Toxoplasma Cathepsin Protease B and Aspartyl Protease 1 Are Dispensable for Endolysosomal Protein Digestion

Christian McDonald, a,*, David Smith, a,*, Manlio Di Cristina, b Geetha Kannan, a Zhicheng Dou, c Vern B. Carruthers a

a Department of Microbiology and Immunology, University of Michigan Medical School, Ann Arbor, Michigan, USA
b Department of Chemistry, Biology and Biotechnology, University of Perugia, Perugia, Italy
c Department of Biological Sciences, Clemson University, Clemson, South Carolina, USA

Christian McDonald, David Smith, and Zhicheng Dou contributed equally to this article. Their order was determined alphabetically and by seniority.

ABSTRACT The lysosome-like vacuolar compartment (VAC) is a major site of proteolysis in the intracellular parasite Toxoplasma gondii. Previous studies have shown that genetic ablation of a VAC-residing cysteine protease, cathepsin protease L (CPL), resulted in the accumulation of undigested protein in the VAC and loss of parasite viability during the chronic stage of infection. However, since the maturation of another VAC localizing protease, cathepsin protease B (CPB), is dependent on CPL, it remained unknown whether these defects result directly from ablation of CPL or indirectly from a lack of CPB maturation. Likewise, although a previously described cathepsin D-like aspartyl protease 1 (ASP1) could also play a role in proteolysis, its definitive residence and function in the Toxoplasma endolysosomal system were not well defined. Here, we demonstrate that CPB is not necessary for protein turnover in the VAC and that CPB-deficient parasites have normal growth and viability in both the acute and chronic stages of infection. We also show that ASP1 depends on CPL for correct maturation, and it resides in the T. gondii VAC, where, similar to CPB, it plays a dispensable role in protein digestion. Taken together with previous work, our findings suggest that CPL is the dominant protease in a hierarchy of proteolytic enzymes within the VAC. This unusual lack of redundancy for CPL in T. gondii makes it a single exploitable target for disrupting chronic toxoplasmosis.

IMPORTANCE Roughly one-third of the human population is chronically infected with the intracellular single-celled parasite Toxoplasma gondii, but little is known about how this organism persists inside people. Previous research suggested that a parasite proteolytic enzyme, termed cathepsin protease L, is important for Toxoplasma persistence; however, it remained possible that other associated proteolytic enzymes could also be involved in the long-term survival of the parasite during infection. Here, we show that two proteolytic enzymes associated with cathepsin protease L play dispensable roles and are dependent on cathepsin L to reach maturity, which differs from the corresponding enzymes in humans. These findings establish a divergent hierarchy of proteases and help focus attention principally on cathepsin protease L as a potential target for interrupting Toxoplasma chronic infection.

KEYWORDS Toxoplasma gondii, autophagy, cathepsin, lysosome, proteases

The apicomplexan parasite Toxoplasma gondii is an obligate intracellular protozoan that chronically infects ~2 billion people worldwide (1). Toxoplasmosis can be fatal for those who are immunocompromised, such as AIDS or organ transplant patients (1, 2). In pregnant women, Toxoplasma can overcome barriers at the maternal-fetal interface and parasitize the developing fetus, which can cause a spontaneous abortion or neurological pathologies in the surviving offspring (3). Strikingly, most immunocom-
petent infected individuals are asymptomatic despite being persistently infected. In-
sufficient understanding of the underlying pathways that permit long-term infection
precludes the development of viable treatment options to limit disease in at-risk
individuals.

The endolysosomal pathway is a vital system in eukaryotic organisms. Proteases
associated with these systems are of particular interest due to their role in various
biological processes, including catabolism and enzyme activation (2, 4, 5). The lys-
osomal cysteine proteases cathepsins L and B are ubiquitously expressed in eukaryotic
cells and together are integral for proteolytic activity within lysosomes (6). Single-
 knockout experiments in murine models show only a modest or indistinguishable
phenotype for cathepsin L or B deficiency, respectively (7, 8). For example, mouse
embryonic fibroblasts lacking cathepsin L present a nonlethal deficit in the turnover of
autophagosomes (7). This indicates that the absence of one of these proteases can be
overcome in mice. However, double-knockout experiments of both CPL and CPB had a
profound effect on mouse health, including pronounced neurodegeneration, defects in
motility, and, most significantly, lethality 3 weeks after birth (6, 9). In addition, mice
deficient in the lysosomal aspartyl protease cathepsin D die within 4 weeks of birth (10).
Cathepsin D also regulates the activation of cathepsin B (11). Thus, whereas cathepsins
L and B are individually dispensable in mammals, cathepsin D is critical for maintaining
proteolytic activity and survival in mammals.

Lysosomal cathepsin proteases are expressed in multicopy families in some patho-
genic protozoa, such as Plasmodium falciparum. P. falciparum is one of the causative
agents of malaria and, like many other protozoan parasites, relies on proteases for
various aspects of its pathogenesis (2). Plasmodium obtains host hemoglobin and
digests it in a specialized organelle, known as the food vacuole (FV), to procure
necessary nutrients (12, 13). Three cathepsin L-like cysteine proteases (falcipains 2a, 2b,
and 3) and four cathepsin D-like aspartyl proteases (plasmepsins I, II, and IV and
histoaspartic protease) localize to the FV and are key players in hemoglobin degrada-
tion (14–16). As with the mammalian systems mentioned above, functional redundancy
exists between these proteases (13, 17, 18). Knockout of falcipain-2a in one study
resulted in an accumulation of unprocessed hemoglobin in the FV; however, this was
a transient phenotype and was subsequently not seen in later stages of the parasite life
cycle, presumably due to the expression of falcipains 2b and 3 (17). Likewise, parasites
lacking any one of the four plasmepsins were slower growing yet viable (18). Taken
together, these findings suggest these proteases work cooperatively in the turnover of
host hemoglobin.

Like its Plasmodium kin, Toxoplasma possesses a recently characterized lysosome-
like organelle known as the vacuolar compartment (VAC), which houses proteases
canonically associated with lysosomes (19–21). Acute-stage parasites (tachyzoites) have
been shown to ingest host cytosolic proteins for the duration of the parasite cell cycle,
and the catabolism of these host-derived constituents occurs in the VAC (19, 22). In
contrast to Plasmodium, Toxoplasma encodes single copies of cathepsin L (CPL),
cathepsin B (CPB), and cathepsin D (aspartyl protease 1, or ASP1) in its genome. Of the
five cathepsin-like cysteine proteases encoded by the Toxoplasma genome, the VAC is
the major site for CPB and CPL (23–25). Recent investigations of the VAC uncovered
how chronic-stage parasites (bradyzoites) sustain themselves in the host indefinitely
following encystation. First, the genetic ablation of CPL has shown its contribution to
acute-stage virulence and invasion. Second, cystogenic strains of T. gondii deficient in
this protease accumulate large quantities of undigested material in the VAC of brady-
zoites in vitro. Finally, the viability of CPL-deficient bradyzoites was starkly reduced in
vitro, consistent with vastly reduced cyst burden in vivo (26). It remains unclear whether
this is due solely to CPL disruption or due to CPB maturation being dependent on CPL
(23). In other words, it is unknown whether the phenotypes observed in CPL-deficient
parasites are due to a lack of active CPB.

