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Identification of three novel Toxoplasma gondii rhoptry proteins

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2 **Identification of three novel *Toxoplasma gondii* rhoptry proteins**

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18 Note: Supplementary data associated with this article.

19 **Abstract**

20 The rhoptries are key secretory organelles from apicomplexan parasites that contain proteins
21 involved in invasion and modulation of the host cell. Some rhoptry proteins are restricted to the
22 posterior bulb (ROPs) and others to the anterior neck (RONs). As many rhoptry proteins have been
23 shown to be key players in *Toxoplasma* invasion and virulence, it is important to identify,
24 understand and characterize the biological function of the components of the rhoptries. In this
25 report, we identified putative novel rhoptry candidate genes by identifying *Toxoplasma* genes with
26 similar cyclical expression profiles as known rhoptry protein encoding genes across its cell cycle.
27 Using this approach we identified two new rhoptry bulb (ROP47 and ROP48) and one new rhoptry
28 neck protein (RON12). ROP47 is secreted and traffics to the host cell nucleus, RON12 was not
29 detected at the moving junction during invasion. Deletion of ROP47 or ROP48 in a type II strain
30 did not show major influence in in vitro growth or virulence in mice.

31

32 *Keywords: Toxoplasma gondii, Rhoptry, Rhoptry neck, Host-pathogen interaction*

33

34 **1. Introduction**

35 *Toxoplasma gondii* is a highly successful parasite infecting approximately 30% of people
36 worldwide and is the second largest cause of death due to foodborne illness in the United States
37 (Scallan et al., 2011). Toxoplasmosis can be fatal in the unborn fetus and in immunosuppressed
38 individuals if the disease is not recognized and treated early, in contrast to immunocompetent
39 patients in whom it is mainly self-limited (Montoya and Liesenfeld, 2004; Scallan et al., 2011).

40 *Toxoplasma* resides within a non-fusogenic parasitophorous vacuole and has three apical secretory
41 organelles: the dense granules, micronemes and rhoptries. Rhoptries are club-shaped organelles
42 divided into two distinct compartments, the posterior bulb and the more anterior duct (neck)
43 through which rhoptry proteins are secreted. Proteins derived from the rhoptry secretory organelles
44 are crucial for the invasion and survival of apicomplexan parasites within host cells and thus
45 rhoptry protein targeting is a vital process for *Toxoplasma*. Some rhoptry proteins are restricted to
46 the bulb (ROPs) and others to the neck (RONs). The role of rhoptries in the invasion process,
47 virulence and/or host cell modulation has been well documented, although the molecular
48 mechanisms remain only partially understood. RONs 2, 4, 5 and 8 have been shown to be involved
49 in parasite invasion of the host cell (Besteiro et al., 2009; Straub et al., 2009); ROP16 activates the
50 transcription factors STAT3 and STAT6 (Saeij et al., 2007; Yamamoto et al., 2009; Ong et al.,
51 2010); ROP38 downregulates Mitogen Activated Protein Kinase (MAPK) pathway activation
52 (Peixoto et al., 2010); and ROP5 and ROP18 act jointly to block immunity-related GTPase (IRG)
53 mediated clearance of the parasite by the host cell (Behnke et al., 2012; Fleckenstein et al., 2012;
54 Niedelman et al., 2012). Once rhoptry proteins are secreted into the host cell cytosol they traffic to
55 distinct cellular destinations. For example, ROP16 and the rhoptry protein phosphatase 2 C (PP2C-
56 hn) carry a nuclear localization signal (NLS), which mediates their trafficking to the host nucleus
57 (Gilbert et al., 2007; Saeij et al., 2007), ROP5 and ROP18, following secretion into the host cell,
58 traffic back to the outside of the parasitophorous membrane vacuole (PVM) via an arginine-rich
59 amphipathic helix domain that is essential for this localization (Reese and Boothroyd, 2009);

60 Fentress et al., 2012). Other rhoptry proteins, such as Toxofilin, remain in the host cell cytosol upon
61 secretion (Lodoen et al., 2010).

62 Rhoptry proteins contain a classic eukaryotic signal peptide for entrance into the secretory
63 pathway and are trafficked from the endoplasmic reticulum through the Golgi by a conserved
64 pathway before being packaged into the apically located secretory organelles (Sadak et al., 1988;
65 Bradley and Boothroyd, 1999; Bradley et al., 2004; Carey et al., 2004; Hajj et al., 2006b, 2007;
66 Turetzky et al., 2010). N-terminal pro-domains have been implicated in rhoptry protein sorting and
67 indeed several rhoptry proteins exhibit N-terminal processing. However, the failure to remove the
68 pro-domain does not seem to disrupt targeting (Bradley et al., 2002; Miller et al., 2003; Turetzky et
69 al., 2010). Other rhoptry proteins do not appear to be processed and the mechanism by which they
70 are targeted to the rhoptries is unknown. Additionally, any soluble protein recombinantly fused to a
71 signal peptide is delivered to the dense granules by default (Joiner and Roos, 2002). Therefore,
72 motifs that mediate trafficking of proteins to the rhoptry organelles are insufficiently defined, or
73 insufficiently specific, to allow genome-wide identification of rhoptry proteins.

74 Proteomic and genomic approaches have been widely used to identify the contents of the
75 rhoptries of apicomplexan parasites (Hoppe et al., 2000; Bradley et al., 2005; Peixoto et al., 2010;
76 Marugán-Hernández et al., 2011; Reid et al., 2012; Oakes et al., 2013). A proteomic study of
77 *Toxoplasma* rhoptry contents led to the identification of 38 rhoptry proteins (Bradley et al., 2005).
78 Twenty of these proteins were shown to localize to the rhoptry organelles, 11 to the rhoptry bulb
79 and nine to the rhoptry neck (Bradley et al., 2005; Taylor et al., 2006; Gilbert et al., 2007;
80 Proellocks et al., 2009; Straub et al., 2009; Peixoto et al., 2010; Lamarque et al., 2012). As
81 expected, several previously known rhoptry proteins were readily detected in this proteomic
82 analysis. However, TgNHE2 and TgSUB2, previously characterized as rhoptry proteins, were
83 missed. The absence of these known rhoptry proteins is likely due to limitations of the technique,
84 such as size cut-off or low amounts of protein.

85 Because several rhoptry proteins, such as the aforementioned ROP5, ROP16, ROP18 and
86 ROP38, contain kinase-like domains (Hajj et al., 2006a; Peixoto et al., 2010), another study
87 exploited a phylogenomic approach to characterize the *Toxoplasma* kinome, defining a 44-member
88 family of kinase-like rhoptry proteins based on sequence similarities, including all previously
89 reported kinase-like rhoptry proteins (Peixoto et al., 2010). To evaluate the accuracy of these
90 predictions, nine of the identified kinase-like rhoptry proteins were confirmed to localize to the
91 rhoptries, whereas two (ROP21 and ROP22) did not but were still annotated as rhoptry proteins.
92 Surprisingly, the overlap between these two studies is relatively small, with only 11 genes found in
93 common, which emphasizes the complementarity of different methodologies and the likelihood that
94 there are still rhoptry proteins yet to be identified.

95 Previous characterization of the cell cycle transcriptome of *Toxoplasma* covering 12 h post-
96 synchronization and nearly two tachyzoite replication cycles showed that the mRNA levels of
97 proteins secreted from the rhoptry organelles display a cyclical expression profile, reaching peak
98 levels in late S phase/early mitosis followed by a rapid and dramatic decline of these transcripts in
99 early G1, before peaking again in the next S phase (Behnke et al., 2010). Inner membrane complex
100 (IMC) mRNAs presented a very similar cyclical expression profile. Microneme mRNAs were offset
101 by 1–2 h from rhoptry mRNAs and defined a distinct temporal class. By contrast, dense granule
102 mRNAs largely were not regulated in the tachyzoite cell cycle.

103 Here we combined in silico and in vivo methods to identify novel rhoptry proteins likely to
104 be involved in parasite modulation of host cells and found two novel rhoptry bulb proteins and one
105 novel rhoptry neck protein.

106

107 **2. Materials and methods**

108 *2.1. Parasites and cell lines*

109 Parasites were maintained in vitro by serial passage on monolayers of human foreskin
110 fibroblasts (HFFs) at 37°C in 5% CO₂. HFFs were grown in DMEM supplemented with 10% FBS.

111 C57BL6/J mouse embryonic fibroblasts (MEFs) were a gift from A. Sinai (University of Kentucky
112 College of Medicine, Lexington, KY, USA) and were grown in HFF media supplemented with 10
113 mM HEPES, 1 mM sodium pyruvate and 1X non-essential amino acids.

114

115 *2.2. Identification of candidate genes*

116 For identification of new rhopty protein coding genes, cell cycle transcriptome data of the
117 *Toxoplasma gondii* during synchronized growth in human foreskin fibroblasts (HFFs) deposited at
118 Gene Expression Omnibus (GEO) with accession number **GSE19092** were used (Behnke et al.,
119 2010). The data were normalized using Robust Multi Array (RMA) algorithm using Affymetrix
120 Expression Console Software version 1.3.1, and all background values less than 6.5 were set to 6.5.
121 Genes with an expression value of less than 6.5 in 12 or more samples were removed from the
122 analysis. K-means clustering of each duplicated sample (time-point) was performed using the
123 default distance metric (Pearson correlation) and a maximum number of 50, 45, 40, 35, 30, 25, 20,
124 12 and 10 clusters in Multiple Array Viewer 4.6. The cluster with the largest number of previously
125 annotated rhopty proteins was chosen for further analysis. The candidate genes were chosen using
126 the following criteria: i) contained a predicted signal peptide, ii) had unannotated function and iii)
127 unknown subcellular localization.

128

129 *2.3. Gene tagging*

130 Genomic sequences were obtained from the ToxoDB database (Version 8.0, ToxoDB.org)
131 (Kissinger et al., 2003). For endogenous gene tagging (Huynh and Carruthers, 2009), primers were
132 designed to amplify 1 - 3 kb of the predicted 3' ends of genes. For heterologous expression of
133 ROP47, the coding regions of TGME49_261740, together with putative promoter (~1500 bp
134 upstream of ATG start codon) were amplified by PCR. All forward primers contained the 5'-
135 CACC-3' sequence required to perform directional TOPO cloning in pENTR/D-TOPO (Invitrogen,
136 USA) and all reverse primers contained the hemagglutinin (HA) tag sequence followed by a stop

137 codon (Table 1). HA-tagged sequences were cloned in vector pTKOatt by Gateway Recombination
138 Cloning Technology (Invitrogen, USA). For endogenous gene tagging, the resulting vectors were
139 linearized using a restriction enzyme with a unique restriction site within the cloned fragment.
140 Linearized vector was transfected into RH Δ *hxgprt* Δ *ku80* (a gift from V. Carruthers, University of
141 Michigan, Ann Arbor, MI, USA) parasites by electroporation. For heterologous expression of
142 ROP47, vector was not linearized and was transfected into RH Δ *hxgprt*. Electroporation was done in
143 a 2 mm cuvette (Bio-Rad Laboratories, USA) with 2 mM ATP (MP Biomedicals, USA) and 5 mM
144 glutathione (EMD, Germany) in a Gene Pulser Xcell (Bio-Rad Laboratories), with the following
145 settings: 25 μ FD, 1.25 kV, ∞ Ω . Stable integrants were selected in media with 50 μ g/ml of
146 mycophenolic acid (Axxora, USA) and 50 μ g/ml of xanthine (Alfa Aesar, USA) and cloned by
147 limiting dilution. The correct tagging of each gene was confirmed by PCR, using a primer upstream
148 of the plasmid integration site and a primer specific for the HA tag and/or by immunofluorescence
149 (IF) analysis.

