Toxoplasma gondii serine hydrolases regulate parasite lipid mobilization during growth and replication within the host

Highlights

- *T. gondii* serine hydrolases with similar catalytic folds prefer different substrates
- *T. gondii* ASH proteins are lipid-metabolizing enzymes
- *T. gondii* lipid metabolism pathways use enzymes that are viable therapeutic targets

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In brief

The intracellular parasite *Toxoplasma gondii* relies on lipid scavenging and processing to infect its mammalian hosts. In this study, Onguka et al. characterize a set of enzymes important for lipid metabolism and identify small-molecule inhibitors that perturb lipid regulation in the parasite.
Toxoplasma gondii serine hydrolases regulate parasite lipid mobilization during growth and replication within the host

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https://doi.org/10.1016/j.chembiol.2021.05.001

SUMMARY

The intracellular protozoan parasite Toxoplasma gondii must scavenge cholesterol and other lipids from the host to facilitate intracellular growth and replication. Enzymes responsible for neutral lipid synthesis have previously been thought to be depalmitoylating enzymes. We find they do not cleave palmitoyl thiol esters but rather hydrolyze short-chain lipid esters. Deletion of one of the hydrolases results in alterations in levels of multiple lipids species. We also identify small-molecule inhibitors of these hydrolases and show that treatment of parasites results in phenotypic defects reminiscent of parasites exposed to excess cholesterol or oleic acid. Together, these data characterize enzymes necessary for processing lipids critical for infection and highlight the potential for targeting parasite hydrolases for therapeutic applications.

INTRODUCTION

The phylum Apicomplexa contains multiple human parasites, including Plasmodium falciparum and Toxoplasma gondii. T. gondii, which is an obligate intracellular parasite, asymptptomatically infects approximately 30% of the world’s population (Black and Boothroyd, 2000; Boothroyd, 2009; Robert-Gangneux and Darde, 2012). Infection of immuno-compromised individuals, such as HIV patients, chemotherapy recipients, and organ transplant recipients can result in toxoplasmosis (Montoya and Liesenfeld, 2004). In infants, toxoplasmosis can result in seizures, enlarged liver, enlarged spleen, jaundice, and blindness. Unfortunately, current treatments (pyrimethamine, sulfadiazine, clindamycin, trimethoprim-sulfamethoxazole) for toxoplasmosis are not effective against acute infections and have side effects that include teratogenic or embryocidal effects, decrease in fertility, and fatal “gasping syndrome” in premature infants (Montoya and Liesenfeld, 2004). For these reasons, a greater understanding of the factors that regulate the T. gondii asexual lytic cycle has the potential to provide valuable new avenues for therapeutic development for toxoplasmosis.

Previously, our laboratory used a chemical genetic screen to identify a T. gondii serine hydrolase (TgPPT1) as a depalmitoylating enzyme (Child et al., 2013; Hall et al., 2011). This approach utilized a small-molecule library that contained ~1,200 compounds designed to inhibit cysteine and serine hydrolases (Ara-stu-Kapur et al., 2008). Inhibition of TgPPT1 resulted in increased host cell invasion as a result of increased parasite motility. However, deletion of the TgPPT1 gene resulted in viable parasites, suggesting that other depalmitoylating enzymes are likely to exist that may be able to compensate for loss of TgPPT1 (Kemp et al., 2013). In further support of this assessment, a homology search by hidden Markov modeling identified three putative depalmitoylase enzymes in T. gondii. These serine hydrolases were named active serine hydrolases Tg(ASH)2, 3, and 4 (TgPPT1 is TgASH1) and were selected based on the presence of an α/β-hydrolase domain and clear sequence similarity to TgPPT1 (Kemp et al., 2013). However, genetic deletion studies provided little insight into the function of these enzymes. While deletion of TgASH2 or TgASH3 did not affect growth or infectivity (Kemp et al., 2013), deletion of TgASH4 caused specific defects in formation of ordered rosettes inside the parasitophorous vacuole, and subsequent inability of parasites to disperse from the site of egress (Foe et al., 2018). In addition, in vitro characterization of the purified TgASH4 enzyme demonstrated that it prefers to process short acyl chains (acetate), suggesting that it is likely not a depalmitoylase. Interestingly, the TgASH4 deletion
phenotype is similar to phenotypes observed when parasites are grown under conditions of excess cholesterol or excess oleic acid (Nolan et al., 2017, 2018), suggesting that TgASH4 and possibly TgASH2 and TgASH3 might play a role in lipid metabolism in T. gondii.