Of the seven aspartyl proteases identified in the Toxoplasma genome, only one,
ASP1, shares a clade with the P. falciparum FV plasmepsins (27, 28). This phylogenetic
relationship may point to a digestive function within the *T. gondii* endolysosomal system, including the VAC. In interphase parasites, ASP1 was reported to localize to vesicular structures in the parasite apical region that are distinct from apical secretory organelles (micronemes, rhoptries, and dense granules) (27). During daughter cell formation, ASP1 redistributed to the inner membrane complex (IMC) that delineates developing daughter parasites. Thus, although these findings suggest a dynamic localization of ASP1, the extent to which its phylogenetic heritage aligns with the occupation of an endolysosomal compartment remains unknown.

Although a critical requirement for protein turnover in the VAC in chronic-stage *T. gondii* was shown previously, key questions remain regarding the contribution of specific proteases to this degradation system. Therefore, in this study our aim was to define the role of *T. gondii* CPB and ASP1 in protein turnover in the VAC, delineate their position within a protease hierarchy, and determine their contribution to infection.

**RESULTS**

**Determining the extent of CPB’s role in tachyzoite growth and host-derived protein turnover.** Maturation of the *Toxoplasma* protease CPB was shown to be dependent on another VAC-residing protease, CPL. Since CPL-deficient parasites exhibit numerous phenotypes and do not produce mature CPB, it is unclear whether the phenotypes described are attributed to inactive CPB. To assess the role of CPB in *T. gondii*, we knocked out CPB in the cystogenic type II Prugniaud Δku80 strain (29) (termed Pru here) and confirmed this by PCR analysis (see Fig. S1 in the supplemental material), immunoblotting, and immunofluorescence (Fig. 1A and B). The inclusion of the previously described PΔcpl strain (26) in the immunoblot confirmed that maturation of CPB is dependent on expression of CPL, which is consistent with previous findings in a type I strain (23). Similarly, we found that CPB localizes to the VAC of tachyzoites (Fig. 1B), which is also consistent among strains (23). To test how tachyzoite replication was affected following CPB knockout, Pru, PΔcpb, and PΔcpl tachyzoites were grown in human foreskin fibroblast (HFF) cells, followed by the enumeration of parasites per parasitophorous vacuole (PV) to quantify replication. At 24 h postinfection, no measurable differences were seen between these strains; however, at 48 h postinfection, PΔcpl tachyzoites had slower growth than Pru, whereas replication of PΔcpb was indistinguishable from that of Pru (Fig. 1C). As mentioned above, the ingestion of host-derived material has been shown in the VAC of acute-stage parasites. Using a Chinese hamster ovarian (CHO)-K1 cell line inducibly expressing mCherry, we found that the accumulation of host-derived mCherry was significantly higher in PΔcpl tachyzoites than in both Pru and PΔcpb strains, whereas PΔcpb parasites showed accumulation comparable to that of Pru (Fig. 1D). In all, these findings suggest that CPB has a limited role in tachyzoite VAC digestive function.

**CPB maturation depends on CPL activity in bradyzoites.** We used a previously described series of strains (26) to explore the role of CPL activity in the maturation of itself and CPB in chronic-stage bradyzoites. PΔcplS1CPL and PΔcplB1CPL express CPL exclusively in tachyzoites and bradyzoites, respectively. PΔcplB1CPL* exclusively expresses a catalytically inactive form of CPL in bradyzoites. Using immunoblotting, we found CPL catalytic activity is necessary for normal self-maturation based on observing a pseudomature (*pmatCPL*) species in PΔcplB1CPL* bradyzoites (Fig. 2A). These results are consistent with previous findings in tachyzoites (23). When probed for CPB, immunoblots also displayed bands for mature CPB only in strains that express active CPL, namely, Pru and PΔcplB1CPL bradyzoites (Fig. 2B). Immature CPB was observed in ΔcplB1CPL*, confirming that CPL activity is necessary for maturation of CPB in bradyzoites.

**CPB is not necessary for turnover of autophagic material in bradyzoites.** We have previously demonstrated that CPL deficiency in *T. gondii* leads to the accumulation of autophagic material in the VAC in bradyzoites (26). However, due to CPB maturation being dependent on the proteolytic activity of CPL, parasites deficient in CPL will also be deficient in active CPB. Therefore, it remained a possibility that this
accumulation was actually due to a lack of active CPB in the VAC. To address this, we first used surface antigen 1 (SAG1) as a tachyzoite stage-specific marker as well as green fluorescent protein (GFP), which is expressed following bradyzoite differentiation (29), and found that differentiation was similar among the Pru, PΔcpb, and PΔcpl strains (Fig. 3A). Consistent with our observations of CPB and CPL colocalization in tachyzoites, these two proteases were also found to colocalize in bradyzoite cysts (Fig. 3B). To determine the extent to which CPB independently contributed to the turnover of autophagic material, we assessed the accumulation of dense puncta, indicative of undigested autophagosomes in PΔcpb bradyzoites, and compared this accumulation to that of Pru and PΔcpl bradyzoites. Our results showed that after 7 and 14 days of differentiation, punctate structures were larger in PΔcpl bradyzoites than Pru ones (Fig. 3C and D). However, no difference in puncta size was seen in PΔcpb bradyzoites compared to Pru bradyzoites. Collectively, our findings suggest that CPB does not play a major digestive role in the endolysosomal system of Toxoplasma during acute or chronic infection in vitro.

ASP1 resides in the Toxoplasma VAC and is dependent on CPL for normal maturation. Unlike the related apicomplexan, Plasmodium, Toxoplasma possesses one group A aspartyl protease, ASP1 (27). T. gondii ASP1 was shown to have punctate localization that extensively changes during tachyzoite replication in daughter cells and...
was predicted to be compartmentalized (27). The Toxoplasma VAC displays similar fragmentation during replication (21); therefore, we hypothesized that ASP1 resides within the VAC. To address this definitively, we generated antibodies to full-length recombinant ASP1 and created an ASP1-deficient mutant in the RH strain background (RΔasp1) along with genetically complementing the mutant (RΔasp1ASP1) (Fig. S2).

Immunoblots displayed mature ASP1 in parental strain RH and RΔasp1ASP1 complement strain but not in RΔasp1 (Fig. 4A). Note that despite affinity purifying the antibody on recombinant ASP1, several nonspecific bands were observed in the RΔasp1 lysate, including one that nearly comigrates with mature ASP1. PCR analysis (Fig. S2) and immunofluorescence staining confirmed the absence of ASP1 expression in newly invaded RΔasp1 tachyzoites. Immunofluorescence staining also revealed colocalization of ASP1 with CPL in RH and RΔasp1ASP1 (Fig. 4B).