150

151 2.4. IF analysis

152 Parasites were allowed to invade cells on coverslips and incubated for 16 – 24 h. The cells
153 were then fixed with 3% (vol/vol) formaldehyde in PBS for 20 min at room temperature and
154 blocked in PBS with 3% (wt/vol) BSA and 5% (vol/vol) goat serum. Coverslips were incubated
155 with primary antibody for 1 h at room temperature or overnight at 4°C, and fluorescent secondary
156 antibodies and Hoechst dye were used for antigen and DNA visualization, respectively. Coverslips
157 were mounted on a glass slide with Vectashield (Vector Laboratories, USA), and photographs were
158 taken using NIS-Elements software (Nikon, Japan) and a digital camera (CoolSNAP EZ; Roper
159 Industries, USA) connected to an inverted fluorescence microscope (model eclipse Ti-S; Nikon). To
160 determine seroconversion, peripheral blood serum was used as a primary antibody. For co-
161 localization experiments, GFP-expressing parasites were fixed with methanol or 3% (vol/vol)
162 formaldehyde. For moving junction staining, this standard IF protocol was modified slightly.

163 Parasites were added to HFFs on coverslips, spun down to bring them into contact with host cells,
164 and allowed to attach to and invade host cells for 5 min at 37°C. Unattached parasites were washed
165 off with PBS, and cells were fixed with 3% (vol/vol) formaldehyde in PBS for 20 min at room
166 temperature, blocked in PBS with 5% (vol/vol) FBS and 5% (vol/vol) normal goat serum for 1 – 2 h
167 at room temperature, and permeabilized by incubation in PBS with 0.2% (wt/vol) saponin at 37°C
168 for 20 min.

169

170 2.5. Antibodies

171 Recombinant ROP47 peptide antigen was generated by PCR amplification of sequences
172 immediately downstream of the predicted ROP47 signal peptide and immediately upstream of the
173 ROP47 stop codon from *PruΔhxgprt* genomic DNA. The amplicon was cloned in frame into
174 pET102/D-TOPO (Invitrogen), creating a Thioredoxin-ROP47-V5-His6 construct. This protein was
175 expressed in BL21 Star (Invitrogen) cells and purified on a Ni-NTA column (Invitrogen) eluted
176 with 250 mM imidazole and dialyzed under native conditions according to the manufacturer's
177 directions. Rabbit polyclonal antibodies (Covance, USA) were raised against the purified peptide
178 antigen. Antibodies were affinity purified from the ROP47 antiserum with Affi-Gel 10 resin (Bio-
179 Rad) covalently coupled to the antigen peptide and eluted with 100 mM Glycine, pH 2.5, then
180 immediately neutralized with 1 M Tris pH 8.0. The eluate was subsequently incubated with Affi-
181 Gel 10 resin covalently coupled to a Thioredoxin-V5-His6 peptide that was expressed and purified
182 as above to remove antibodies in the serum reacting against thioredoxin and the epitope tags. The
183 flow-through was collected, tested for specificity and used for immunogenic assays.

184 Antibodies against HA (Roche), *Toxoplasma* surface antigen (SAG)-1 (DG52, (Burg et al.,
185 1988)), *Toxoplasma* rhoptry protein ROP1 (Tg49; (Ossorio et al., 1992)), *Toxoplasma* dense
186 granule protein GRA7 (Dunn et al., 2008), *Toxoplasma* rhoptry neck protein RON4 (generously
187 provided by P. Bradley, University of California, Los Angeles, CA, USA), *Toxoplasma* inner
188 membrane complex proteins IMC1 and MLP1 (generously provided by M.J. Gubbels, Boston

189 College, Boston, MA, USA) and mouse TGTP (A-20; Santa Cruz Biotechnology, USA) were used
190 in the IF assay. IF secondary antibodies were coupled with Alexa Fluor 488 or Alexa Fluor 594
191 (Invitrogen).

192

193 2.6. *Western blot*

194 Parasites were syringe lysed from infected HFFs with lysis buffer, boiled for 5 min and
195 subjected to 10% SDS-PAGE. Proteins were transferred to a polyvinylidene difluoride membrane,
196 which was blocked in PBS/0.1% Tween-20/5% non-fat dry milk and incubated with primary and
197 secondary antibodies. The blot was incubated with a luminal-based substrate (Immun-Star
198 WesternC; Bio-Rad Laboratories) and chemiluminescence was detected using a charge-coupled
199 device camera (Chemidoc XRS; Bio-Rad Laboratories). The bands were visualized using Quantity
200 One 1-D analysis software.

201

202 2.7. *Generation of gene knockouts (KOs)*

203 The 5' and 3' flanking regions of the genes to be knocked out were cloned in pTKO2
204 (Rosowski et al., 2011) around the hypoxanthine-xanthine-guanine ribosyl transferase (HXGPRT)
205 selectable marker using Multisite Gateway Pro 3-Fragment Recombination (Invitrogen). Flanking
206 regions (5' and 3') of TGME49_201860, TGME49_218270 and TGME49_261740 were cloned
207 from type II genomic DNA. Primers contained *att* recombination sites (denoted in primer sequence
208 with italics, Table 1) and amplified ≈ 2 kb upstream of the start codon and downstream of the stop
209 codon. These flanking regions were then cloned around the HXGPRT selectable marker flanked by
210 5' and 3' untranslated region (UTRs) from dihydrofolate reductase (DHFR), as previously described
211 (Rosowski et al., 2011). Before transfection, the KO vector was linearized. Pru Δ *hxgp*rt Δ *ku80* (a gift
212 from D. Bzik, Dartmouth Medical School, Lebanon, NH, USA) parasites were transfected with the
213 KO construct by electroporation, and stable integrants were selected and cloned by limiting
214 dilution, as described in Section 2.3. PCR with a forward primer upstream of the 5' flanking region

215 (P1) and a reverse primer within the HXGPRT cassette (P2) confirmed the disruption in the desired
216 loci (Supplementary Fig. S1A). Additionally, PCR was performed to confirm the inability to
217 amplify the target genes (P3 and P4) (Supplementary Fig S1A).

218

219 *2.8. Plaque assays*

220 For the plaque assays, 100–500 parasites per well were added to monolayers of MEFs
221 seeded the day before and either previously stimulated with 1000 U/mL of mouse IFN γ or left
222 unstimulated for 24 h before infection in a 24 well plate in MEF media. Infections were then
223 incubated for 5 days at 37°C and the number of plaques was counted using a microscope.

224

225 *2.9. Animal infections*

226 Six to 10 week old female C57BL/6J mice (The Jackson Laboratory, USA) were used in all
227 experiments. For i.p. infection, tachyzoites were grown in vitro and extracted from host cells by
228 passage through a 30-gauge needle, washed twice in PBS and quantified with a hemocytometer.
229 Parasites were diluted in PBS and mice were inoculated i.p. with 500 tachyzoites of each strain (in
230 100 μ L) using a 28-gauge needle. For oral infection, brain homogenate of chronically infected mice
231 was stained with dolichos biflorus-FITC (Vector Laboratories) and cysts were enumerated by
232 microscopy. The mice were orally gavaged with 1000 cysts. All of the animals were monitored
233 daily and weighed three times per week. Peripheral blood serum was collected on days 7 and 30 of
234 the experiment and the levels of IFN γ were determined using commercially available ELISA kits,
235 according to the manufacturer's protocol (eBioscience, USA). The Massachusetts Institute for
236 Technology, USA, Committee on Animal Care approved all protocols. All mice were maintained
237 under specific pathogen-free conditions, in accordance with institutional and federal regulations.

238

239 **3. Results and Discussion**

240 *3.1. Identification of novel rhoptry proteins*

241 Previous characterization of the *Toxoplasma* cell cycle transcriptome showed that the
242 mRNA levels of rhoptry encoding genes display the same cyclical expression profile (Behnke et al.,
243 2010). We hypothesized that unidentified rhoptry encoding genes would show a similar expression
244 profile. Genes were identified that showed the same expression pattern as established rhoptry
245 encoding genes by performing K-means clustering of 13 duplicate samples spanning 12 h post-
246 thymidine release into the tachyzoite cell cycle. The cluster with the largest number of previously
247 annotated rhoptry protein encoding genes (67 out of 75 annotated rhoptry encoding genes were
248 expressed above background and were present on the *Toxoplasma* array; Supplementary Table S1)
249 was chosen for further analysis (Table 2, Supplementary Fig. S2). This cluster contained 190 unique
250 genes, of which 45 encoded proteins annotated as rhoptries (out of 67 possible genes) and four
251 encoded IMC proteins. Thirty-one out of these 45 have been previously confirmed to localize to the
252 rhoptry organelles.

253 The *Toxoplasma* genome is predicted to encode 8127 genes, 1920 of which have a signal
254 peptide (23.4%). Our analysis imposed no explicit selection for genes coding for proteins with
255 signal peptide sequences; however, the resulting list is enriched (65 genes with a predicted signal
256 peptide out of 190, $P = 0.0003$, Hypergeometric distribution) in signal-peptide-containing proteins
257 (34%).

258 Interestingly, 22 out of the 75 tachyzoite rhoptry proteins encoding genes never clustered
259 with the remainder of the rhoptries (Supplementary Table S1). Two of these genes (RON2L1 and
260 BRP1 (Schwarz et al., 2005; Fritz et al., 2012)) encode bradyzoite- or sporozoite-specific rhoptry
261 proteins and six encode confirmed tachyzoite rhoptries (ROP9, ROP38, ROP39, TgARO, Toxolysin
262 1 and Subtilisin 2). According to the cell cycle expression data (Behnke et al., 2010), ROP9, ROP38
263 and ROP39 do not display an obvious cyclical expression profile. In addition, ROP9 was shown to
264 be secreted with micronemal proteins, in a calcium-dependent manner (Kawase et al., 2007). The
265 cellular localization of the proteins encoded by the other 14 genes annotated as rhoptry protein

266 genes (Supplementary Table S1) was never confirmed by IF and it is therefore possible that these
267 are not localized to the rhoptries.

268 These results underscore the relevance of our approach and strongly suggest that our rhoptry
269 protein cluster might contain many of the remainder of the unidentified rhoptry protein-encoding
270 genes of tachyzoites. To identify new putative rhoptry protein encoding genes that could modulate
271 host cell functions, we chose eight genes that encoded a protein with a predicted signal peptide,
272 unannotated function and unknown subcellular localization from within our rhoptry cluster. The
273 eight candidate genes are described in Table 2, highlighted in grey, and all display the cyclic
274 expression profile described above (Fig. 1).

275 To determine the localization of the candidate gene products within the parasite, we
276 genetically engineered *Toxoplasma* strains expressing an HA-tagged version of the candidate genes
277 at their endogenous loci. We were unable to endogenously tag TGME49_261740 and therefore
278 determined its localization by heterologous expression of a C-terminal HA-tagged copy of the
279 candidate gene, including at least 1500 bp of the putative endogenous promoter. The correct tagging
280 of each endogenously tagged gene was confirmed by PCR (Supplementary Fig. S3). IF analysis was
281 performed on HFFs infected with each of the HA-tagged parasites. The staining pattern of four out
282 of eight candidate gene products was unlike that of the secretory organelles and appeared to label
283 the whole parasite (Supplementary Fig. S4). The genes whose labeling appeared consistent with that
284 of the secretory organelles were TGME49_201860, TGME49_218270, TGME49_232020 and
285 TGME49_261740 (Fig. 2A). Some characteristics of the proteins encoded by these genes are
286 described in Table 3.

287 The product of TGME49_201860-HA appears to have a punctate distribution, present at
288 both the posterior and the apical end of the parasite (Fig. 2A). TGME49_218270-HA seems to label
289 the rhoptry bulb. We were unable to detect these two proteins using an anti-HA western blot, and
290 therefore do not know whether they undergo post-transcriptional processing.

291 TGME49_232020-HA localizes to the apical pole of the parasite, a labeling consistent with
292 the rhoptry neck (Fig. 2A). This gene is predicted to encode a protein with no signal peptide in type
293 II parasites. However, as first exon prediction is notoriously difficult, in type I and type III
294 parasites, this protein is predicted have a signal peptide. Moreover, TGME49_232020 encodes a
295 protein with a predicted molecular weight of 135 kDa. However, western blot analysis of the
296 TGME49_232020-HA strain detected a band of approximately 40 kDa, instead of the predicted 135
297 kDa (Supplementary Fig. S5). Indeed, several rhoptry proteins exhibit N-terminal processing and
298 that is also probably the case for TGME49_232020. The rhoptry subtilisin TgSUB2 recognizes and
299 cleaves after the consensus sequence SΦXE (Miller et al., 2003). There is a putative SUB2
300 cleavage site (SPQE) between amino acids 968 and 971 of TGME49_232020. Cleavage at this site
301 would generate a ~32 kDa tagged product, which could be consistent with the band observed.
302 Longer exposures did not reveal a 135 kDa pro-protein, suggesting that the half-life of the pro-
303 protein is very short.