T. gondii encodes three genes that produce enzymes involved in neutral lipid synthesis. Two enzymes are responsible for generating cholesterol esters (acyl-CoA:cholesterol acyltransferases [TgACAT1 and 2]) (Lige et al., 2013; Nishikawa et al., 2005; Sonda et al., 2001) and one enzyme generates triacylglycerides (acyl-CoA:diacylglycerol acyltransferase [TgDGAT1] (Quittnat et al., 2004), which are both major components of lipid droplets. Deletion of TgDGAT is lethal (Quittnat et al., 2004), and deletion of either TgACAT1 or 2 results in severe growth defects (Nishikawa et al., 2005), while deletion of both is lethal (Lige et al., 2013; Nishikawa et al., 2005). These results suggest that the pathway for neutral lipid generation is essential for parasite survival. Although the importance of neutral lipid synthesis has been demonstrated for parasite growth and replication, little is known about how parasites utilize scavenged lipids during their intracellular growth.

The T. gondii genome contains several genes annotated as putative lipases and carboxy-esterases whose functions have not been experimentally verified. Several or all of these enzymes could be bona fide acyl-triglyceride lipase(s), even though they lack clear homology with the known mammalian or yeast lipases involved in acyl-triglyceride metabolism. Furthermore, how and when the parasite utilizes scavenged lipids and what enzymes are required for the mobilization of stored neutral lipids is not clear. While it has been proposed that T. gondii relies on host enzymes for mobilizing scavenged lipids (Nolan et al., 2017), it remains possible that the parasite has its own internal mechanism for metabolism of these lipids.

In this study, we biochemically characterized the four previously identified ASH α/β-serine hydrolases (TgPPT1 [also known as TgASH1], TgASH2, TgASH3, and TgASH4) that are conserved across multiple phyla with no known biological function. A BLAST search for conserved motifs identified an acyltransferase motif, a calcium-independent phospho-lipase motif, a pentapeptide (GxSxG) lipase motif, and a lipid binding motif. In vitro biochemical assays confirm that the enzymes all have both esterase and thioesterase activity with different substrate chain-length preferences. Using a liquid chromatography-mass spectrometry (LC-MS) lipidomics approach, we show that TgASH4 has a role in parasite lipid metabolism. We also screened and identified inhibitors of recombinant TgASH2, 3, and 4 and show that treatment of parasites photocopies lipid defects observed in T. gondii when grown under excess cholesterol or oleic acid. Together, these results define the biochemical properties of the TgASH2, 3, and 4 enzymes, and suggest that, due to their role in lipid metabolism, they represent viable targets for treatment of T. gondii infections.

**RESULTS**

**Sequence analysis and modeling of parasite serine hydrolases**

To gain insight into the potential functions of TgASH proteins, we used sequence alignments and computational structure analysis. T. gondii ASH proteins share high protein sequence similarity with members of the α/β-hydrolase superfAMILY, specifically with serine hydrolases and serine esterases, which include the carboxylesterases, thioesterases, cholinesterases, and lipases (Long and Cravatt, 2011). Sequence alignments confirmed the presence of a conserved catalytic triad (Ser-His-Asp) in each TgASH protein that can also be found in various homologs in P. falciparum and humans (Figures 1A and S1A). This analysis also revealed the presence of a putative hydrophobic/lipid binding motif (Ghosh et al., 2009; Rajakumari and Daum, 2010) and a calcium-independent phospholipase motif (Jenkins et al., 2004) in TgASH2, TgASH3, and TgASH4. In addition, TgAH3 and TgASH4 each has an additional putative acyltransferase motif (Rajakumari and Daum, 2010).