Since CPL expression is required for maturation of CPB, we reasoned that CPL might also contribute to the maturation of ASP1. Accordingly, immunoblotting of RH strain parasites lacking CPL (RΔcpl) with anti-ASP1 revealed the accumulation of proASP1 and intermediate species of ASP1 (Fig. 5A), suggesting a role for CPL expression in ASP1 maturation. Note that the nonspecific band nearly comigrating with mature ASP1 confounded interpretation of how much mature ASP1 is present in RΔcpl tachyzoites. We also observed a corresponding increase in the staining for ASP1 and CPB in RΔcpl tachyzoites (Fig. 5B), indicating the accumulation of immature species of these proteases in the absence of CPL expression. Collectively, our findings suggest that ASP1 resides in the VAC and that its maturation involves CPL. The findings do not, however, rule out the possibility of other proteases contributing to the maturation of ASP1.

**ASP1 is not required for tachyzoite replication or digestion of host-derived protein in the VAC.** To further investigate the maturation and function of ASP1 in infection, we generated Pru parasites lacking ASP1 alone (PΔasp1) or lacking ASP1 and CPL (PΔasp1Δcpl) (Fig. S3). Immunoblotting of tachyzoite (Fig. 6A, left) or bradyzoite (Fig. 6A, right) corroborated a role for CPL in ASP1 maturation. Because the nonspecific band that comigrated with mature ASP1 is absent from bradyzoites, it also is apparent that maturation of ASP1 is strongly dependent on the expression of CPL in bradyzoites. The increased expression of both ASP1 and CPL in bradyzoites compared to that of tachyzoites is also evident (Fig. 6A, right). To assess the role of ASP1 in tachyzoite replication, parasites were allowed to infect HFF monolayers for 24 and 48 h. Enumeration of the parasites per PV displayed significant differences only after 48 h postinfection in RΔcpl and PΔasp1Δcpl strains (Fig. 6B). Interestingly, PΔasp1Δcpl parasites did
not replicate more slowly than Δcpl parasites, implying that the absence of CPL is principally responsible for the replication phenotype. We next tested for a possible role for ASP1 in digestion of host-derived protein in the VAC. Following the incubation of parasites with mCherry-expressing CHO-K1 cells, PΔasp1 failed to accumulate host-derived protein in the VAC, as indicated by the lack of mCherry signal. This suggests that ASP1 plays a role in the turnover of autophagosomes in the VAC in bradyzoites.

**FIG 3** CPL, not CPB, is the major protease necessary for the turnover of autophagosomes in the VAC in bradyzoites. (A) The rate of parasite differentiation from the tachyzoite stage to the bradyzoite stage was determined in vitro by assessment of the tachyzoite-specific antigen SAG1 and GFP under the control of a bradyzoite LDH2 promoter. Infected monolayers were cultured for the indicated days, fixed, probed for SAG1, and quantified. Bars indicate means ± standard deviations from 3 biological replicates. All strains were compared, and only statistical significance is shown. Unpaired two-tailed t test with Welch’s corrections was performed on the means from 3 biological replicates. (B) Immunofluorescence localization of CPB and CPL in bradyzoites. Parasite strains were differentiated for 7 days, fixed, and stained with anti-CPB (violet) and anti-CPL (blue). The inset in the merged image of Pru shows enlargement of the boxed region. Scale bars, 5 μm. (C) Phase-contrast microscopy was used to image bradyzoites in vitro following 7- and 14-day cultures. Enlarged dark puncta seen in PΔcpl bradyzoites are indicative of defective protein degradation in the VAC (arrows). Scale bars, 5 μm. (D) The size of the puncta in bradyzoites was measured. The following numbers of cysts across 3 biological replicates were used to analyze each strain: Pru (day 7, 109 cysts; day 14, 98 cysts), PΔcpb (day 7, 106 cysts; day 14, 102 cysts), and PΔcpl (day 7, 105 cysts; day 14, 97 cysts). Lines represent medians with 95% confidence intervals. For this data set, ROUT with a Q value of 0.1% was used to identify and remove 1 outlier each for Pru and PΔcpl for 14 days postdifferentiation. Since the data did not fit a normal distribution, a nonparametric Kruskal-Wallis test with Dunn’s multiple comparisons was used to compare the medians of data combined from 3 biological replicates. All strains were compared, and only significant differences are shown.
derived mCherry, whereas accumulation was seen in PΔcpl and PΔcplΔasp1 tachyzoites (Fig. 6C). These results mirror what was seen in CPB and indicate that neither CPB nor ASP1 is a major contributor to tachyzoite growth or endolysosomal digestive function, whereas CPL has a dominant role.

ASP1 plays a dispensable role in proteolytic turnover of autophagic material. It was also important to establish whether ASP1 could have a functional overlap in the turnover of autophagic material in bradyzoites. Therefore, similar to procedures for PΔcpb parasites, we compared the differentiation and size of puncta in PΔasp1 and PΔasp1Δcpl bradyzoites to those of Pru and PΔcpl bradyzoites after 7 and 14 days of differentiation. Differentiation was indistinguishable among the strains based on the decrease in SAG1-positive vacuoles and the corresponding increase of GFP-positive cysts (Fig. 7A). Automated measurement of puncta size also indicated no difference between PΔasp1 and Pru bradyzoites in turnover of autophagosomes at day 7 or 14 of differentiation, which is distinct from the enlarged puncta observed in PΔcpl (Fig. 7B and C). Interestingly, PΔasp1Δcpl bradyzoites showed an intermediate phenotype, with puncta that were smaller than those of PΔcpl but potentially larger than those of Pru.

FIG 4  T. gondii ASP1, a cathepsin D ortholog that localizes to the VAC. (A) Immunoblot detection of T. gondii ASP1 in tachyzoites. Protein bands are labeled as proform (pro), intermediate form (int), and mature form (mat) based on their molecular weights. Polypeptides marked with asterisks are nonspecific bands. (B) Immunofluorescence assay of tachyzoites after fixation, staining, and probing for T. gondii ASP1 (green) and CPL (red). Arrowheads indicate colocalization of ASP1 with CPL in the VAC. Scale bars, 2 μm.

FIG 5  T. gondii immature ASP1 accumulates in tachyzoites lacking active CPL in the VAC. (A) Immunoblot of tachyzoite lysates probed for ASP1 and CPL along with actin as a loading control. For ASP1, based on molecular weights, protein bands are labeled as proform (pro), intermediate form (int), and mature form (mat). (B) Immunofluorescence assay to detect T. gondii ASP1 (green) and CPB (red) in RH, RΔcpl, and RΔcplCPL tachyzoites. Arrowheads show localization of ASP1 with CPB in the VAC. Scale bars, 2 μm.
Although the basis for this intermediate phenotype remains to be determined, the findings indicate that turnover of autophagosomes is not exacerbated by deletion of both ASP1 and CPL. Taken together, our results suggest that, similar to CPB, ASP1 does not play a major role in the proteolytic turnover of autophagic material in the VAC in bradyzoites, whereas CPL has a dominant role in this activity.