304 TGME49_261740-HA appears to label the entire rhoptry organelle (Fig. 2A). Interestingly,
305 TGME49_261740 is in the top five of the most highly expressed genes across the *Toxoplasma* cell
306 cycle (Behnke et al., 2010) and is one of the most polymorphic *Toxoplasma* genes (Minot et al.,
307 2012). Anti-HA western blotting of this protein detected a band of approximately 15 kDa,
308 consistent with the predicted size of 14 kDa (Supplementary Fig. S1B).

309 These four proteins are conserved between *Toxoplasma* and *Neospora* (Table 3), but protein
310 BLAST analysis did not reveal the presence of close (30% or more protein identity) homologues in
311 other apicomplexans. Additionally, no known conserved domains were detected, giving no
312 indication about the potential function of these proteins.

313

314 3.2. TGME49_218270, TGME49_232020 and TGME49_261740 encode new rhoptry proteins

315 To further confirm the localization of TGME49_201860, TGME49_218270,
316 TGME49_232020 and TGME49_261740 within the parasite, we performed co-staining of each

317 protein with a rhoptry bulb marker, ROP1, on intracellular parasites (Fig. 2A). TGME49_218270-
318 HA shows a perfect co-localization with ROP1.

319 TGME49_201860-HA does not overlap with this rhoptry marker. To further investigate the
320 cellular localization of this protein, we co-stained the TGME49_201860-HA expressing parasites
321 with dense granule (GRA7) and inner membrane complex (IMC1 and MIP1-like protein-1 (MLP1))
322 markers (Fig. 2B). TGME49_201860-HA appears to partially overlap with GRA7. In addition,
323 while it does not co-localize with IMC1, it co-localizes with MLP1, a protein that is present at the
324 apical cap and basal complex of mature as well as budding daughter parasites (MJ Gubbels,
325 personal communication).

326 TGME49_232020-HA showed labeling of the apical pole anterior to the rhoptry bulb
327 protein ROP1, as well as co-localization with RON4 staining (Fig. 2C), confirming rhoptry neck
328 localization in intracellular parasites. Interestingly, TGME49_232020-HA does not localize to the
329 moving junction in invading parasites (Fig. 2C). It was recently shown that RON9, RON10 and
330 RON11, in contrast to the other RONS described to date, do not relocalize from the rhoptry neck to
331 the moving junction during host invasion (Lamarque et al., 2012; Beck et al., 2013).

332 TGME49_232020 localization also seems to be independent of the moving junction. Similar to
333 TGME49_232020, RON10 has a predicted signal peptide in its N-terminus but no known domains
334 or motifs have been identified. In contrast, RON9 harbors several protein-protein interaction
335 domains. The disruption of either RON9 or RON10 led to the retention of either protein in the
336 endoplasmic reticulum (ER), suggesting an interaction between RON9 and RON10 during their
337 trafficking through the secretory pathway on the way to the rhoptries. Whether TGME49_232020
338 can interact with the RON9/RON10 complex or any other rhoptry neck protein in a similar fashion
339 remains to be determined. TGME49_261740 co-localizes with ROP1 (Fig. 2B) and is also found in
340 the nucleus of infected host cells (Fig. 2D). NLS Mapper (Kosugi et al., 2009) predicts that
341 TGME49_261740 (ROP47) encodes a bipartite NLS with a score of 3.4. Moreover, this protein is
342 predicted to have a size of 15 kDa and is probably able to diffuse through nuclear pores.

343 Altogether, these results suggest that TGME49_261740 and TGME49_218270 encode new
344 rhoptry bulb proteins, hereafter referred to as ROP47 and ROP48, respectively. TGME49_232020
345 codes for a new rhoptry neck protein, from now on referred to as RON12.

346

347 3.3. *TGME49_201860 and ROP48 are not implicated in evasion of the IFN γ response*

348 To investigate the role of TGME49_201860, ROP47 and ROP48 in parasite biology, we
349 removed these genes in the *Pru Δ hxgp Δ ku80* strain using double homologous recombination. PCR
350 confirmed both the absence of the target gene coding sequences and the insertion of the *hxgp Δ* gene
351 in the parasite genome. The deletion of ROP47 was confirmed by western blot (Supplementary Fig.
352 S1B). Our attempts to generate a KO of RON12 in the *Pru Δ hxgp Δ ku80* strain have to date been
353 unsuccessful.

354 To determine a potential growth phenotype, monolayers of MEF were infected with
355 *Pru Δ hxgp Δ ku80*, *Δ tgme49_201860* or *Δ rop48* parasites, the parasites were allowed to grow for 5
356 days, and then the areas of the plaques formed on the monolayers were quantified (Fig. 3A).
357 *Pru Δ hxgp Δ ku80*, *Δ tgme49_201860* and *Δ rop48* strains did not form significantly different sized
358 plaques. These data indicate that deletion of TGME49_201860 or ROP48 has no major influence on
359 in vitro *Toxoplasma* growth.

360 It is well established that IFN γ is the main mediator of resistance against *Toxoplasma*
361 (Suzuki et al., 1988). An important class of downstream effectors of this immune activation is the
362 IFN γ -inducible immunity-related GTPases (IRGs), which belong to the dynamin family of GTPases
363 and can cooperatively oligomerize to vesiculate membranes. The IRGs are able to disrupt the PVM
364 and kill the parasite (Butcher et al., 2005). Recently, two rhoptry proteins were shown to mediate
365 *Toxoplasma* evasion of the IFN γ -induced IRGs in murine cells (Fentress et al., 2010; Steinfeldt et
366 al., 2010; Niedelman et al., 2012). To study a potential role of ROP48 and TGME49_201860 in
367 *Toxoplasma* resistance to IFN γ and the IRGs, we measured the percentage of vacuoles coated with
368 Irgb6 by IF in IFN γ -stimulated MEFs infected with *Pru Δ hxgp Δ ku80*, *Δ tgme49_201860* or *Δ rop48*.

369 The percentage of coated vacuoles was approximately 50% in both *Δtgme49_201860* and *Δrop48*
370 and was not significantly different from that of the parental strain, *PruΔhxgprrtΔku80* (Fig. 3B).
371 Although it is generally assumed that once the PVM is coated, it will eventually lead to killing of
372 the parasite inside, it has also been shown that *Toxoplasma* can escape a coated vacuole and invade
373 a new cell (Zhao et al., 2009). Therefore, to measure killing of *Toxoplasma*, a plaque loss assay was
374 performed, as described in Niedelman et al., (2012). Briefly, 100 parasites were seeded on a
375 monolayer of MEF, either previously stimulated for 24 h with IFN γ or left untreated, and the
376 number of plaques that form after 5 days of growth was determined. The three strains had an
377 average of ~30% plaque loss when comparing plaques formed on IFN γ -stimulated MEFs with
378 unstimulated MEFs. This percentage of plaque loss was lower than the percentage of vacuoles
379 coated with Irgb6, suggesting that some coated vacuoles can escape destruction by the IRGs.
380 Overall, the results suggest that TGME49_201860 and ROP48 are not implicated in evading the
381 IFN γ response in MEFs. Indeed, ROP5 and ROP18 were recently reported to mediate *Toxoplasma*
382 evasion of the murine IFN γ response (Niedelman et al., 2012). These two proteins seem to
383 determine the majority of strain differences in mouse IRG evasion, even for non-clonal strains for
384 which virulence determinants have not been studied. However, neither ROP18 nor ROP5 markedly
385 affect survival in IFN γ -activated human cells and it is reasonable to speculate that one or more
386 still unidentified, secreted, potentially a rhoptry, protein could be involved in escaping IFN γ -
387 mediated killing in other host cell types. Whether this (these) protein(s) can be found within the
388 rhoptry cluster will be the object of future studies.

389

390 3.4. *TGME49_201860*, *ROP47* and *ROP48* are not implicated in *Toxoplasma* virulence in mice

391 To examine the role of *TGME49_201860*, *ROP47* and *ROP48* on parasite virulence, we
392 infected C57BL/6 mice by i.p. injection with 500 tachyzoites of *PruΔhxgprrtΔku80*,
393 *Δtgme49_201860*, *Δrop47*, *Δrop48* or a heterologous control strain (Het) and assessed mouse

394 morbidity (through monitoring of weight loss) and survival during the initial phase of infection
395 (days 0-32) (Fig. 4A, B). Weight loss started at day 4 p.i. The mice reached their lowest weight
396 between days 14 and 18 (~80% of their initial body weight) and did not regain their original weight.
397 The survival of C57BL/6 mice infected with either strain did not show significant differences
398 compared with Pru Δ hxgp Δ ku80 infected mice ($P = 0.9060$, Log-rank Mantel-Cox test).
399 Seroconversion was examined at day 30 p.i. and all the animals tested seropositive (data not
400 shown). The level of IFN γ in peripheral blood serum was measured at day 7 and day 30 p.i. (Fig.
401 4D). IFN γ levels were higher at day 7 than day 30 and similar for all the strains. Prior to infection,
402 animals did not display detectable levels of IFN γ . At day 35 p.i., the surviving mice were sacrificed
403 and the number of brain cysts per mouse was determined (Fig. 4C). The numbers of cysts generated
404 by the Δ tgme49_201860, Δ rop47 and Δ rop48 parasites were not significantly different from that of
405 their parental strain. The number of brain cysts observed in the mice injected i.p. with the
406 heterologous control strain was significantly lower (~1.6 fold, $P = 0.0011$, Student's t -test) than that
407 observed in mice infected with Pru Δ hxgp Δ ku80 parasites. Accordingly, it was previously reported
408 that Pru Δ ku80::hxgp Δ exhibited lower cyst burdens than Pru Δ hxgp Δ ku80 (Fox et al., 2011), which
409 suggests the presence of HXGPRT might affect generation or viability of brain cysts. Additionally,
410 we infected C57BL/6 mice by oral gavage with 1000 brain cysts of the same strains. We found no
411 difference in mouse morbidity and survival (data not shown).

412 Overall, the results suggest that TGME49_201860, ROP47 and ROP48 are not implicated in
413 *Toxoplasma* virulence in mice. Interestingly, relatively few rhoptry proteins seem to have been
414 investigated on their ability to affect virulence in the mouse model. Of the 10 proteins that were
415 reported, ROP5, ROP16, ROP13, ROP18, RON8, RON9/10, TLN1, BRP1 and PP2C-hn (Saeij et
416 al., 2006; Gilbert et al., 2007; Turetzky et al., 2010; Hajagos et al., 2011; Straub et al., 2011;
417 Lamarque et al., 2012; Niedelman et al., 2012;), only four (RON8, ROP5, ROP16, ROP18) have an
418 effect on mouse survival. However, mouse virulence is often tested in tachyzoites, the asexually
419 reproducing form of the parasite, leaving out events that are restricted to the sexual life cycle of the

420 parasite. Alternatively, the mouse model used may not be the optimal setting to reveal an essential
421 role for such proteins in infection. For instance, a role in virulence could only be apparent in other
422 intermediate hosts and/or when cysts are ingested naturally. Elucidation of this question will be
423 extremely challenging due to the remarkable host range of *Toxoplasma*.

424

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434

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- 624
- 625

626 **Figure legends**

627

628 **Fig. 1.** Identification of new *Toxoplasma* rhopty proteins. The gene expression profile of the entire
629 genome of *Toxoplasma* was compared with the cyclical gene expression profile of rhopty genes
630 throughout the cell cycle by performing clustering of duplicate samples spanning 12 h post-
631 synchronization (Supplementary Fig. S2). Shown are eight genes that displayed an expression
632 profile signature similar to the rhopty pattern and that were chosen for further analysis. The
633 expression profile of ROP1 is also shown.