To explore TgASH proteins in the context of serine hydrolases from humans and the related parasite pathogen P. falciparum, we performed phylogenetic analysis of putative serine hydrolase protein sequences. We performed HMMer searches of these SHs against the SwissProt database to generate a list of significant matches (sequence e < 0.01; supplemental information spreadsheet 1) in the human proteome and subsequently performed a multiple alignment of the entire set of hits with the Toxoplasma and Plasmodium sequences. This analysis indicated that TgASH2, TgASH3, and TgASH4 are more similar to each other than they are to TgPPT1 (Figures 1B and S1B). TgPPT1 clustered with known acyl-protein thioesterases LYP1 and LYP2 (Foe et al., 2015; Kemp et al., 2013), in agreement with our previous studies demonstrating its depalmitoylase activity (Child et al., 2013). Contrast, TgASH2-4 and the P. falciparum serine hydrolases P0818600 and P0805000 clustered with the α/β-hydrolase domain containing protein 17 enzymes (ABHD17A, B, and C) and with ABHD13 (ABHDD), all of which are suggested to have depalmitoylase activity. ABHD17s in particular, are thought to be responsible for depalmitoylating N-Ras, thereby regulating the dynamic cycling of Ras localization (Conibear and Davis, 2010). These observations suggested that, while TgASH2-4 also align to human proteins with depalmitoylase activity, their lack of similarity to TgPPT1 suggests that they might have distinct substrate preferences.

To gain further insight into the potential unique properties of the parasite serine hydrolases, we generated structural predictions using homology modeling. We aligned parasite proteins to similar proteins for which high-resolution crystallographic structures were available. We then selected the best templates chosen using Fold & Function Assignment (Jaroszewski et al., 2011), and generated structural models with MODELER (Sali and Blundell, 1993). For each protein, we built two structures and selected the model that best satisfied the expected geometries of the catalytic triad (Table S1). The final structural models of PPT1, ASH2, ASH3, and ASH4 predict that the catalytic residues all fall within a defined active site pocket (Figures 1C–1F and S1C). They also reveal a potential lipid binding domain that overlaps with the acyltransferase motif highlighted in the structures (Ghosh et al., 2009), located near the active site, reinforcing the assignment of the catalytic residues (Figures 1A, S1A, and S1Ci–v). Although sequence-based analyses are not sufficient to predict specific enzymatic activity or biological function, in this case they did reveal conserved hydrolase motifs in our T. gondii serine hydrolases of interest, consistent with our
Figure 1. Sequence analysis and modeling of parasite serine hydrolases

(A) Sequence alignment of TgPPT1, TgASH2, TgASH3, and TgASH4 using Clustal Omega.

(B) Dendrogram of T. gondii and P. falciparum serine hydrolases generated using the Clustal Omega algorithm. Branch length represents sequence relatedness.

(C–F) Structural models of (C) TgPPT1, (D) TgASH2, (E) TgASH3, and (F) TgASH4 showing the active site cavity surface (red volume) generated using PyMOL. See also Figure S1.
Figure 2. Purified TgASH enzymes exhibit esterase and thioesterase activities

(A–C) (A) Structures of the quenched fluorogenic substrate for esterase (QSE) and thioesterase (QStE) activities for measuring depalmitoylase activity. Quantification of the rate of hydrolysis of (B) QStE and (C) QSE by recombinant TgPPT1, TgASH2, TgASH3, and TgASH4 enzymes. Data represent the mean ± standard deviation of three independent experiments from three technical triplicates.

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hypothesis that these enzymes share a common catalytic mechanism. Furthermore, phylogenetic clustering suggests that TgASH2-4 and TgPPT1 have distinct biochemical functions (Figures 1B and S1B). To confirm this hypothesis, the best models for TgASH2-4 were aligned to TgPPT1 using PyMOL “super” (Ho and Gruswitz, 2008). This analysis allows prediction of the size of the active site “cavity and pockets” and correlate that to the size of the substrate that would optimally bind. Based on this structural analysis, we observe that the size/volume of the catalytic cleft is the largest for TgPPT1 (Figure 1C), intermediate for TgASH2 (Figure 1D), and is the smallest for TgASH3 (Figure 1E) and TgASH4 (Figure 1F), reflecting possible differences in the sizes of potential substrates that could be accommodated by each enzyme.

Biochemical analysis of TgASH esterase and thioesterase activities

Based on homology modeling and the presence of a pentapeptide lipase motif (GxSxG), we hypothesized that TgASH2-4 are likely to have distinct substrate preferences compared to TgPPT1. In addition, we previously reported that TgPPT1 and TgASH4 have distinct substrate preferences in vitro (Foe et al., 2018), however, the biochemical properties of TgASH2 and TgASH3 have not yet been determined. To determine the enzymatic functions of all of the TgASH proteins, we recombinantly expressed the full-length proteins in Escherichia coli, purified them (Figure S3D), and subjected them to biochemical analyses.