**CPL plays a principal role in bradyzoite viability.** To quantify the extent to which CPL, CPB, and ASP1 contribute to bradyzoite viability, we measured the ability of bradyzoites lacking these proteases to invade, differentiate to tachyzoites, and replicate within HFF monolayers, which, if left undisturbed, results in the formation of plaques (Fig. 8A). Because proteolytic deficiency in the VAC is associated with accumulation of puncta and no increases in puncta were observed for PΔcpb or PΔasp1 between 7 and 14 days of differentiation, we only measured bradyzoite viability at 7 days of differentiation. As expected, PΔcpl bradyzoites showed a marked loss of viability based on plaque formation (Fig. 8B and C). In contrast, viability of PΔcpb and PΔasp1 bradyzoites
**FIG 7** *T. gondii* ASP1 has a limited role in protein turnover of autophagosomes. (A) The rate of parasite differentiation from the tachyzoite stage to the bradyzoite stage was determined *in vitro* by assessment of the tachyzoite-specific antigen SAG1 and GFP under the control of a bradyzoite LDH2 promoter. Infected monolayers were cultured for the indicated days, fixed, probed for SAG1, and quantified. Bars indicate means ± standard deviations from 3 biological replicates. Experiments analyzed 105, 106, 153, and 151 cysts for Pru, 75, 108, 155, and 130 cysts for PΔcpl, 119, 124, 189, and 146 cysts for PΔasp1 and 78, 110, 112, and 161 cysts for PΔasp1Δcpl on days 1, 2, 3, and 4, respectively. All strains were compared, and only statistical significance is shown. Unpaired two-tailed *t* test with Welch’s corrections was performed on the means across 3 biological replicates. (B) Phase-contrast microscopy was used to image bradyzoites *in vitro* following 7- and 14-day culture under differentiation conditions. Enlarged dark puncta indicative of defective protein degradation in VACs are visible in PΔcpl bradyzoites and PΔasp1Δcpl cysts (arrows). Scale bars, 5 μm. (C) The size of the punctae in bradyzoites was measured. The following numbers of cysts across 3 biological replicates were used to analyze each strain: Pru (day 7, 109 cysts; day 14, 98 cysts), PΔcpl (day 7, 105 cysts; day 14, 97 cysts), PΔasp1 (day 7, 85 cysts; day 14, 97 cysts), PΔasp1Δcpl (day 7, 90 cysts; day 14, 97 cysts). Lines represent medians with 95% confidence intervals. For this data set, ROUT with a Q value of 0.1% was used to identify and remove 1 outlier from PΔasp1 for 7 days postdifferentiation and 1 outlier each for Pru and PΔcpl for 14 days postdifferentiation. Since the data did not fit a normal distribution, we used a nonparametric Kruskal-Wallis test with Dunn’s multiple comparisons to compare the medians of data combined from 3 biological replicates. All strains were compared, and only significant differences are shown.
was no different from that of Pru. Moreover, the decreased viability of $P_{Δasp1Δcpl}$ bradyzoites was no greater than that of $P_{Δcpl}$ bradyzoites compared to that of Pru bradyzoites. Together, these findings suggest that whereas CPL is a key enzyme in bradyzoites, CPB and ASP1 do not play a substantial role in bradyzoite survival.

**DISCUSSION**

*Toxoplasma gondii* cathepsin L (CPL) was previously shown to have a role in protein turnover within the parasite VAC (19, 26). CPL is also necessary for the maturation of *T. gondii* CPB (23). Extending these findings here, we demonstrate that CPB and CPL colocalize in chronic-stage bradyzoites. Accordingly, it was necessary to assess whether a block in protein turnover in the VAC of CPL-deficient parasites was actually due to a
lack of active CPB. Using ingestion assays to assess their contribution to protein digestion in the VAC in tachyzoites, we found that those lacking the \textit{cpb} gene showed the accumulation of host-derived mCherry comparable to that of the Pru parental strain. Similarly, we found comparable punctae in \textit{Δcpb} and Pru bradyzoites, suggesting that the observed effect of protein accumulation in the VAC of CPL-deficient tachyzoites and bradyzoites is not due to a defect in CPB maturation.

Interestingly, we also found that an aspartyl protease (ASP1), which also occupies the VAC, was not necessary for protein degradation in the VAC or for parasite viability. Further, although ASP1 abundance increases in CPL-deficient parasites, this is apparently due to a lack of its maturation rather than a compensatory mechanism. That parasites lacking both ASP1 and CPL are no worse off than those lacking CPL alone essentially rules out compensation by ASP1 in the absence of CPL. Transcriptomic data sets show that CPL, CPB, and ASP1 each are upregulated in chronic-stage bradyzoites (30) compared to acute-stage tachyzoites, which is further supported by our immunoblotting results for CPB and ASP1 in tachyzoites and bradyzoites. Although we found no evidence to suggest CPB and ASP1 are necessary for the turnover of proteins in the VAC in bradyzoites, their increased expression in bradyzoites suggests they have another role in chronic infection. Despite this, we found that neither CPB nor ASP1 is essential for bradyzoite survival. This is in stark contrast to CPL, reinforcing the notion that CPL is the major protease of the VAC in \textit{T. gondii} bradyzoites.

The nonredundant functions between CPL and other VAC proteases in \textit{T. gondii} are unlike what has been seen in other eukaryotic organisms previously described. In some other systems cathepsin L is dispensable. For example, in murine models it is only when cathepsins B and L both are deleted from the genome that a strong detrimental phenotype is observed (6, 9).

Cathepsin D is the major aspartyl protease in the mammalian endolysosomal system and is essential for mouse survival (10). This is in contrast to what has been observed in apicomplexans (previously in \textit{P. falciparum} and reported here in \textit{T. gondii}), whereby no single aspartyl protease resident within lysosome-like compartments has been shown to be essential to parasite fitness (18). Although not essential for survival, a quadruple knockout of all four FV plasmepsins in \textit{P. falciparum} resulted in a slower growth rate, delayed schizont maturation, and reduced hemozoin formation (31, 32). This differs from our findings in the related parasite \textit{T. gondii}, whereby no defect was observed in stage differentiation and protein turnover in \textit{Δasp1} parasites. It also has been shown here and elsewhere that ASP1 is dispensable for tachyzoite growth (28). Whether ASP1 functions exclusively in the VAC or also plays a secondary role in daughter cell formation associated with the IMC remains unclear. We could not independently validate the association of ASP1 with the IMC in forming daughter cells, because the antibody was insufficiently specific when used to probe replicating parasites despite its specificity for ASP1 in newly invaded parasites (Fig. 4B and Fig. S3E).