634

635 **Fig. 2.** Cellular localization of C-terminally tagged genes. Human foreskin fibroblasts (HFFs) were
636 infected with RH expressing hemagglutinin (HA)-tagged TGME49_201860, TGME49_218270
637 (*rop48*), TGME49_232020 (*ron12*) or TGME49_261740 (*rop47*), fixed and stained with (A) α -HA
638 and rhopty protein marker α -ROP1, (B) α -HA and dense granule protein marker α -GRA7, α -HA
639 and inner membrane complex marker α -IMC1, α -HA and inner membrane complex marker α -
640 MLP1. (C) α -HA and rhopty neck/moving junction marker, α -RON4 and (D) α -HA, α -ROP47
641 and Hoechst dye. (A-C) Scale bars represent 5 μ m. (D) Scale bars represent 20 μ m.

642

643 **Fig. 3.** Deletion of TGME49_201860 and ROP48 does not affect plaque loss in response to IFN γ .
644 (A) Monolayers of mouse embryonic fibroblasts (MEF) were infected with Pru Δ *hxgprt* Δ *ku80*,
645 Δ TGME49_201860 or Δ *rop48* parasites. The plaque area was quantified after 5 days. Mean \pm S.D.;
646 $n \geq 30$ plaques. (B) Quantification of the localization of IFN γ -inducible immunity-related GTPase
647 B6 (*Irgb6*) on the parasite containing vacuole and of the percentage of plaque loss after 5 days on
648 MEF stimulated with IFN γ compared with unstimulated MEF. Mean \pm S.D.; $n = 4$ experiments.

649 **Fig. 4.** Deletion of the genes encoding TGME49_201860, ROP47 and ROP48 does not affect
650 *Toxoplasma gondii* virulence in mice. C57BL/6 mice were infected with 500 tachyzoites and
651 weight and survival of mice was monitored. (A) Average percentage change in weight over time for
652 mice i.p. infected with the indicated strains; $n \geq 4$ for each strain, mean + S.D. (B) Mouse survival
653 after i.p. infection with indicated strains; $n \geq 4$ for each strain. (C) IFN γ cytokine levels in
654 peripheral blood serum of surviving animals ($n \geq 2$ for each strain) was determined by ELISA at
655 days 7 (D7) and 30 (D30) p.i., mean \pm S.D. Pru - *Pru* Δ *hxgprrt* Δ *ku80*. Het – Heterologous control.
656 (D) Average number of brain cysts at day 35 following i.p. infection with the indicated strains; $n \geq$
657 2 for each strain; mean \pm S. D. Pru - *Pru* Δ *hxgprrt* Δ *ku80*. Het – Heterologous control. * $P = 0.0011$,
658 Student's *t*-test.

659

660 **Supplementary Figure legends**

661

662 **Supplementary Fig. S1.** Generation of gene knockouts (KO) in *Toxoplasma gondii*. (A) Scheme
663 depicting the strategy used to obtain the KO strains. The 5' and 3' flanking regions (FR) of our
664 genes of interest were cloned on both sides of hypoxanthine-xanthine-guanine ribosyl transferase
665 (HXGPRT) selection marker. The vector was linearized prior to transfection of $\Delta ku80$ parasites.
666 Following a double homologous recombination event, the genes of interest were replaced by
667 HXGPRT. The arrows represent the primers P1, P2, P3 and P4 used to verify the gene replacement.
668 (B) To confirm the disruption of TGME49_201860 and ROP48, PCR was performed with a
669 forward primer upstream of the 5' FR (P1) and a reverse primer within the HXGPRT cassette (P2).
670 A second PCR was executed to confirm the inability to amplify the target genes (P3 and P4). The
671 deletion of ROP47 was confirmed by western blot using an anti-ROP47 antibody on human
672 foreskin fibroblasts (HFF) lysates that were either uninfected, infected with the parental strain or
673 with the $\Delta rop47$ strain. Top band, indicated by *, is non-specific.

674

675 **Supplementary Fig. S2.** Identification of new *Toxoplasma* rhoptry protein-encoding genes. The
676 gene expression profiles of all expressed *Toxoplasma* genes were compared with the cyclical gene
677 expression profile of rhoptry genes throughout the cell cycle by performing K-means clustering of
678 13 duplicate samples spanning 12 h post-synchronization. The cluster with the highest number of
679 known rhoptry protein-encoding genes is shown. Genes analyzed in detail in this report are
680 highlighted. The complete list of genes is described in Table 2.

681

682 **Supplementary Fig. S3.** Validation of endogenously tagged parasites. (A) Scheme depicting the
683 strategy used to obtain the endogenously tagged strains. The hemagglutinin (HA) tag was added to
684 the 3' regions of genes of interest by PCR. The resulting amplicon was cloned upstream of the 3'
685 untranslated region (UTR) of GRA2 in a vector harboring the hypoxanthine-xanthine-guanine

686 ribosyl transferase (HXGPRT) gene. The vector was linearized prior to transfection of *Δku80*
687 parasites. By reciprocal recombination, the region of homology in the construct recombined with
688 the homologous sequences on the chromosomal gene, introducing the HA tag to the 3' end of the
689 endogenous gene. The arrows represent the primers forward (F) and reverse (R) used to verify the
690 gene replacement. GOI – gene of interest. (B) Endogenous gene tagging of seven candidate genes
691 was confirmed by PCR, using a primer upstream of the plasmid integration site and a primer
692 specific for the HA tag.

693

694 **Supplementary Fig. S4.** Cellular localization of endogenously tagged genes in *Toxoplasma*.

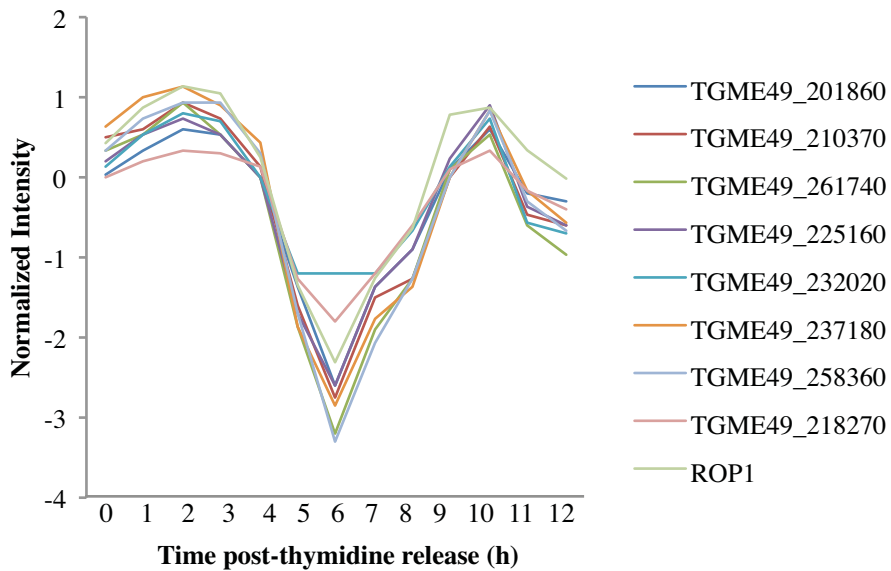
695 Representative picture of human foreskin fibroblasts (HFFs) infected with RH $\Delta ku80$ expressing
696 GFP (green) and endogenously tagged TGME49_210370, TGME49_225160, TGME49_237180 or
697 TGME49_258360. Endogenously tagged TGME49_210370 is shown. Cells were fixed after 24 h
698 and stained with α -hemagglutinin (α -HA, red). Scale bar represents 5 μ m.

699

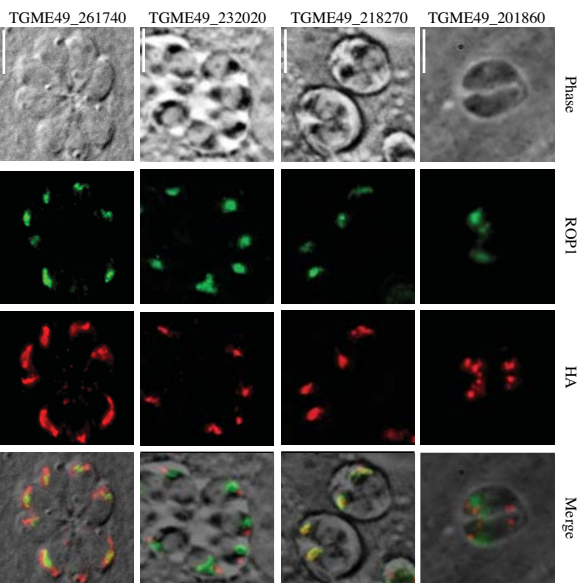
700 **Supplementary Fig. S5.** The protein encoded by *Toxoplasma* gene TGME49_232020 is most
701 likely processed. Western blot of RH $\Delta hxgp rt \Delta ku80$ and RH $\Delta ku80$ with hemagglutinin (HA)-tagged
702 TGME49_232020 lysates with α -HA shows migration of TGME49_232020-HA at \sim 40 kDa,
703 whereas its predicted size is \sim 135 kDa. Blotting for the major surface antigen of *Toxoplasma*,
704 SAG-1, was used as a loading control.