We first tested the purified enzymes for esterase and thioesterase activities using quenched substrate probes that mimic palmitoylated Ser (QSE) or Cys (QSTE) residues (Figure 2A) (Amara et al., 2019; Foe et al., 2018). TgPPT1 showed the highest activity of all the enzymes for these lipid substrates with similar activity for both QSE and QSTE. TgASH2 and TgASH3 weakly processed both substrates but preferred the thioester substrate. Both enzymes were still 6-fold less effective at processing this palmitate mimetic compared to TgPPT1. TgASH4 showed no activity toward either substrate consistent with its preference for short acyl substrates (Figures 2B and 2C) (Foe et al., 2018). We next tested whether the enzymes are capable of processing thioesters in general using a substrate with a short acyl chain, S-(4-nitrophenyl)thioacetate (Figure 2D). In contrast to the palmitoyl substrates, all enzymes were able to process the thioacetate, although TgPPT1 showed substantially reduced activity for the acyl substrate compared to ASH2-4, again highlighting the difference in substrate specificity of these serine hydrolases and supporting the hypothesis that only TgPPT1 is a depalmitoylase. We also wanted to determine if these hydrolases (and ASH4 in particular due to its preference for acetyl groups) could be capable of hydrolyzing acetyl-protein substrates. Because most acetyl modifications on proteins are found on lysine residues, we tested the ability of the recombinant proteins to cleave simple coumarin substrates modified with an acetyl group. We found that all four hydrolases TgPPT1, TgASH2, TgASH3, and TgASH4 processed the acetyl ester but not the acetyl amide substrate (Figure 1E). These results suggest that all four hydrolases primarily cleave esters and thioesters and are not likely to de-acylate amides, such as acyl lysine residues found on proteins.

To further define the substrate preference for each of the enzymes, we tested a panel of fluorogenic substrates that contain lipids of increasing carbon chain lengths (Figures 3A–3C). Consistent with its known function as a depalmitylase and with its relatively high activity toward QSTE, TgPPT1 preferred longer acyl chains (eight and ten carbons). Interestingly, TgASH2 was able to process both short acyl substrates and longer lipids but showed a preference for medium length lipids (seven carbons). TgASH3 and TgASH4 preferred short acyl chain substrates (two and four carbons) and were unable to process long-chain lipid acid esters (Figure 3C; Table S2). These classifications based on substrate specificity also fit with the alignment of the enzymes based on sequence similarities (Figures 1A, 1B, S1A, and S1B) and homology modeling showing the different sizes of the active site pockets (Figures 1C–1F), suggesting that the enzymes likely have distinct functions in vivo.

TgASH4 deletion alters the T. gondii lipid profile

We previously reported that deletion of TgASH4 results in parasites that form disordered rosettes (Foe et al., 2018), a phenotype similar to that observed when parasites are grown in the presence of excess cholesterol or oleic acid (Nolan et al., 2017, 2018; Robibaro et al., 2002) or when the phosphatidic acid (PA)/diacylglycerol (DAG) ratio is dysregulated in T. gondii (Bullen et al., 2016). Our sequence analysis revealed that TgASH2-4 each contain a calcium-independent phospholipase motif (Figures 1A, S1A, and S1C–S1I), suggesting that these enzymes are likely lipases (Jenkins et al., 2004). Combined with our in vitro biochemical data, which show that TgASH2-4 can cleave fatty esters, these observations strongly support a role for all three hydrolases in parasite lipid metabolism (Figures 1A and supplemental information Figure S1A). Because deletion of TgASH2 or TgASH3 does not result in any phenotypes (Kemp et al., 2013), we believe that these enzymes may play redundant or overlapping functions, making genetic studies of function difficult. Therefore, we focused on TgASH4, which has a clear defect in parasite replication inside the vacuole (Foe et al., 2018). To test our hypothesis that TgASH4 is involved in lipid metabolism, we performed lipid-profiling studies using wild-type and ΔTgASH4 parasites. We infected primary human foreskin fibroblast cells with wild-type T. gondii parasites as well as parasites in which the TgASH gene was disrupted (ΔTgASH4). We also generated parasites in the knockout background in which TgASH4 expression was rescued with WT (ΔTgASH4-S124A) TgASH4. After 24 h of infection, we collected parasites and extracted lipids for analysis by thin-layer chromatography (TLC). We found that loss of TgASH4 resulted in accumulation of both phospholipids and neutral lipids compared to wild-type parasites (Figures 4A and 4B). This

Please cite this article in press as: Onguka et al., Toxoplasma gondii serine hydrolases regulate parasite lipid mobilization during growth and replication within the host, Cell Chemical Biology (2021), https://doi.org/10.1016/j.chembiol.2021.05.001
accumulation was rescued by expression of wild-type (ΔTgAsh4-ASH4) but not catalytically dead mutant TgASH4 (Figures 4A and 4B).