Limiting parasite access to essential nutrients by blocking protein turnover has become a major focus of anti-parasite drug development (33–35). In \textit{P. falciparum}, the hydrolysis of hemoglobin involves the cooperative action of two classes of proteases. In addition to their direct role in protein degradation, falcipains have an indirect role via activation of plasmepsins; however, autoprocessing of plasmepsins has also been observed following falcipain disruption (15). This has implications for the development of inhibitors against \textit{P. falciparum} proteases involved in protein degradation in the FV, as they will have to block the function of multiple cathepsins simultaneously. Our work reveals a salient aspect of the \textit{Toxoplasma} endolysosomal system in that, unlike \textit{Plasmodium}, there exists an opportunity to target a single nonredundant protease, CPL.

Although our findings suggest noncritical roles for CPB and ASP1 \textit{in vitro}, the contributions of these proteases to infection \textit{in vivo} have been only partly explored. For example, antisense RNA inhibition of CPB expression or treatment with a protease inhibitor significantly decreased \textit{T. gondii} infection of chicken embryos (36). However, possible off-target effects from antisense inhibition or treatment are difficult to rule out in such studies. Although targeted deletion of ASP1 in the RH strain caused lethal
infection of mice with the same kinetics as those for wild-type parasites (28), the high virulence of the RH strain often renders it difficult to measure moderate contributions to infection. Additional studies are necessary to better define the role of CPB and ASP1 during in vivo infection.

*T. gondii* chronically infects one-third of the worldwide population (1). Despite drug development against the infection being an active area of research, there is currently no effective treatment for chronic toxoplasmosis. Although previous work has focused on developing inhibitors against more than one cathepsin protease in *T. gondii* (37), it is now clear that CPL can be the focus of future efforts targeting disruption of VAC proteolysis. Of particular interest is that the development of compounds that specifically inhibit *T. gondii* CPL might not have severe side effects resulting from inhibition of host cathepsin L (34).

**MATERIALS AND METHODS**

Para*site* cell culture and differentiation into bradyzoites. Throughout this investigation, human foreskin fibroblast (HFF) cells were used to propagate parasite strains in Dulbecco's modified Eagle medium (DMEM) supplemented with 10% (vol/vol) cosmic calf serum (CCS) or fetal bovine serum (FBS).

For all bradyzoite conversions, tachyzoites were mechanically lysed by scraping infected HFF monolayers that then were passed through 20- and 23-gauge syringes and a 3-μm filter. Filtered parasites then were counted and allowed to infect fresh monolayers of HFF cells for 24 h. Bradyzoite differentiation was induced using alkaline pH medium and ambient CO₂ (38–40). Briefly, 24 h after parasites were applied to HFF monolayers, DMEM was replaced with an alkaline differentiation medium (RPMI without NaHCO₃, 50 mM HEPES, penicillin-streptomycin, 3% FBS, pH 8.25). Differentiation medium was replaced daily.

Generation of transgenic *T. gondii* strains. Strategy, vectors, and repair templates employed to create all the transgenic strains used in this study are described in the supplemental material. The transfection of parasites was performed as described previously (41). Briefly, 2 × 10⁷ tachyzoites were harvested, washed twice in cytosome (120 mM KCl, 0.15 mM CaCl₂, 10 mM K₂HPO₄-KH₂PO₄, [pH 7.6], 25 mM HEPES, pH 7.6, 2 mM EGTA, 5 mM MgCl₂, 2 mM ATP, 5 mM glutathione), and resuspended in 700 μl of cytosome containing either 5 μg of PCR repair template or 50 μg of plasmid. The entire mixture was then transferred to an electroporation cuvette (4-mm gap) and exposed to an electric pulse with an electroporator (BTX 600) set at 2.0 kV and 48 ohms. Electroporated parasites were transferred to fresh HFF monolayers and subjected to drug selection. Clones were isolated by limiting dilution.

Expression and purification of recombinant proform ASP1. An ~1.5-kb DNA fragment encoding amino acids 142 to 610 of the recombinant proform of ASP1 (proASP1) was PCR amplified from a *T. gondii* RH cDNA library using Q5 high-fidelity DNA polymerase (NEB) and cloned into a pET22b vector with a 6× His epitope tag fused to the N-terminal end of ASP1. This lacks the first 141 amino acids of ASP1, corresponding to the cytosolic N-terminal domain and transmembrane domain, which could affect protein solubility. The resulting pET22b-6× His-ASP1 expression construct was transformed into the *Escherichia coli* protein expression strain, ER2566 (NEB), induced with 1 mM isopropyl 1-thio-β-D-galactopyranoside (IPTG) at 37°C for 4 h, and purified using a standard purification procedure as described previously (25). The purified denatured ASP1 protein was refolded in phosphate-buffered saline (PBS) overnight by dialysis. Insoluble protein was removed by centrifugation at 20,000 × g for 10 min, and the soluble fraction was concentrated using an Amicon Ultra 30-kDa-cutoff concentrator (Millipore) and stored at ~20°C for future use.

Immunization and affinity purification of anti-ASP1. Two New Zealand White rabbits were injected with 200 μg of purified recombinant proASP1 in Freund’s complete adjuvant and boosted 3 times at days 14, 21, and 56 with 50 μg of recombinant protein in Freund’s incomplete adjuvant (Cocalico Biologicals, Inc.). The final sera were collected and further affinity purified. Purified recombinant proASP1 was cross-linked to 0.5 ml of AminoLink coupling resin (Thermo Fisher), and affinity-purified anti-ASP1 antibodies were eluted with 0.2 M glycine, pH 3.0. The elution fractions containing the antibodies were pooled, buffer exchanged, and concentrated by using Amicon Ultra 30-kDa concentrators.