705

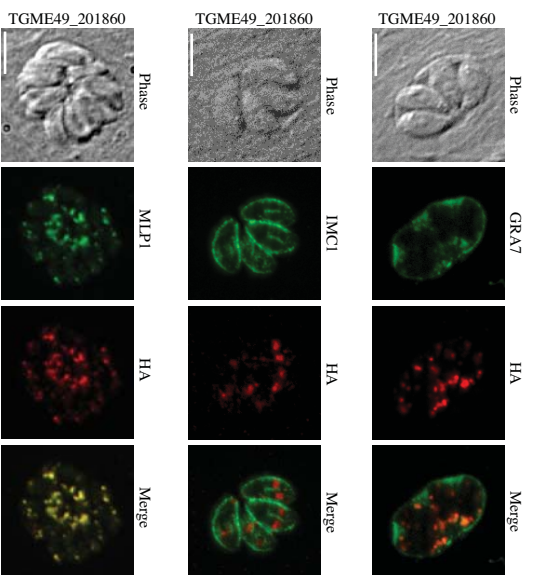
706



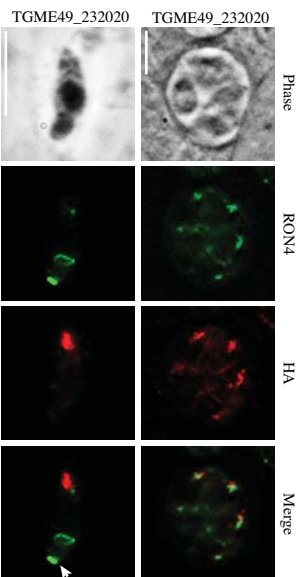
A



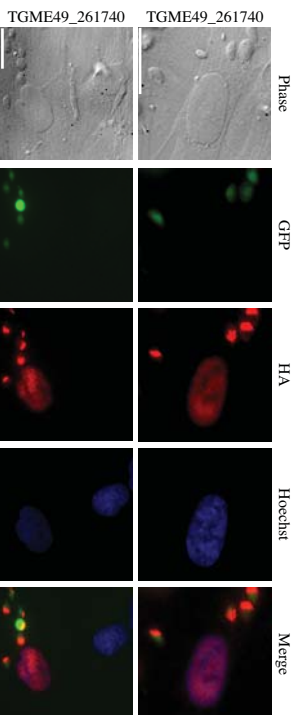
B



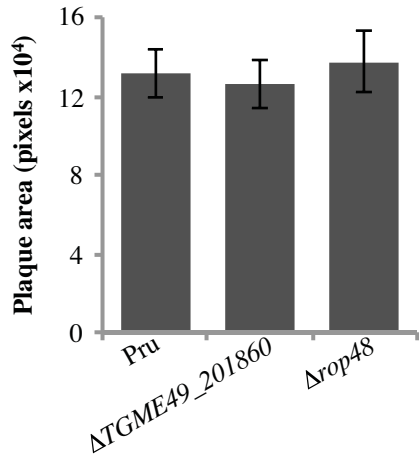
C



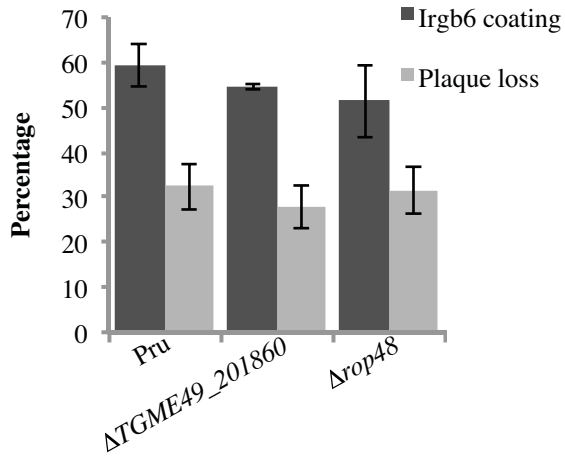
D

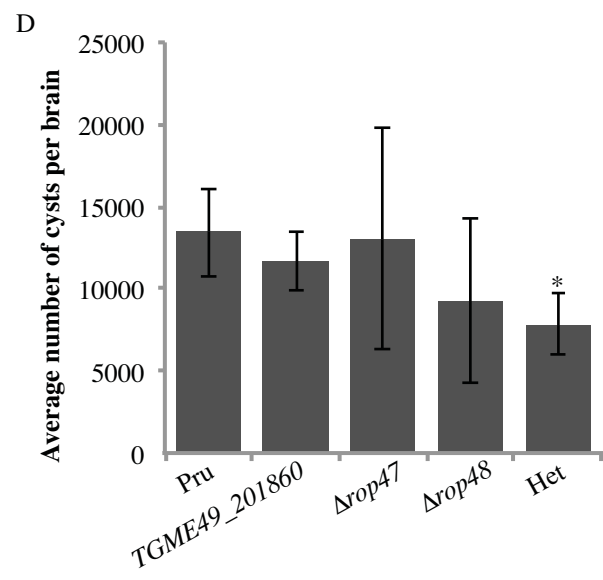
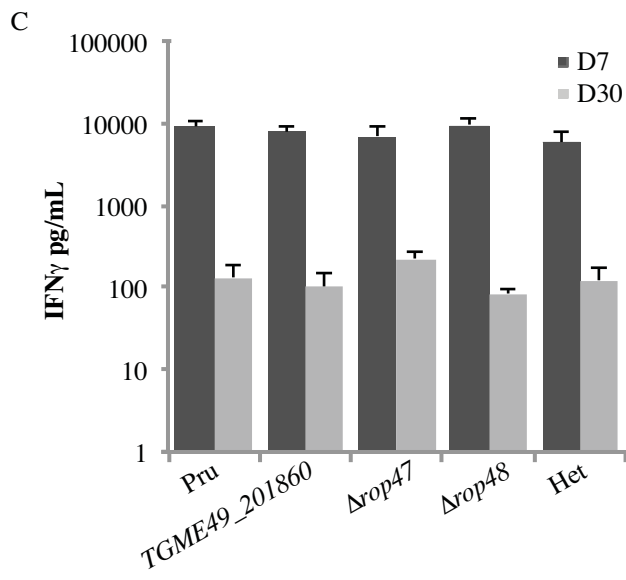
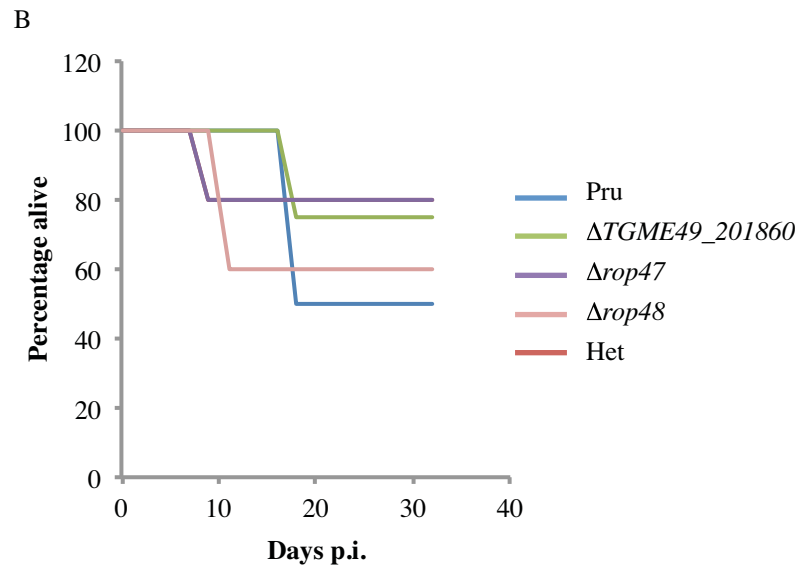
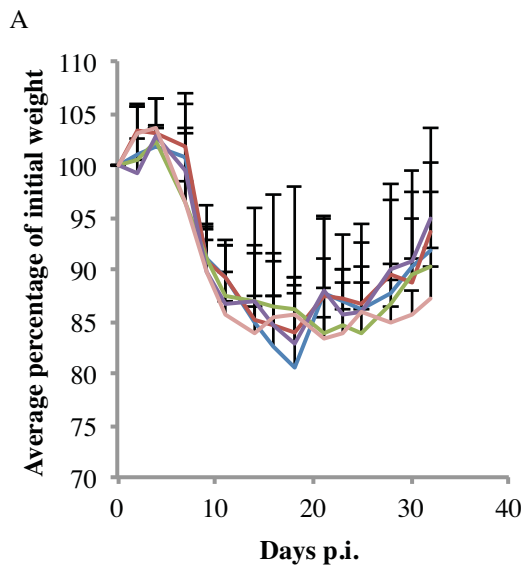


A

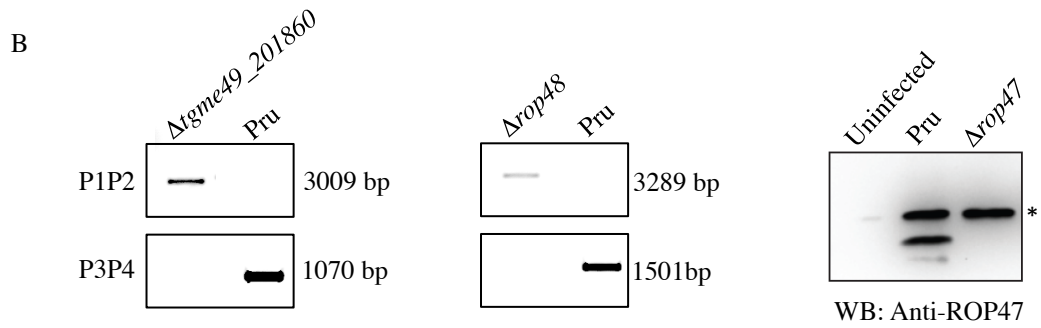
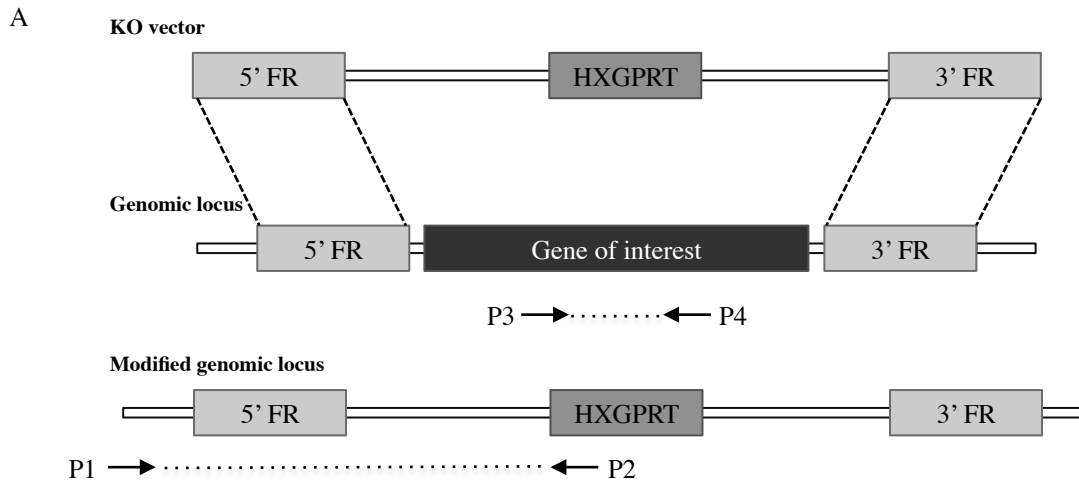


B





Supplementary Fig. S1

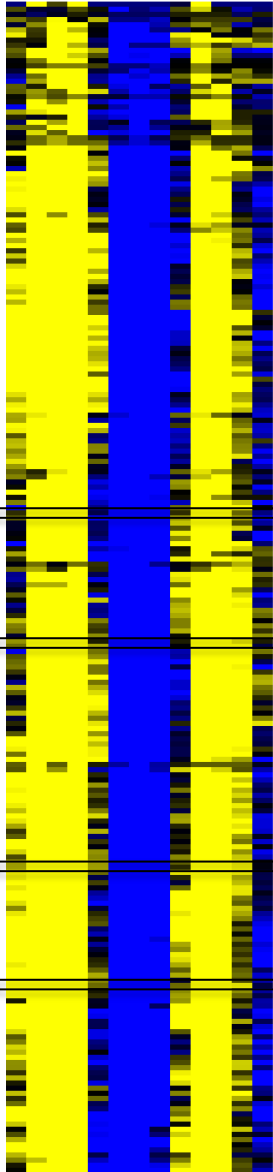


Supplementary Fig. S2



Time post-thymidine release (h)

