGGT1\_297880\_3'UTR\_attB3 and TGGT1\_297880\_3'UTR\_attB2. An additional GRA23 gene targeting construct was constructed using the above GRA23 UTRs with a DHFR-TS selectable marker cassette (Donald and Roos, 1994). The DHFR-TS cassette was cloned into the Gateway pDONR 221 P4r-P3r vector by amplifying the cassette with the following forward and reverse primers: DHFR-TS\_attB4r and DHFR-TS\_attB3r. Electroporation, selection by mycophenolic acid/xanthine and cloning by limiting dilution were performed as described above into RHΔHXGPRT/ΔKu80 (Fox et al., 2009). The parasites were confirmed to contain gene knockouts by our ability to amplify the knockout construct and inability to amplify the coding region by PCR in these clones (Fig. S2A, B, and C). The RHΔGRA17 strain was complemented by disrupting the *Toxoplasma uracil phosphoribosyl transferase (UPRT)* locus with GRA17-HA or *Toxoplasma* codon-optimized *Plasmodium falciparum* EXP2-HA and positively selecting for the absence of UPRT with floxuridine (FUDR) (Bohne and Roos, 1997). The targeting construct was generated into pTKO2 using the approach outlined above, with the GRA17-HA transgene flanked by the 5' and 3' UTRs from *UPRT*. The GRA17-HA transgene was amplified from the cloned GRA17 sequence contained in the pTKO-att plasmid described above using the TGGT1\_222170\_attB4r and TGGT1\_222170\_attB3r primers for cloning into pDONR221 P4r-P3r. The 5' and 3'UTRs of *UPRT* were amplified with the following forward and reverse primer pairs from isolated *Toxoplasma* RH genomic DNA: (5'UTR, forward: UPRT\_5'UTR\_attB1 and UPRT\_5'UTR\_attB4; 3'UTR, forward: UPRT\_5'UTR\_attB3 and reverse: UPRT\_5'UTR\_attB2) for cloning into the pDONR221 P4-P1 and P3-P2, respectively. The chimeric gra1-5'UTR-GRA23-HA-FLAG transgene was generated by a 2-step “sewing PCR” where the gra1 5'UTR fragment (#1) was PCR amplified with the gra1\_5'UTR\_attB4r and gra1\_5'UTR\_GRA23bridge\_R
primers from a RH-derived genomic DNA template such that it contains a 3’ extension of
GRA23 coding sequence. A GRA23-HA-FLAG coding sequence fragment (#2) was
generated by PCR amplification using the gra1_5'UTR_GRA23bridge_F and DHFR-
TS_attB3r with the pLIC-GRA23-HF-dhfr plasmid as template. Fragment #2 contains a
5’extension containing gra1 5’UTR sequence that overlaps with the first fragment,
allowing a second round of PCR amplification using the gra1_5’UTR_attB4r and DHFR-
TS_attB3r primers and fragments #1 and #2 as templates to generate the chimeric gra1-
5’UTR-GRA23-HF-dhfr sequence to combine with the UPRT flanking sequences as
described above.

Endogenous tagging of Toxoplasma GRA24 was performed by digesting the pLIC-PGRA1-
GRA24-HA-FLAG (Braun et al., 2013) plasmid DNA with a restriction endonuclease
within the coding sequence and transforming the target strains as previously described
(Huynh and Carruthers, 2009).

The Plasmodium falciparum EXP2 cDNA was codon optimized for E. coli by DNA 2.0
(Menlo Park, CA) using proprietary parameters that optimize expression by codon usage
as well as RNA stability using secondary structure predictions. The resulting EXP2
cDNA (below) was PCR amplified, adding an HA tag to the C-terminus with
EXP2_Nsil_F and EXP2_Paci_R primers and cloned into the pGRAGFP/HPT plasmid
(Rome et al., 2008) in which GFP was removed and the EXP2-HA coding sequence
added using Nsil and PacI restriction sites.

This PfEXP2-HA expression cassette was then amplified with the following forward and
reverse primers for Gateway cloning into the pDONR 221 P4r-3r: EXP2_attB4r and
EXP2_attB3r. Toxoplasma-codon optimized PfEXP2 coding sequence:

5'atgcatAAAGTTAGCTACATCTTTAGCTTCTTCTTGCTGTTCTTCGTCTACAAGAACAC
GAATACCGTCGTTTGCGACAATGGTTACGGTGACTTGGCTGCGACCTCCCGCGTTA
CGACGTTATCAAAAGATCCGATTTCCGATTTCCCGAAGACATTTACGAGCACCGG

Design and cloning of guide RNA for Cas9-mediated gene disruption

The GRA17gRNA_L32stop+ and GRA17gRNA_L32stop- oligonucleotides encoding a target sequence within GRA17 were annealed and ligated into the guide RNA (gRNA) expression cassette in a BsaI-digested pSS013 vector (Sidik et al., 2014), also encoding Cas9-3xFLAG. The GRA17donor_L32stop+ and GRA17donor_L32stop- oligonucleotides were annealed for co-electroporation with the pSS013 vector to serve as a template for homologous recombination. As above, the following oligos were used to create gDNA/oligonucleotide donor template combinations: Gly20stop [gRNA oligos: GRA17gRNA_G20stop+ and GRA17gRNA_G20stop-; donor template oligos: GRA17donor_G20stop+ and GRA17donor_G20stop-]; Lys54stop [gRNA oligos: GRA17gRNA_K54stop+ and GRA17gRNA_K54stop-; donor templates: GRA17donor_K54stop+ and GRA17donor_K54stop-].