Immunoblotting. Toxoplasma gondii parasites were purified as described above, counted using a hemocytometer, and pelleted by centrifugation at 1,800 × g for 10 min. Parasite pellets were resuspended in SDS-PAGE sample buffer containing 5 mM dithiothreitol and heated at 95°C for 5 min. Parasite proteins were separated by SDS-PAGE on 15% polyacrylamide gels and then transferred to an Immobilon-P polyvinylidene difluoride membrane using semidy electroblotting. Blots were blocked using 0.1% (vol/vol) Tween 20 in PBS (PBST) containing 5% (wt/vol) nonfat dried milk. Blots were probed with the primary antibodies rabbit anti-CPL (1:8,000), mouse anti-CPB (1:2,000), rabbit anti-CPL (1:8,000), affinity-purified rabbit anti-ASP1 (1:2,500), and mouse anti-MIC2 (1:5,000). Secondary horseradish peroxidase-conjugated goat-anti-mouse/rabbit IgG (Jackson Immunoresearch) diluted 1:10,000 in PBST, was applied. Bound antibody was visualized with the chemiluminescence substrate SuperSignal West Pico (Thermo Scientific) and imaged using a GBOX Chemi XRQ imager (Syngene). Fluorescence immunoblots were as described above, except they were blocked in 50 mM Tris-HEC1 pH 7.4, 1.25% (vol/vol) fish gelatin, 150 mM NaCl for 30 min, and secondary antibodies were IRDye 680CW-conjugated goat anti-rabbit IgG conjugated and IRDye 800CW-conjugated goat-anti mouse IgG conjugated. Blots were air-dried before being imaged by a Li-COR Odyssey CLx instrument.
Immunofluorescence assay. To confirm protease localization in tachyzoites, indirect immunofluorescence was performed. Parasites were allowed to infect HFF cells for 30 min in 8-well chamber slides at 37°C, 5% CO₂. Infected cells then were fixed using 4% (vol/vol) methanol-free formaldehyde in PBS. Cells were permeabilized with 0.1% (vol/vol) Triton X-100 for 10 min, followed by 20 min of blocking with PBS containing 10% FBS. Blocked monolayers were incubated at room temperature (RT) for 40 min with the primary antibody mouse anti-CPB (1:200), rabbit anti-CPL (1:500), or affinity-purified rabbit anti-ASP1 (1:200). Following washes in PBS–1% FBS–1% normal goat serum, slides were probed with secondary fluorophore-conjugated antibodies (Jackson Immunoresearch), mounted using Mowiol, and imaged on a Zeiss Axio inverted fluorescence microscope equipped with a Zeiss AxioCam 305 scMOS digital camera.

For indirect immunofluorescence on bradyzoites, freshly lysed parasites were placed onto an HFF cell monolayer and left at 37°C and 5% CO₂ overnight. Invaded parasites were subsequently stimulated to differentiate into bradyzoites by culturing in alkaline differentiation media and incubating at 37°C and 0% CO₂ for 7 days. Bradyzoite cultures were fixed, probed with antibodies, and mounted as described above.

Parasite ingestion assay. To assess the degradation of proteinaceous material within the lysosome-like VAC in tachyzoites, the previously described ingestion assay was used (42). Briefly, modified CHO-K1 cells were induced to express cytosolic mCherry following treatment with doxycycline (2 μg/ml) for 5 days prior to infection. On the sixth day, freshly harvested parasites were allowed to infect and ingest cytosolic mCherry from doxycycline-treated CHO-K1 cells for 4 h. After incubating, parasites were harvested from these cells. Purified parasitophorous vacuoles were treated at 12°C for 1 h with freshly prepared 1 mg/ml pronase (Roche)–0.01% saponin–PBS before washing, placing them on slides precoated with Cell-Tak (Corning), and fixing with 4% formaldehyde in PBS (20 min at RT). Fixed parasites were washed three times with PBS and permeabilized by incubation with PBS with 0.1% Triton X-100 for 10 min at RT. Slides then were mounted with Mowiol and imaged as described above. Samples were coded and enumerated blindly.

Tachyzoite growth assays. To determine the contributions of CPB, CPL, and ASP1 to intracellular tachyzoite growth, the relative growth rate of Pru and each knockout strain was compared based on the number of parasites per parasitophorous vacuole 24 and 48 h postinvasion. Freshly harvested tachyzoites were placed onto a monolayer of confluent HFF cells grown on coverslips in a 6-well plate containing DMEM supplemented with 10% CCS. Parasites were allowed to invade for 1 h, at which point parasites that had not invaded were washed away with DMEM supplemented with 10% CCS. Parasites were left to grow for a further 24 and 48 h before washing with PBS three times and fixing with 4% (vol/vol) methanol-free formaldehyde in PBS. Cells were permeabilized, blocked, and washed as described above. To visualize individual parasites for counting, the prepared slides were incubated for 40 min at RT with rat anti-SAG1 antibodies, which were generated by a commercial service (Pierce) using baculovirus-derived recombinant SAG1 (kindly provided by Martin Boulanger), followed by secondary Alexa-S94-conjugated anti-rat antibody (Jackson Immunoresearch). The number of parasites per parasitophorous vacuole was counted for each strain.

Puncta assays. To assess the formation of puncta (accumulation of dense material) in each strain, we followed the puncta quantification assay described previously (26). Briefly, HFF cell monolayers on coverslips were infected with T. gondii tachyzoites and differentiated to bradyzoites over the course of 7 days (as described above). For direct immunofluorescence of bradyzoite cyst walls, fluorescein-labeled Dolichos biflorus agglutinin (DBA; Vector Labs) was diluted 1:400 and incubated with infected monolayers that had been fixed and made permeable (as described above). The fluorescent signal derived from DBA staining allowed for automatic detection of the bradyzoite-containing cyst area using ImageJ software. Cyst images were captured with a 100× oil objective as described above. Automatic bradyzoite cyst identification and puncta quantification were performed in ImageJ using the following parameters, which were described previously (26). Maximum entropy thresholding on the GFP channel was used to identify cysts. This was followed by the identification of objects with areas between 130 and 1,900 μm². Particles (punctae) under the GFP mask (and, therefore, within a cyst) were analyzed by automatic local thresholding on the phase image using the Phansalkar method, with the following parameters: radius, 5.00 μm; k value, 0.5; r value, 0.5. Puncta were measured from the resulting binary mask by particle analysis according to the following parameters: size, 0.3 to 5.00 μm; circularity, 0.50 to 1.00.

Bradyzoite differentiation assay. Immunofluorescence assays were performed to determine the relative rate at which parasites differentiated from the tachyzoite stage to the bradyzoite stage, based on the amount of tachyzoite-specific SAG1 and bradyzoite-specific GFP present within parasitophorous vacuoles/developing cysts. Freshly lysed tachyzoites were placed onto confluent HFF monolayers on coverslips (in 6-well plates) and left to incubate at 37°C and 5% CO₂ for 24 h. Following this, the medium was removed from each well and alkaline medium was added, along with placement in incubators lacking CO₂ to induce bradyzoite conversion (still maintained at 37°C). Coverslips were fixed and stained with the tachyzoite-specific marker rat anti-SAG1 (1:1,000) following 1, 2, 3, and 4 days of differentiation. Strains used in this study express GFP under the bradyzoite-specific LDH2 promoter, and this is used as a marker for bradyzoites. Cysts that displayed 50% or more staining by either rat anti-SAG1 antibodies or GFP were considered positive.