0 1 2 3 4 5 6 7 8 9 10 11 12



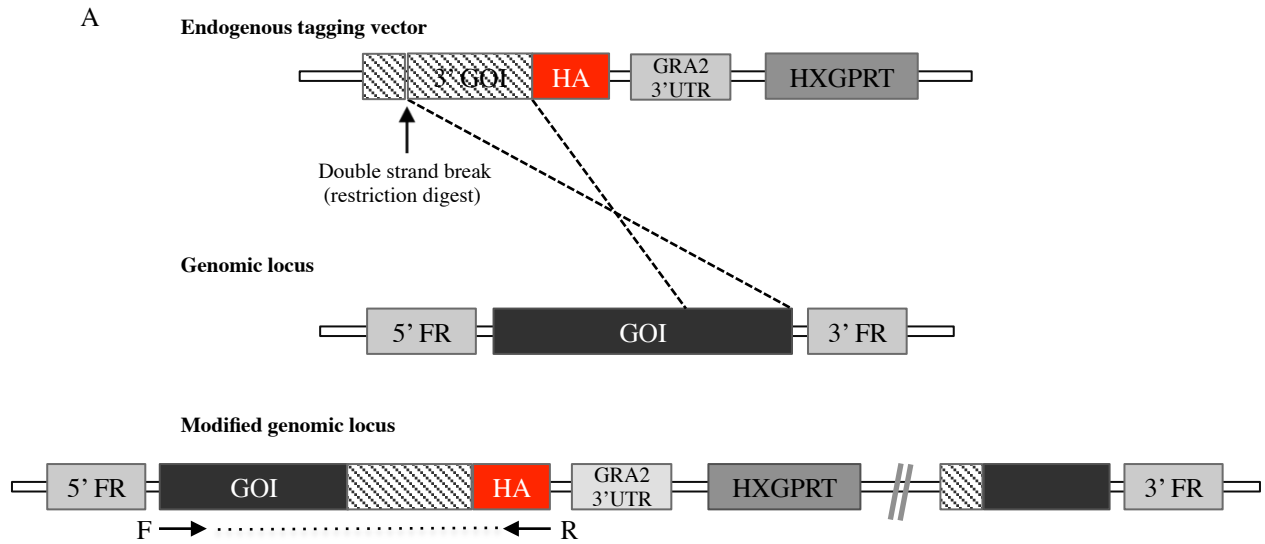
TGME49_232020

TGME49_201860

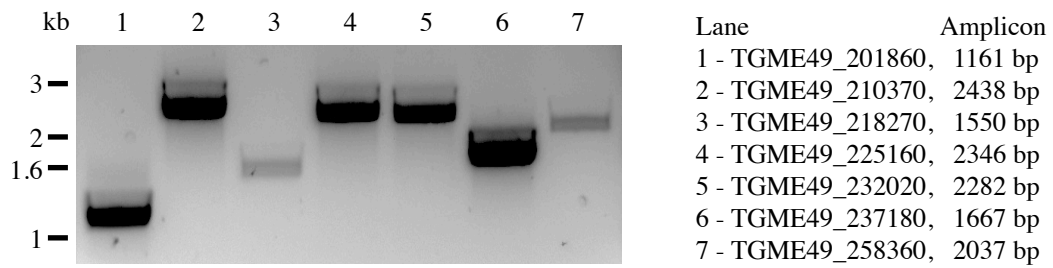
TGME49_261740

TGME49_218270

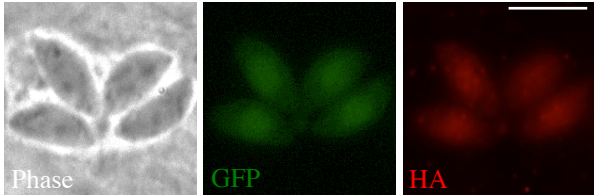
Supplementary Fig. S3



B



Supplementary Fig. S4



Supplementary Fig. S5

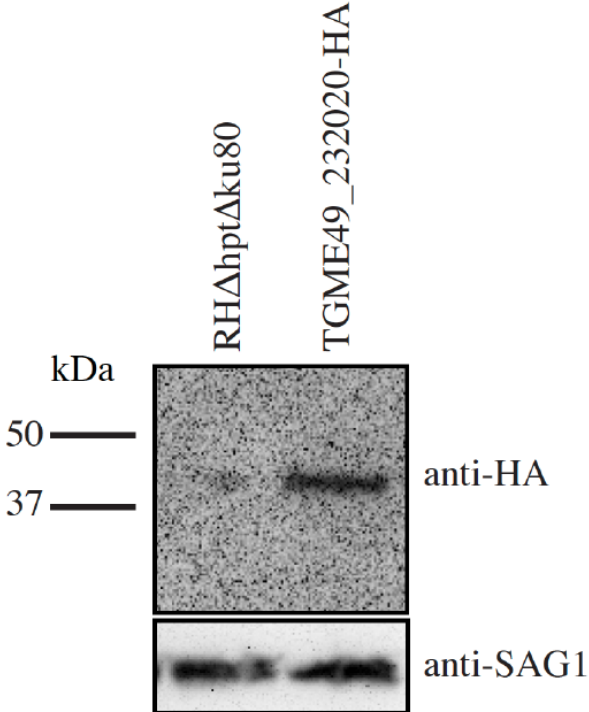


Table 1. Primers used in this study.

Gene ID	Forward primer	Reverse primer		
Primers used for gene tagging (5'-3')				
TGME49_201860	CACCTCGCGACCACCTGCGTTTTGA	TTACGCGTAGTCCGGGACGTCGTACGGGTAGAGTTCAGCGAAAGCACCTGC		
TGME49_210370	CACCACAGCCCGTTACGCTTGCAGC	TTACGCGTAGTCCGGGACGTCGTACGGGTAAACGGAGGGAAGAAACGGGG		
TGME49_218270	CACCGGCTGGAGATTTTTCCCGGAACT	TTACGCGTAGTCCGGGACGTCGTACGGGTAAAGCCGACTTGCAGAAGGCAC		
TGME49_225160	CACCCCATGCTTCTGTTCCGAGAAAT	TTACGCGTAGTCCGGGACGTCGTACGGGTATGAATCCTTGAAACTGCGAATC		
TGME49_232020	CACCTGGCTCTCCAGCCGCGGCGATT	TTACGCGTAGTCCGGGACGTCGTACGGGTATCGTCGCCGGCGCCTTCCGC		
TGME49_237180	CACCAGAAACCGCTGCTGAGGAATGG	TTACGCGTAGTCCGGGACGTCGTACGGGTACATAAGAAATTTATTTATGGAGGCG		
TGME49_258360	CACCGCGTGTTCGTCAGGATCTGCT	TTACGCGTAGTCCGGGACGTCGTACGGGTACCCGTTAAGATGCGCAACGAC		
TGME49_261740	CACCACAAAACGGGGAGCA	TTACGCGTAGTCCGGGACGTCGTACGGGTACGGTCTTTTCCACCTTTCACAG		
Primers used for gene knock-out (5'-3')				
TGME49_201860	<i>GGGGACAAGTTTGTACAAAAAGCAGGCTTATGCTTCGTACGACCAACG</i>	<i>GGGGACAACCTTTGTATAGAAAAGTTGTGGCTAATGTATCCACGTGG</i>		
TGME49_201860	<i>GGGGACAACCTTTGTATAATAAAGTTGCTGAGGTCCAAGCAGCACCGAA</i>	<i>GGGGACCACCTTTGTACAAGAAAAGCTGGGTAGAGAGAGTCGGGTTTCGTTCCG</i>		
TGME49_218270	<i>GGGGACAAGTTTGTACAAAAAGCAGGCTTAAAGACCTCTCGCCTCGGATT</i>	<i>GGGGACAACCTTTGTATAGAAAAGTTGCTTAGGCCTAAGTGTGGAGC</i>		
TGME49_218270	<i>GGGGACAACCTTTGTATAATAAAGTTGCTCTAGTCGTCTCGGATGATGG</i>	<i>GGGGACCACCTTTGTACAAGAAAAGCTGGGTATGTCGCTGGAGGATTCATGG</i>		
TGME49_261740	<i>GGGGACAAGTTTGTACAAAAAGCAGGCTGTAGTAACCTCGGATACACTTC</i>	<i>GGGGACAACCTTTGTATAGAAAAGTTGGGTGGACCGCCGAACATCATTTGTTTC</i>		
TGME49_261740	<i>GGGGACAACCTTTGTATAGAAAAGTTGGGTGGACCGCCGAACATCATTTGTTTC</i>	<i>GGGGACCACCTTTCTTGTACAAAAGTGGTAAAGGGCATGTTTTGACACGGG</i>		
	P1	P2	P3	P4
TGME49_201860	CGTGACTTGAAATGCAGTCC	GATCCAGACGCTTCAATGC	CACCTCGCGACCACCTGCGTTTTGA	TTACGCGTAGTCCGGGACGTCGTACGGGTAGAGTTCAGCGAAAGCACCTGC
TGME49_218270	CACCTCGACTGACATCTCAG	GATCCAGACGCTTCAATGC	CACCGGCTGGAGATTTTTCCCGGAACT	TTACGCGTAGTCCGGGACGTCGTACGGGTAAAGCCGACTTGCAGAAGGCAC
Primers used for generation of recombinant ROP47 peptide antigen (5'-3')				
TGME49_261740	CACCATTGTCTCGCCGC		CITTCTTTTTCGACCTTTCACAG	

Table 2. *Toxoplasma* genes in the rhopty cluster.

Probeset_id	ToxoDB V8_ID	Product Description	Subcellular localization	Reference	Confirmed	Exons	Transmembrane Domains	Predicted Signal Peptide	Cell Cycle Microarray RMA	
									Max	Min
20.m08222	TGME49_203990	rhopty protein ROP12 (ROP12)	RHOPTRY	Bradley et al., 2005	Confirmed	3	1	Yes	12.0	7.7
20.m03896	TGME49_205250	rhopty protein ROP18 (ROP18)	RHOPTRY	Taylor et al., 2006	Confirmed	1	1	Yes	13.8	10.6
27.m00091	TGME49_211290	rhopty protein ROP15 (ROP15)	RHOPTRY	Bradley et al., 2005	Confirmed	6	0	Yes	12.7	8.9
33.m02185	TGME49_214080	toxofilin	RHOPTRY	Bradley et al., 2005	Confirmed	1	1	Yes	12.2	8.9
33.m01398	TGME49_215775	rhopty protein ROP8 (ROP8)	RHOPTRY	Beckers et al., 1997	Confirmed	1	1	Yes	13.0	9.4
42.m00026	TGME49_223920	rhopty neck protein RON3 (RON3)	RHOPTRY	Bradley et al., 2005	Confirmed	21	1	Yes	12.3	7.9
42.m03584	TGME49_227810	rhopty kinase family protein ROP11 (incomplete catalytic triad) (ROP11)	RHOPTRY	Bradley et al., 2005	Confirmed	1	0	Yes	13.1	9.0
44.m06355	TGME49_229010	rhopty neck protein RON4 (RON4)	RHOPTRY	Bradley et al., 2005	Confirmed	20	0	Yes	11.8	7.7
44.m00026	TGME49_230350	rhopty neck protein RON11 (RON11)	RHOPTRY	Beck et al., 2013	Confirmed	16	4		10.4	6.5
49.m03277	TGME49_242240	rhopty kinase family protein ROP19A (ROP19A)	RHOPTRY	Peixoto et al., 2010	Confirmed	1	0	Yes	10.7	7.4
52.m01543	TGME49_252360	rhopty kinase family protein ROP24 (incomplete catalytic triad) (ROP24)	RHOPTRY	Peixoto et al., 2010	Confirmed	1	0	Yes	11.8	8.0
55.m04748	TGME49_258230	rhopty kinase family protein ROP20 (ROP20)	RHOPTRY	Peixoto et al., 2010	Confirmed	1	1	Yes	10.1	6.5
55.m08191	TGME49_258580	rhopty protein ROP17 (ROP17)	RHOPTRY	Peixoto et al., 2010	Confirmed	1	0	Yes	13.1	9.8
55.m00092	TGME49_258660	rhopty protein ROP6 (ROP6)	RHOPTRY	Sohn et al., 1999	Confirmed	5	1	Yes	13.3	9.5
55.m00167	TGME49_261750	rhopty neck protein RON10 (RON10)	RHOPTRY	Lamarque et al., 2012	Confirmed	7	0		11.3	6.7
55.m08219	TGME49_262730	rhopty protein ROP16 (ROP16)	RHOPTRY	Bradley et al., 2005	Confirmed	1	0	Yes	11.9	8.5
59.m03479	TGME49_269885	rhopty metalloprotease toxolysin TLN1 (TLN1)	RHOPTRY	Hajagos et al., 2011	Confirmed	12	1	Yes	11.6	6.9
74.m00767	TGME49_282055	protein phosphatase PP2C-hn (PP2CHN)	RHOPTRY	Gilbert et al., 2007	Confirmed	13	0		11.5	6.9
80.m02343	TGME49_291960	rhopty kinase family protein ROP40 (incomplete catalytic triad) (ROP40)	RHOPTRY	Peixoto et al., 2010	Confirmed	3	0	Yes	13.1	9.2
83.m02145	TGME49_295110	rhopty protein ROP7 (ROP7)	RHOPTRY	Hajj et al., 2006	Confirmed	1	1	Yes	12.7	8.6