Western blotting and immunofluorescence
Infected HFFs were lysed and samples were prepared for Western blotting as described (Rosowski et al., 2011). Extracellular parasites were derived from cells that were newly lysed out and washed 3× with PBS and lysed for Western blotting as above. IF was performed as previously described (Rosowski et al., 2011).

**Extracellular tachyzoite staining**

Extracellular parasites were derived from cells that were freshly lysed out and washed 3× with PBS, centrifuged onto coverslips at 163 × g for 3 minutes and subjected to IF.

**RNA sequencing**

Bone marrow-derived mouse macrophages were isolated and infected as previously described (Hassan et al., 2012). Total RNA excluding miRNA was extracted with the Qiagen RNeasy Plus kit (Qiagen, USA) and assayed for integrity, size distribution and concentration with the Agilent 2100 Bioanalyser. The mRNA was isolated with the Dynabead mRNA purification kit (Invitrogen) and processed for sequencing on an Illumina HiSeq 2000 machine according to standard Illumina protocols. The sequences were assembled and aligned to the ME49 8.0 genome as previously described (Hassan et al., 2012). The accession number for the mRNA sequencing data reported in this paper is GEO: GSE65980.

**In vivo mouse imaging and physiology**

5 female BALB/c (age 6 weeks, The Jackson Laboratory) were infected intraperitoneally with 10,000 tachyzoites and imaged *in vivo* as described previously (Rosowski et al., 2011). Brain homogenate of chronically infected mice was stained with dolichos biflorus-FITC (Vector Laboratories), and cysts were enumerated by microscopy. For all infections, plaque assays were performed after the experiment to assess relative parasite MOI and found to yield an equivalent number of plaques across strains per experiment and all surviving mice were verified to be seropositive for *Toxoplasma* antigen. Mice determined to be below a body condition score of 2 were euthanized and
counted as dead in accordance with institutional and federal regulations. All mice were maintained in specific pathogen-free conditions. The MIT Committee on Animal Care approved all protocols (assurance number A-3125-01).

**Cloning and preparation of cRNA for Xenopus oocyte injection**

To clone cDNAs of GRA17, GRA23, and GRA15, total RNA was isolated from infected cells and a cDNA pool was created as described (Rosowski et al., 2011) and PCR-amplified with the following forward and reverse primers, respectively (GRA17: GRA17_-SigP_BamHI_F and GRA17_NotI_R or GRA17_HA_NotI_R; GRA23: GRA23_-SigP_BamHI_F and GRA23_NotI_R or GRA23_HA_NotI_R; GRA15: GRA15_-SigP_BamHI_F and GRA15_SpeI_R or GRA15_HA_SpeI_R; EXP2: EXP2_-SigP_BamHI_F and EXP2_NotI_R or EXP2_HA_NotI_R) for ligation into pBluescript KS+ with BamHI and NotI restriction sites for GRA17, GRA23, and EXP2 and BamHI and SpeI restriction sites for GRA15. The start sites for each cDNA were chosen immediately after the predicted signal peptide cleavage site and a heterologous Kozak sequence and start codon were added to the upstream primers. An additional heterologous glycine residue was added after the start codon of the GRA15 cDNA for stability. The plasmids were digested with PvuI endonuclease and cRNA was generated and prepared for oocyte injection as described (Bett et al., 2012).

**Isolation of total membranes and cytosol from Xenopus oocytes**

As described in (Engelund et al., 2013; Kamsteeg and Deen, 2001) injected oocytes (~12 oocytes per tube) were mechanically lysed with a p200 pipette tip in HbA buffer [20 mM Tris-HCl (pH7.4), 5 mM MgCl2, 5 mM NaH2PO4, 1 mM EDTA, 80 mM sucrose and 25× of a protease inhibitor cocktail (Cat# 04693116001, Roche)]. The lysates were centrifuged at 200 × g at 4 °C for 5 minutes and the resulting supernatant was taken and centrifuged at 20,000 × g at 4 °C for 20 minutes. The supernatant (cytosolic fraction) and
the pellet (the total membrane fraction) were separated and treated with 2× Laemmli Sample Buffer, boiled, and subjected to SDS-PAGE and Western blotting.

**Xenopus oocytes immunohistochemistry**

cRNA-injected *Xenopus* oocytes were snap frozen unfixed in OCT freezing media. The blocks were cryo-sectioned into 6 µm slices onto glass slides. The slides were then fixed with 4% paraformaldehyde for 15 minutes at room temperature. The slides were washed 2× with PBS and then incubated with 0.25% Triton X-100 in PBS for 10 minutes. The slides were washed 3× with PBS and blocked for 1 hour in IHC Blocking Buffer (IHC-BB) [1 mg/ml Blocking Reagent (Perkin Elmer, Cat# FP1012), 2% heat-inactivated goat serum, in 0.1 M Tris, pH 7.5, and 0.15 M NaCl]. The slides were incubated with primary antibody diluted in IHC-BB overnight at 4 °C and subsequently washed 3× with PBS. The slides were incubated with goat anti-rat Alexa-594 (Life Technologies) secondary antibody diluted 1:500 in IHC-BB for 1 hour at room temperature. The slides were washed 3× with PBS and mounted for imaging.

**Supplemental References**