Bradyzoite viability assays. Bradyzoite viability was assessed by combining plaque assay and quantitative PCR (qPCR) analysis of genome number, as previously described (26). Briefly, HFF cell monolayers were infected with T. gondii tachyzoites in 6-well plates. Following host cell invasion, tachyzoites underwent differentiation to bradyzoites as described above, resulting in the generation of intracellular cysts. Differentiation was carried out over the course of 7 days, replacing the alkaline medium daily. Following differentiation, the culture medium in each well was replaced with 2 ml Hanks...
balanced salt solution, and cysts were liberated from the infected HFF monolayers by mechanical
extraction by lifting cells with a cell scraper and syringing several times through 25-gauge needles. In the
next step, 2 ml of prewarmed 2× pepsin solution (0.026% pepsin in 170 mM NaCl and 60 mM HCl, final
concentration) was added to each sample, and samples were left to incubate at 37°C for 30 min.
Reactions were stopped by adding 94 mM Na2CO3, removing the supernatant after centrifugation at
1,500 × g for 10 min at RT, and resuspending pepsin-treated parasites in 1 ml of DMEM without serum.
Parasites were enumerated, and 1,000 parasites per well were added to 6-well plates containing
confluent monolayers of HFFs in D10 medium in triplicate. To allow for the formation of bradyzoite
plaques, parasites were left to grow undisturbed for 10 days. After this period, the number of plaques in
confluent monolayers of HFFs in D10 medium in triplicate. To allow for the formation of bradyzoite
Parasites were enumerated, and 1,000 parasites per well were added to 6-well plates containing
in centimeter (diameter) was normalized to the parental control, and this normalized
value was used for statistical analysis.

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.
FIG S1, TIF file, 0.5 MB.
FIG S2, TIF file, 0.3 MB.
FIG S3, TIF file, 1 MB.

ACKNOWLEDGMENTS
We thank Aric Schultz for help with the automated puncta analysis, Martin Bou-
langer for providing recombinant SAG1, and My-Hang (Mae) Huynh for proofreading the
manuscript prior to submission.

This work was supported by National Institutes of Health grant R01AI120627 (to V.B.C. and M.D.C.), R01AI143707 (to Z.D.), and P20GM109094 (to Z.D.). Funding was also
provided by grants from the Stanley Medical Research Institute (to V.B.C.) and the
Knights Templar Eye Foundation (to Z.D.), along with support from the University of
Perugia Fondo Ricerca di Base 2018 program of the Department of Chemistry, Biology, and
Biotechnology (to M.D.C.).

We declare that we have no conflicts of interest concerning the contents of this
article.

V.B.C., M.D.C., and Z.D. conceived the study. C.M., D.S., and Z.D. performed most of
the experiments and analyzed the data. M.D.C. made and confirmed by PCR PΔcpb,
PΔasp1, and PΔasp1PΔcp1 strains. G.K. performed the ingestion experiments, with some
assistance from C.M., and made bradyzoite lysates for immunoblots. C.M., D.S., Z.D.,
M.C.D., and V.B.C. made the figures. C.M., D.S., Z.D., and V.B.C. wrote the paper, and all
authors revised the paper.

REFERENCES
1. Montoya J, Liesenfeld O. 2004. Toxoplasmosis. Lancet 363:1965–1976.
https://doi.org/10.1016/S0140-6736(04)67361-6.
2. Siqueira-Neto JL, Debnath A, McCaffrey L, Bernatchez JA, Ndao M, Reed SL, Rosenthal PJ. 2018. Cysteine proteases in protozoan parasites. PLoS Negl Trop Dis 12:e0006512. https://doi.org/10.1371/journal.pntd.0006512.
3. Robbins JR, Zeldovich VB, Poukchanski A, Boothroyd JC, Bakardjiev AL. 2012. Tissue barriers of the human placenta to infection with Toxoplasma gondii. Infect Immun 80:418–428. https://doi.org/10.1128/IAI.05899-11.
4. Stoka V, Turk V, Turk B. 2016. Lysoosomal cathepsins and their regulation in aging and neurodegeneration. Ageing Res Rev 32:22–37. https://doi.org/10.1016/j.arr.2016.04.010.
5. Neurath H, Walsh KA. 1976. Role of proteolytic enzymes in biological regulation (a review). Proc Natl Acad Sci U S A 73:3825–3832. https://doi.org/10.1073/pnas.73.11.3825.
6. Felbo B, Kessler B, Mothes W, Goebel HH, Ploegh HL, Bronson RT, Olsen BR. 2002. Neuronal loss and brain atrophy in mice lacking cathepsins B and L. Proc Natl Acad Sci U S A 99:7883–7888. https://doi.org/10.1073/pnas.112632299.
7. Den nemärker J, Lohmüller T, Müller S, Aguilar SV, Tobin DJ, Peters C, Rein heckel T. 2010. Impaired turnover of autophagolysosomes in cathepsin L deficiency. Biol Chem 391:913–922. https://doi.org/10.1515/bc .2010.097.
8. Halangk W, Lerch WM, Brandt-Nedelev B, Roth W, Ruthenbuerg M, Rein-heckel T, Domschke W, Lippert H, Peters C, Deussing J. 2000. Role of cathepsin B in intracellular trypsinogen activation and the onset of acute pancreatitis. J Clin Invest 106:773–781. https://doi.org/10.1172/JCI9411.
9. Sevenich L, Pennacchio LA, Peters C, Reinheckel T. 2006. Human cathepsin L rescues the neurodegeneration and lethality in cathepsin B double-deficient mice. Biol Chem 387:885–891. https://doi.org/10.1515/ BC.2006.112.
10. Saftig P, Hetman M, Schmidt-Walter K, Heine L, Messmann H, Köster A, Hess B, Evers M, von Figura K. 1995. Mice deficient for the lysozymal proteinase cathepsin D exhibit progressive atrophy of the intestinal
mucoza and profound destruction of lymphoid cells. EMBO J 14: 3590–3608. https://doi.org/10.1092/1400-2705.1995.tb0029.x

11. Aghdassi AA, John DS, Senderl M, Weiss FU, Reineckel T, Mayerle J, Lerch MM. 2018. Cathepsin D regulates cathepsin B activation and disease severity predominantly in inflammatory cells during experimental pancreatitis. J Biol Chem 293:1018–1029. https://doi.org/10.1074/jbc.M117.184772.

12. Sijwali PS, Kato K, Seydel KB, Gut J, Lehman J, Klemba M, Goldberg DE, Miller LH, Rosenthal PJ. 2004. Plasmodium falciparum cysteine protease falcipain-1 is not essential in erythrocystic stage malaria parasites. Proc Natl Acad Sci U S A 101:8721–8726. https://doi.org/10.1073/pnas.0402738101.

13. Rosenthal PJ. 2011. Falcipains and other cysteine proteases of malaria parasites, p 30–48. In Robinson MW, Dalton JP (ed), Cysteine proteases and their suitability as drug targets. PLoS Negl Trop Dis 12:e0005639. https://doi.org/10.1371/journal.pntd.0005639.

14. Banerjee R, Liu J, Beatty W, Pelosof L, Klemba M, Goldberg DE. 2002. Four aspartic proteases and a novel, dynamic and cell-cycle-dependent gene disruption confirms a critical role for the cysteine protease falcipain-1 in Plasmodium falciparum. J Biol Chem 278:12870–12876. https://doi.org/10.1074/jbc.M708942900.

15. Ersmark K, Samuelsson B, Hallberg A. 2006. Plasmepsins as potential targets for new antimarial therapy. Med Res Rev 26:626–666. https://doi.org/10.1002/med.20082.