113.m00009	TGME49_297960	rhopty neck protein RON6 (RON6)	RHOPTRY	Proellocks et al., 2006	Confirmed	10	1	Yes	11.5	7.0
129.m00252	TGME49_299060	sodium/hydrogen exchanger NHE2	RHOPTRY	Karasov et al., 2005	Confirmed	12	13		10.4	6.5
145.m00331	TGME49_300100	rhopty neck protein RON2 (RON2)	RHOPTRY	Bradley et al., 2005	Confirmed	10	3	Yes	11.4	6.5
541.m00141	TGME49_306060	rhopty neck protein RON8 (RON8)	RHOPTRY	Straub et al., 2009	Confirmed	15	1	Yes	11.9	7.0
551.m00238	TGME49_308090	rhopty protein ROP5 (ROP5)	RHOPTRY	Bradley et al., 2005	Confirmed	1	0	Yes	13.5	10.2
583.m09207	TGME49_308810	rhopty neck protein RON9 (RON9)	RHOPTRY	Lamarque et al., 2012	Confirmed	9	1		10.8	6.5
583.m00003	TGME49_309590	rhopty protein ROP1 (ROP1)	RHOPTRY	Ossorio et al., 1992	Confirmed	1	0	Yes	12.6	9.4
583.m00597	TGME49_310010	rhopty neck protein RON1 (RON1)	RHOPTRY	Bradley et al., 2005	Confirmed	5	1	Yes	11.3	6.8
583.m00636	TGME49_311470	rhopty neck protein RON5 (RON5)	RHOPTRY	Straub et al., 2009	Confirmed	38	0	Yes	12.3	7.6
583.m00692	TGME49_315220	rhopty protein ROP14 (ROP14)	RHOPTRY	Bradley et al., 2005	Confirmed	11	11		10.9	6.5
583.m05686	TGME49_315490	rhopty protein ROP10 (ROP10)	RHOPTRY	Bradley et al., 2005	Confirmed	1	0	Yes	11.1	6.6
49.m05689	TGME49_242118	myosin-light-chain kinase	RHOPTRY	Peixoto et al., 2010		1	1		10.8	6.5
20.m00331	TGME49_202200	hypothetical protein	RHOPTRY	Bradley et al., 2005		7	0		12.0	7.9
27.m00846	TGME49_211260	rhopty kinase family protein ROP26 (incomplete catalytic triad) (ROP26)	RHOPTRY	Peixoto et al., 2010		2	0		12.7	9.8
41.m01337	TGME49_222100	hypothetical protein	RHOPTRY	Bradley et al., 2005		1	0	Yes	11.0	6.5
49.m03276	TGME49_242230	rhopty kinase family protein ROP29 (ROP29)	RHOPTRY	Peixoto et al., 2010		1	1		10.8	6.9
49.m03399	TGME49_244250	hypothetical protein	RHOPTRY	Bradley et al., 2005		4	0		11.2	6.9
52.m01529	TGME49_252200	<i>Toxoplasma</i> palmitoyl acyltransferase TgDHHC7	RHOPTRY	Beck et al., 2013		8	4		10.1	6.5
52.m01582	TGME49_253370	hypothetical protein (RON4L1)	RHOPTRY	Boothroyd and Dubremetz, 2008		30	0	Yes	10.0	6.5
55.m04788	TGME49_258800	rhopty kinase family protein ROP31 (ROP31)	RHOPTRY	Peixoto et al., 2010		1	1	Yes	10.0	6.9
55.m05020	TGME49_262920	trypsin domain-containing protein	RHOPTRY	Bradley et al., 2005		11	1	Yes	9.8	6.5
83.m01271	TGME49_294560	rhopty kinase family protein ROP37 (incomplete catalytic triad) (ROP37)	RHOPTRY	Peixoto et al., 2010		1	0	Yes	10.6	6.5
83.m01285	TGME49_294790	hypothetical protein	RHOPTRY	Bradley et al., 2005		1	0		13.0	9.7
113.m00755	TGME49_297070	hypothetical protein	RHOPTRY	Bradley et al., 2005		2	0	Yes	12.6	8.2

583.m00694	TGME49_315210	rhoptyr protein, putative (ROP14B)	RHOPTRY	Reid et al., 2012	11	7		10.7	6.5
20.m00355	TGME49_201520	protein phosphatase 2C domain-containing protein			9	0	Yes	11.0	6.6
20.m05880	TGME49_201520	protein phosphatase 2C domain-containing protein			9	0	Yes	11.0	6.5
20.m03673	TGME49_201760	hypothetical protein			4	0		10.5	6.5
20.m03682	TGME49_201860	hypothetical protein			2	1	Yes	13.3	9.8
20.m03718	TGME49_202420	hypothetical protein			2	1		9.2	6.5
20.m00003	TGME49_202500	GAPM1a			4	6		13.7	10.2
20.m03760	TGME49_203010	aurora kinase			1	0		9.9	6.5
20.m05981	TGME49_203930	hypothetical protein			6	5		10.6	6.6
20.m03880	TGME49_204880	hypothetical protein			1	0	Yes	9.3	6.5
20.m03902	TGME49_205330	hypothetical protein			2	8		12.7	8.6
20.m03905	TGME49_205360	hypothetical protein			20	2	Yes	9.7	6.5
20.m03977	TGME49_206710	hypothetical protein			5	0		8.5	6.5
25.m01833	TGME49_208910	hypothetical protein			3	0		9.2	6.5
25.m01852	TGME49_209170	hypothetical protein			1	0	Yes	9.6	6.5
25.m01855	TGME49_209200	hypothetical protein			12	0		9.0	6.5
25.m01859	TGME49_209250	hypothetical protein			7	1		10.8	7.8
26.m00235	TGME49_210270	hypothetical protein			3	0		10.1	7.6
26.m00242	TGME49_210370	hypothetical protein			1	0	Yes	12.2	8.2
26.m00370	TGME49_210420	hypothetical protein			1	1		11.3	7.1
27.m00828	TGME49_210820	hypothetical protein			3	10		9.3	6.5
28.m00429	TGME49_211850	hypothetical protein			3	0		11.8	9.1
31.m00934	TGME49_212980	hypothetical protein			2	1		11.8	8.6
33.m02670	TGME49_214400	hypothetical protein			2	0		10.7	6.7

35.m00004	TGME49_216080	apical complex lysine methyltransferase	4	0		11.3	8.3
35.m00895	TGME49_216620	EF hand domain-containing protein	39	7		9.0	6.5
37.m00748	TGME49_217520	hypothetical protein	2	0	Yes	12.5	8.8
38.m01037	TGME49_218240	hypothetical protein	2	1	Yes	9.6	6.5
38.m01040	TGME49_218270	hypothetical protein	1	8	Yes	12.7	8.4
41.m01283	TGME49_221250	hypothetical protein	8	0		9.8	6.5
41.m00036	TGME49_221620	beta-tubulin, putative	4	0		13.1	9.7
41.m01316	TGME49_221675	hypothetical protein	9	1	Yes	8.9	6.5
42.m03399	TGME49_225020	hypothetical protein	4	0		9.5	6.5
42.m03409	TGME49_225160	hypothetical protein	1	2	Yes	11.5	7.8
42.m07438	TGME49_225200	hypothetical protein	33	0	Yes	9.2	6.5
42.m00061	TGME49_225320	hypothetical protein	5	0	Yes	10.8	6.5
42.m00060	TGME49_225330	hypothetical protein	6	0	Yes	9.8	6.5
42.m03456	TGME49_225860	hypothetical protein	4	0		9.8	6.5
42.m03481	TGME49_226220	alveolin domain containing intermediate filament IMC9 (ALV6/IMC9)	9	0		9.7	6.5
42.m00093	TGME49_227000	hypothetical protein	16	0		9.9	6.5
44.m02549	TGME49_229500	hypothetical protein	2	3		9.8	6.6
44.m02600	TGME49_230480	hypothetical protein	1	2		11.4	7.5
44.m02630	TGME49_231000	START domain-containing protein	18	1		9.0	6.5
44.m02636	TGME49_231070	protein kinase	1	0		10.8	6.8
44.m02644	TGME49_231160	hypothetical protein	3	0		12.7	9.5
44.m02678	TGME49_231840	hypothetical protein	6	0		9.7	6.5
44.m02696	TGME49_232020	hypothetical protein	2	1		8.8	6.5

44.m02714	TGME49_232260	hypothetical protein	3	3		10.0	6.5
44.m02750	TGME49_232780	hypothetical protein	8	0		9.5	6.5
46.m01616	TGME49_234540	hypothetical protein	7	9		10.2	6.5
46.m02875	TGME49_235130	transmembrane protein	4	0	Yes	9.3	6.5
46.m01643	TGME49_235380	hypothetical protein	11	0		8.9	6.5
46.m01716	TGME49_236860	haloacid dehalogenase family hydrolase domain-containing protein	2	12		9.8	6.5
46.m01722	TGME49_236960	transporter, major facilitator family protein	10	10		10.8	6.8
46.m01740	TGME49_237180	hypothetical protein	1	0	Yes	12.1	7.8
46.m01741	TGME49_237190	hypothetical protein	4	0		9.7	6.5
46.m01743	TGME49_237210	Tyrosine kinase-like (TKL) protein	4	0		10.2	6.5
49.m03087	TGME49_238150	hypothetical protein	1	8		12.7	10.2
49.m03179	TGME49_239830	TBC domain-containing protein	13	0		10.7	6.5
49.m03213	TGME49_240460	AP2 domain transcription factor AP2VI-1 (AP2VI1)	2	0		10.9	6.8
49.m07198	TGME49_240730	hypothetical protein	2	4		9.0	6.5
49.m07245	TGME49_241000	hypothetical protein	1	4		11.0	7.0
49.m03332	TGME49_243200	hypothetical protein	5	0		12.2	6.6
49.m00056	TGME49_243690	hypothetical protein	4	1		12.1	8.0
49.m03388	TGME49_244080	hypothetical protein	3	1	Yes	10.7	6.5
49.m03412	TGME49_244470	hypothetical protein	8	0		10.0	7.0
50.m03074	TGME49_245550	hypothetical protein	1	4	Yes	10.0	6.5
50.m03107	TGME49_246182	hypothetical protein				10.1	7.2
50.m03131	TGME49_246710	hypothetical protein	1	3		9.0	6.5
50.m03132	TGME49_246720	hypothetical protein	1	0		10.2	7.2
50.m03154	TGME49_247195	microneme protein MIC15 (MIC15)	42	1	Yes	9.8	6.5

50.m03253	TGME49_248690	hypothetical protein	7	0		9.4	6.8
50.m03302	TGME49_249440	hypothetical protein	2	0		9.9	6.5
50.m03315	TGME49_249570	hypothetical protein	20	1	Yes	11.4	6.8
52.m01567	TGME49_253140	hypothetical protein	3	0		9.3	6.5
52.m01583	TGME49_253380	AP2 domain transcription factor AP2III-2 (AP2III2)	1	0		10.5	6.8
52.m01598	TGME49_253600	hypothetical protein	11	0		9.6	6.5
52.m01644	TGME49_254290	hypothetical protein	1	0		9.0	6.5
55.m04629	TGME49_255700	hypothetical protein	6	0		9.7	6.5
55.m08188	TGME49_256030	hypothetical protein	6	0		12.1	7.6
55.m04752	TGME49_258360	hypothetical protein	7	0	Yes	11.6	7.0
55.m00096	TGME49_258700	transporter, major facilitator family protein	12	11		9.3	6.5
55.m04796	TGME49_258900	hypothetical protein	4	0		9.3	6.5
55.m04843	TGME49_259700	hypothetical protein	2	3		11.1	6.9
55.m00144	TGME49_260820	IMC sub-compartment protein ISP1 (ISP1)	3	0		12.0	8.5
55.m04955	TGME49_261740	hypothetical protein	1	1	Yes	14.2	11.9
57.m01689	TGME49_264600	hypothetical protein	2	4		12.2	7.5
57.m01720	TGME49_265080	Tubulin-tyrosine ligase family protein	12	0		9.6	6.5
57.m01755	TGME49_265650	protein phosphatase 2C domain- containing protein	1	0	Yes	9.7	6.5
57.m01783	TGME49_266300	hypothetical protein	2	0		12.3	8.0
57.m01792	TGME49_266435	hypothetical protein	34	0		9.4	6.5
57.m01834	TGME49_267070	aquaporin 2	1	11		11.6	7.7
59.m03403	TGME49_268760	hypothetical protein				12.2	8.6
59.m00087	TGME49_268870	tetratricopeptide repeat-containing protein	10	0		10.3	6.5
59.m00092	TGME49_269330	hypothetical protein	10	0		10.1	6.6

59.m00029	TGME49_269340	hypothetical protein	4	1		11.4	6.9
59.m03542	TGME49_270890	hypothetical protein	3	0		9.4	6.5
59.m00038	TGME49_271270	hypothetical protein	10	1		11.2	7.0
59.m03707	TGME49_273860	hypothetical protein	1	0		11.0	8.2
64.m00327	TGME49_275670	alveolin domain containing intermediate filament IMC15 (ALV5/IMC15)	9	0		10.8	6.5
64.m00582	TGME49_276130	cathepsin CPC2 (CPC2)	10	0	Yes	9.7	6.5
65.m01152	TGME49_278130	hypothetical protein	14	0		9.3	6.5
65.m01964	TGME49_278510	protein phosphatase 2C domain- containing protein	10	1	Yes	11.1	7.3
65.m02537	TGME49_278920	hypothetical protein	5	6		9.5	6.5
69.m00143	TGME49_279420	hypothetical protein	1	0		11.3	6.5
72.m00385	TGME49_280480	EF hand domain-containing protein	1	4		11.5	6.8
72.m00685	TGME49_280670	hypothetical protein	7	10	Yes	11.3	6.7
74.m00455	TGME49_282070	hypothetical protein	9	0		10.8	6.5
74.m00465	TGME49_282210	AP2 domain transcription factor AP2VIIa-8 (AP2VIIA8)	2	0		9.3	6.5
76.m01597	TGME49_285290	hypothetical protein	1	3	Yes	10.3	6.5
76.m01608	TGME49_285650	hypothetical protein	2	0		8.9	6.5
76.m01626	TGME49_285870	SAG-related sequence SRS20A (SRS20A)	2	1	Yes	12.8	8.7
76.m01662	TGME49_286500	hypothetical protein	1	10		9.9	6.5
80.m02122	TGME49_287970	hypothetical protein	3	0		10.4	6.5
80.m02181	TGME49_288950	AP2 domain transcription factor AP2IX-4 (AP2IX4)	1	0		9.7	6.5
80.m03982	TGME49_289150	hypothetical protein	5	1		9.4	6.5
80.m03946	TGME49_289970	hypothetical protein	3	0		11.7	9.3
83.m01220	TGME49_293540	hypothetical protein	6	0		10.3	6.5