To determine the identity of the lipids that accumulated in the absence of TgASH4, we extracted lipids from wild-type and ΔTgAsh4 parasites and analyzed the total lipidome by LC-MS. To increase the number of lipids identified, we used both positive and negative electrospray ionization modes for MS analysis. In general, we observe distinct lipid profiles in the parasites lacking TgASH4 compared to wild-type parasites (Figures 4C, 4D, S2A, and S2B), supporting our hypothesis that TgASH4 plays a role in lipid metabolism in T. gondii. A more detailed survey of the lipidomic data confirmed general defects in glycerolipid metabolism (Figures 4D, 4E, and S2B). Specifically, we found that levels of cytidine diphosphate-DAG (CDP-DAG) lipids were, on average, 16-fold decreased in the knockout parasites. We also observed a slight increase in PA upon loss of TgASH4 expression. Both of these classes of lipids are central intermediates in phospholipid and glycerophospholipid metabolism (Figures 4D, 4E, and S2B).

Synthesis of PA is the initiating event in phospholipid biosynthesis (Pascual and Carman, 2013). In eukaryotes, PA is a precursor for both CDP-DAG and DAG. CDP-DAG is used to make phosphoinositol, phosphatidylglycerol, and cardiolipin, while DAG is required for phosphatidylcholine, phosphatidylethanolamine, and triacylglycerol synthesis (Carman and Han, 2019; Henry et al., 2012) (Figure 2E).

Identification of small molecules that inhibit TgASH proteins

Given the likely roles for TgASH2-4 in lipid metabolism, we sought to identify inhibitors of these enzymes that could be used to further explore their biological function during T. gondii infection. Our laboratory has compiled a library of small molecules that covalently react with cysteine and serine hydrolases. This library contains around 1,200 compounds that have diverse scaffolds attached to reactive electrophiles (Arastu-Kapur et al., 2008; Child et al., 2013; Ponder et al., 2011). We therefore performed our screening against the TgASH proteins using the set of approximately 400 compounds that are likely to target serine hydrolases. Since none of the tested lipid substrates were optimal for all four enzymes, we chose to screen the library using...
an ABPP competition assay in which enzymes are pretreated with inhibitors and the residual activity measured by a gel readout of labeling using the general FP-TAMRA probe. This assay also allowed us to pool the recombinant enzymes and perform the screen on all targets at once (Figure S3). We pre-treated the mixture of enzymes with each compound for 30 min, after which we labeled the enzyme mixture with the broad-spectrum serine hydrolase probe, FP-TAMRA.
Compound binding appeared as a loss of labeling by the FP-TAMRA probe (Figure S4). Because the enzymes have distinct molecular weights they can be resolved using SDS-PAGE, so we could quantify the potency and selectivity of each compound for each enzyme target in a single lane of a gel.

As an initial cutoff, we selected compounds that reduced FP-TAMRA labeling by at least 60% at 33 μM (Figure 5A). To validate the hits, we used a secondary assay to measure their ability to inhibit the recombinant enzymes at a range of inhibitor concentrations. We switched to using the fluorogenic substrates for the validation studies because the results are more quantitative than in gel competition assays. However, since TgASH4 only processed the short-chain substrates (acetate and butyrate) that had high overall background signals from baseline hydrolysis, we continued to use the competition assay for this enzyme.

Figure 5. Identification of small molecules that inhibit TgASH proteins

(A) Dot plot of screening results of the gel-based competition assay with FP-TAMRA and serine reactive libraries against TgPPT1, TgASH2, TgASH3, and TgASH4. The enzymes were pre-incubated with 33 μM compound, or DMSO. Results display the residual TAMRA signal, normalized to the DMSO control and plotted as a percentage of TAMRA labeling relative to the DMSO control.

(B) Structures of selected compounds and their percent inhibition of respective enzymes at 10 μM as assayed by turnover of 4MU-octanoate (TgPPT1, TgASH2, and TgASH3) or gel-based competitive FP-TAMRA labeling for TgASH4. Data represent the mean ± standard deviation.