16. Delauw M-F, Weiss LM, Bzik DJ. 2004. Toxoplasma-induced changes in host risk behaviour are independent of parasite-derived AaaH2 tyrosine hydroxylase. Sci Rep 7:1–14. https://doi.org/10.1038/s41598-017-13229-y.

17. Afonso C, Paixão VB, Klaus A, Lunghi M, Piro F, Emiliani C, Di Cristina M, Cost A, Lachwa J, Prats D, Fünkhauser C, Nikolic P, Coelho E, Haas FD, Souza WD, Coppens I, Sibley LD, Moreno SNJ. 2010. Cathepsin proteases in Toxoplasma gondii, p 30–48. In Robinson MW, Dalton JP (ed), Cysteine proteases and their suitability as drug targets. PLoS Negl Trop Dis 12:e0005639. https://doi.org/10.1371/journal.pntd.0005639.

18. Bonilla JA, Bonilla TD, Yowell CA, Fujioka H, Dame JB. 2007. Critical roles of digestive vacuole plasmepsins of Plasmodium falciparum in vacuolar function. Mol Microbiol 65:64–75. https://doi.org/10.1111/j.1365-2958.2007.07568.x.

19. Sijwali PS, Rosenthal PJ. 2004. Gene disruption of the cysteine protease falcipain-2 in hemoglobin hydrolysis by Plasmodium falciparum. J Biol Chem 284:26839–26850. https://doi.org/10.1074/jbc.M108942900.

20. Miranda K, Pace DA, Cintron R, Rodrigues JCF, Fang J, Smith A, Rohloff P, Coelho E, Haas FD, Souza WD, Coppens I, Sibley LD, Moreno SNJ. 2010. Plasmepsin-digestive vacuole plasmepsins in the specificity and antimalarial mode of action of cysteine and aspartic protease inhibitors. Antimicrob Agents Chemother 53:806. https://doi.org/10.1128/AAC.00882-09.

21. Soete M, Camus D, Dubremetz JF. 1994. Experimental induction of T. gondii bradyzoites. J Eukaryot Microbiol 42:150–157.

22. Cost A, Lachwa J, Prats D, Fünkhauser C, Nikolic P, Coelho E, Haas FD, Souza WD, Coppens I, Sibley LD, Moreno SNJ. 2010. Cathepsin L is the primary target of the invasion-inhibitory compound morpholinurea-leucyl-homophenyl-vinyl sulfone phenyl. J Biol Chem 288:3523–3534. https://doi.org/10.1074/jbc.M110.1443697.

23. Dupont P, Mehlman MR, Barrera R, Cost A, Lachwa J, Prats D, Fünkhauser C, Nikolic P, Coelho E, Haas FD, Souza WD, Coppens I, Sibley LD, Moreno SNJ. 2010. Cathepsin L occupies a vacuolar compartment and is a protein maturing within the endo/exocytic system of Toxoplasma gondii. Mol Microbiol 76:1340–1357. https://doi.org/10.1111/j.1365-2958.2010.07165.x.

24. Bonilla JA, Bonilla TD, Yowell CA, Fujioka H, Dame JB. 2007. Critical roles of digestive vacuole plasmepsins of Plasmodium falciparum in vacuolar function. Mol Microbiol 65:64–75. https://doi.org/10.1111/j.1365-2958.2007.07568.x.

25. Afonso C, Paixão VB, Klaus A, Lunghi M, Piro F, Emiliani C, Di Cristina M, Cost A, Lachwa J, Prats D, Fünkhauser C, Nikolic P, Coelho E, Haas FD, Souza WD, Coppens I, Sibley LD, Moreno SNJ. 2010. Cathepsin L is the primary target of the invasion-inhibitory compound morpholinurea-leucyl-homophenyl-vinyl sulfone phenyl. J Biol Chem 288:3523–3534. https://doi.org/10.1074/jbc.M110.1443697.

26. Di Cristina M, Dou Z, Lunghi M, Kannan G, Hyun M-H, McGovern OL, Schulz TL, Schulz AJ, Miller AJ, Hayes BM, van der Linden W, Emiliani C, Bogyo M, Besteiro S, Coppens I, Carruthers VB. 2017. Toxoplasma depends on lysosomal consumption of autopshagosomes for persistent infection. Nat Microbiol 2:17096. https://doi.org/10.1038/nmicrobiol.2017.96.

27. Shea M, Jäkle U, Liu Q, Berry C, Joiner KA, Soldati-Favre D. 2007. A family of aspartic proteases and a novel, dynamic and cell-cycle-dependent protease localization in the secretory pathway of Toxoplasma gondii. Traffic 8:1018–1034. https://doi.org/10.1111/j.1600-0854.2007.00589.x.

28. Polonais V, Shea M, Soldati-Favre D. 2011. Toxoplasma gondii aspartic protease 1 is not essential in tachyzoites. Exp Parasitol 128:454–459. https://doi.org/10.1016/j.exppara.2011.05.003.

29. Couplin M, Fidder H, Lu L, Tomita T, Gligor J, Mercier C, Cesbron-Delauw M-F, Weiss LM, Bzik DJ. 2011. II Type II Toxoplasma gondii KU80 knockout strains enable functional analysis of genes required for cyst development and latent infection. Eur J Cell Biol 10:1193–1206. https://doi.org/10.1002/ecc.20097-10.

30. Pittman KJ, Alloia MT, Knoll LJ. 2014. Dual transcriptional profiling of mice and Toxoplasma gondii during acute and chronic infection. BMC Genomics 15:806. https://doi.org/10.1186/1471-2164-15-806.

31. Ersmark K, Samuelsson B, Hallberg A. 2006. Plasmepsins as potential targets for new antimarial therapy. Med Res Rev 26:626–666. https://doi.org/10.1002/med.20082.

32. Aghdassi AA, John DS, Senderl M, Weiss FU, Reineckel T, Mayerle J, Lerch MM. 2018. Cathepsin D regulates cathepsin B activation and disease severity predominantly in inflammatory cells during experimental pancreatitis. J Biol Chem 293:1018–1029. https://doi.org/10.1074/jbc.M117.184772.

33. Rothenberg PJ. 2011. Falcipains and other cysteine proteases of malaria parasites, p 30–48. In Robinson MW, Dalton JP (ed), Cysteine proteases and their suitability as drug targets. PLoS Negl Trop Dis 12:e0005639. https://doi.org/10.1371/journal.pntd.0005639.

34. Rothenberg PJ. 2018. Update on drug development targeting parasite cysteine proteases. PLoS Negl Trop Dis 12:e0005850. https://doi.org/10.1371/journal.pntd.0005850.

35. Dou Z, McGovern OL, Di Cristina M, Carruthers VB. 2014. Toxoplasma gondii ingests and digests host cytosolic proteins. mBio 5:e01188-14. https://doi.org/10.1128/mBio.01188-14.