83.m01311	TGME49_295100	hypothetical protein	4	0		9.5	6.9
86.m00370	TGME49_295420	hypothetical protein	9	0		10.4	7.0
86.m00377	TGME49_295620	hypothetical protein	4	0		9.7	6.5
113.m01286	TGME49_298010	hypothetical protein	52	0		9.1	6.5
145.m00603	TGME49_300220	hypothetical protein	15	4	Yes	9.4	6.5
145.m00607	TGME49_300360	ADP/ATP translocase	6	2		9.5	6.5
162.m00326	TGME49_301420	hypothetical protein	7	0		11.8	7.5
541.m00127	TGME49_305250	hypothetical protein	6	6		10.5	7.1
541.m01166	TGME49_305270	hypothetical protein	6	0	Yes	10.2	6.5
541.m00131	TGME49_305510	hypothetical protein	7	0	Yes	11.9	7.9
541.m01185	TGME49_305590	ABC transporter transmembrane region domain-containing protein	9	9		9.5	6.5
542.m00226	TGME49_307020	hypothetical protein	3	0		9.4	6.5
551.m00232	TGME49_308010	hypothetical protein	1	0		9.0	6.5
583.m05275	TGME49_309160	IgA-specific metalloendopeptidase	1	0	Yes	10.4	6.5
583.m09102	TGME49_310240	hypothetical protein	1	2		10.3	6.5
583.m11449	TGME49_310740	hypothetical protein	2	7		9.3	6.5
583.m00659	TGME49_311800	endonuclease/exonuclease/phosphatase family protein	6	0		9.1	6.6
583.m00645	TGME49_312150	hypothetical protein	6	1		11.7	7.6
583.m09217	TGME49_313780	hypothetical protein	1	0		11.1	7.3
583.m09134	TGME49_314260	hypothetical protein	1	1		12.1	8.2
583.m05709	TGME49_315780	myosin regulatory light chain, putative	7	0		11.5	8.6
583.m05738	TGME49_316260	hypothetical protein	2	7		12.3	8.1
583.m09158	TGME49_316280	transporter, major facilitator family protein	4	8		11.7	7.3
583.m05758	TGME49_316540	IMC sub-compartment protein ISP3 (ISP3)	1	0		11.1	8.0

611.m00052	TGME49_317705	enoyl-CoA hydratase/isomerase family protein	8	0		10.5	7.2
641.m01483	TGME49_318470	AP2 domain transcription factor AP2IV-4 (AP2IV4)	1	0		9.3	6.5
641.m00181	TGME49_320740	hypothetical protein	5	1	Yes	11.7	8.6
645.m00324	TGME49_321600	hypothetical protein	7	0		9.6	6.5

Genes highlighted in grey were further characterized in this study.

RMA, Robust Multi-array Average

Table 3. Characteristics of candidate genes

Candidate gene ToxoDB V8 ID	Alias	Signal peptide	TM domains	Conserved domains	Paralogue (% protein identity)	Paralogue present in rhoptry cluster	Homologue in <i>Neospora caninum</i> (% protein identity)
TGME49_201860	-	Yes	1	-	TGME49_301390 (31%)	No	NCLIV_022900 (75%)
TGME49_218270	ROP48	Yes	8	-	TGME49_209810 (29%)	No	NCLIV_061950 (71%)
TGME49_232020	RON12	Yes, in GT1 and CEP	1	-	TGME49_244726 (27%)	No	NCLIV_032020 (59%)
TGME49_261740	ROP47	Yes	1	-	-		NCLIV_025740 (43%)

Supplementary Table S1. Rhoptry encoding genes expressed above background and present on the *Toxoplasma* array

ToxoDB V8_ID	Product Description	In Rhoptry Cluster	Reference	Confirmed
TGME49_203990	rhoptry protein ROP12 (ROP12)	Yes	Bradley et al., 2005	Confirmed
TGME49_205250	rhoptry protein ROP18 (ROP18)	Yes	Taylor et al., 2006	Confirmed
TGME49_211290	rhoptry protein ROP15 (ROP15)	Yes	Bradley et al., 2005	Confirmed
TGME49_214080	toxofilin	Yes	Bradley et al., 2005	Confirmed
TGME49_215775	rhoptry protein ROP8 (ROP8)	Yes	Beckers et al., 1997	Confirmed
TGME49_223920	rhoptry neck protein RON3 (RON3)	Yes	Bradley et al., 2005	Confirmed
TGME49_227810	rhoptry kinase family protein ROP11 (incomplete catalytic triad) (ROP11)	Yes	Bradley et al., 2005	Confirmed
TGME49_229010	rhoptry neck protein RON4 (RON4)	Yes	Bradley et al., 2005	Confirmed
TGME49_230350	rhoptry neck protein RON11 (RON11)	Yes	Beck et al., 2013	Confirmed
TGME49_242240	rhoptry kinase family protein ROP19A (ROP19A)	Yes	Peixoto et al., 2010	Confirmed
TGME49_252200	<i>Toxoplasma</i> palmitoyl acyltransferase TgDHHC7	Yes	Beck et al., 2013	Confirmed
TGME49_252360	rhoptry kinase family protein ROP24 (incomplete catalytic triad) (ROP24)	Yes	Peixoto et al., 2010	Confirmed
TGME49_258230	rhoptry kinase family protein ROP20 (ROP20)	Yes	Peixoto et al., 2010	Confirmed
TGME49_258580	rhoptry protein ROP17 (ROP17)	Yes	Peixoto et al., 2010	Confirmed
TGME49_258660	rhoptry protein ROP6 (ROP6)	Yes	Sohn et al., 1999	Confirmed
TGME49_261750	rhoptry neck protein RON10 (RON10)	Yes	Lamarque et al., 2012	Confirmed
TGME49_262730	rhoptry protein ROP16 (ROP16)	Yes	Bradley et al., 2005	Confirmed
TGME49_282055	protein phosphatase PP2C-hn (PP2CHN)	Yes	Gilbert et al., 2007	Confirmed
TGME49_291960	rhoptry kinase family protein ROP40 (incomplete catalytic triad) (ROP40)	Yes	Peixoto et al., 2010	Confirmed
TGME49_295110	rhoptry protein ROP7 (ROP7)	Yes	Hajj et al., 2006	Confirmed
TGME49_297960	rhoptry neck protein RON6 (RON6)	Yes	Proellocks et al., 2006	Confirmed
TGME49_299060	sodium/hydrogen exchanger NHE2 (ROP)	Yes	Karasov et al., 2005	Confirmed

TGME49_300100	rhoptry neck protein RON2 (RON2)	Yes	Bradley et al., 2005	Confirmed
TGME49_306060	rhoptry neck protein RON8 (RON8)	Yes	Straub et al., 2009	Confirmed
TGME49_308090	rhoptry protein ROP5 (ROP5)	Yes	Bradley et al., 2005	Confirmed
TGME49_308810	rhoptry neck protein RON9 (RON9)	Yes	Lamarque et al., 2012	Confirmed
TGME49_309590	rhoptry protein ROP1 (ROP1)	Yes	Ossorio et al., 1992	Confirmed
TGME49_310010	rhoptry neck protein RON1 (RON1)	Yes	Bradley et al., 2005	Confirmed
TGME49_311470	rhoptry neck protein RON5 (RON5)	Yes	Straub et al., 2009	Confirmed
TGME49_315220	rhoptry protein ROP14 (ROP14)	Yes	Bradley et al., 2005	Confirmed
TGME49_315490	rhoptry protein ROP10 (ROP10)	Yes	Bradley et al., 2005	Confirmed
TGME49_242118	myosin-light-chain kinase	Yes	Peixoto et al., 2010	
TGME49_202200	hypothetical protein	Yes	Bradley et al., 2005	
TGME49_211260	rhoptry kinase family protein ROP26 (incomplete catalytic triad) (ROP26)	Yes	Peixoto et al., 2010	
TGME49_222100	hypothetical protein	Yes	Bradley et al., 2005	
TGME49_242230	rhoptry kinase family protein ROP29 (ROP29)	Yes	Peixoto et al., 2010	
TGME49_244250	hypothetical protein	Yes	Bradley et al., 2005	
TGME49_253370	hypothetical protein (RON4L1)	Yes	Boothroyd and Dubremetz, 2008	
TGME49_258800	rhoptry kinase family protein ROP31 (ROP31)	Yes	Peixoto et al., 2010	
TGME49_262920	trypsin domain-containing protein	Yes	Bradley et al., 2005	
TGME49_294560	rhoptry kinase family protein ROP37 (incomplete catalytic triad) (ROP37)	Yes	Peixoto et al., 2010	
TGME49_294790	hypothetical protein	Yes	Bradley et al., 2005	
TGME49_297070	hypothetical protein	Yes	Bradley et al., 2005	
TGME49_315210	rhoptry protein, putative (ROP14B)	Yes	Reid et al., 2012	
TGME49_243730	rhoptry protein ROP9 (ROP9)	No	Reichman et al., 2002	Confirmed
TGME49_261440	ARM repeats containing protein TgARO	No	Cabrera et al., 2012	Confirmed

TGME49_262050	rhoptry kinase family protein ROP39 (ROP39)	No	Peixoto et al., 2010	Confirmed
TGME49_269885	rhoptry metalloprotease toxolysin TLN1 (TLN1)	No	Hajagos et al., 2011	Confirmed
TGME49_314500	subtilisin SUB2 (SUB2)	No	Miller et al., 2003	Confirmed
TGME49_242110	rhoptry kinase family protein ROP38 (ROP38)	No	Peixoto et al., 2010	Confirmed
TGME49_314250	bradyzoite rhoptry protein BRP1 (BRP1)	No	Schwarz et al., 2005	
TGME49_201130	rhoptry kinase family protein ROP33 (ROP33)	No	Peixoto et al., 2010	
TGME49_202780	rhoptry kinase family protein ROP25 (ROP25)	No	Peixoto et al., 2010	
TGME49_209985	cAMP-dependent protein kinase	No	Bradley et al., 2005	
TGME49_227010	rhoptry kinase family protein ROP30 (ROP30)	No	Peixoto et al., 2010	
TGME49_230470	rhoptry kinase family protein ROP46, putative	No	Peixoto et al., 2010	
TGME49_239600	rhoptry kinase family protein ROP23 (incomplete catalytic triad) (ROP23)	No	Peixoto et al., 2010	
TGME49_240090	rhoptry kinase family protein ROP34, putative	No	Peixoto et al., 2010	
TGME49_249470	rhoptry kinase family protein, truncated (incomplete catalytic triad)	No	Peixoto et al., 2010	
TGME49_266100	rhoptry kinase family protein ROP41 (ROP41)	No	Peixoto et al., 2010	
TGME49_270320	protein phosphatase 2C domain-containing protein	No	Bradley et al., 2005	
TGME49_270920	rhoptry kinase family protein ROP32 (ROP32)	No	Peixoto et al., 2010	
TGME49_281675	rhoptry kinase family protein ROP45 (ROP45)	No	Peixoto et al., 2010	
TGME49_294400	rhoptry neck protein RON2L1 (RON2L1)	No	Fritz et al., 2012	
TGME49_304740	rhoptry kinase family protein ROP35 (ROP35)	No	Peixoto et al., 2010	
TGME49_315940	rhoptry protein, putative	No	ToxoDB.org	

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