See also Figures S3, S4, and S6.
Figure 6. Inhibition of TgASH proteins impairs parasite infection

(A) Representative images of plaques formed by parasites after pretreatment of the parasites with either DMSO or the indicated screening hits (10 μM final).

(B) Quantification of total plaques numbers upon compound treatment. Data represent averages (bar plot) from three independent experiments performed in technical triplicates.

(C) Anti-Toxo, DAPI, Bright field images showing the localization of parasites.

(D) Fraction distribution of parasites/vac from 16pack, 8pack, 4pack, 2pack, and 1pack. Each bar represents the average from three independent experiments with technical triplicates.

(E) Fraction rosette and disordered images showing the effect of different compounds.

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(Figures 5B and S5B). Confirmed hit molecules shared a core chloroisocoumarin scaffold (JCP341, JCP342, JCP343, JCP348, and JCP388; Figure 5B). Chloroisocoumarins are electrophilic traps that form irreversible covalent bonds with active site nucleophiles of serine hydrolases. The compounds are initially attacked by the active site serine residue to form a reversible ester linkage and then by subsequent irreversible covalent modification by a nearby histidine. We validated inhibitors by calculating apparent half-maximal inhibitory concentration (IC_{50}) values following dose-dependent inhibition studies using a fixed time point assay (Figure S5B). Most of the screening hits inhibited all three serine hydrolases, although with overall differences in potency (IC_{50}) and maximum level of inhibition for each hydrolase. However, two of the lead compounds from the screen, JCP343 and JCP348, were potent inhibitors of ASH2-4 but showed virtually no activity for TgPPT1, making them highly valuable for studies of TgASH2-4 function.

To further assess the selectivity of the lead molecules we also performed ABPP competition studies in total parasite extracts using the FP-TAMRA probe (Figure S6). We found that pretreatment of lysates with 10 μM of each probe before labeling with the FP probe resulted in competition of several species. Specifically, we found that the two compounds with significant inhibitory activity against the recombinant TgPPT1 (JCP342 and JCP388) both showed competition of the band corresponding to TgPPT1 in the lysates, while the others did not. Furthermore, two of the top hits (JCP343 and JCP388) showed competition for labeling of species that are consistent in size with the recombinant TgASH2 and TgASH3. These results confirm that none of the identified hits have selectivity for individual TgASH proteins but, because some of them are inactive against TgPPT1, they are useful reagents for assessing the functional roles of the TgASH2-4.

**Inhibition of TgASH proteins impairs parasite infection**

For *T. gondii* to successfully establish an infection, the parasite must invade host cells (Carruthers and Boothroyd, 2007), undergo asexual replication (Clough and Frickel, 2017), egress (exit) from host cells, and invade new cells (Black and Boothroyd, 2000; Blackman and Carruthers, 2013). Therefore, to examine whether the inhibition of TgASH proteins impacts any of the asexual lytic stages of *T. gondii*, we treated extracellular tachyzoite parasites with the newly identified inhibitors of TgASH2-4. We used plaque and replication assays to measure parasite fitness upon compound treatment. Plaque assays can be used to assess the overall viability of the parasites through multiple rounds of the asexual lytic cycle. In these studies, we included the FDA-approved drug, orlistat, as a positive control, since it inhibits fatty acid synthase and has been shown to block lipid metabolism processes in humans and in *P. falciparum* (Yoo et al., 2020). All inhibitors severely impaired parasite growth with 50%–75% reduction in plaque numbers compared with a DMSO control (Figures 6A, 6B, and S5C). These data strongly suggest that TgASH2-4 are important for parasite intracellular growth.

To investigate which stage of the asexual lytic cycle is impacted by TgASH2-4 inhibition, we first analyzed the rate of replication in either the presence or absence of inhibitors. The rate of parasite replication can be determined by quantifying the number of parasites per vacuole upon treatment with either DMSO or inhibitor (see the STAR Methods for further information). Parasites pretreated with the identified hits resulted in significantly slower growth rate (less than 8 or 16 parasites/vacuole) compared with DMSO-treated controls, suggesting a disruption in the rate of parasite replication (Figure 6C). The relative potencies of the compounds in this assay matched the patterns we observed in the plaque assay (Figure 6B) and the potencies for the recombinant TgASH proteins (Figure 5B).

*T. gondii* tachyzoites divide asexually in the parasitophorous vacuole through a process called endodyogeny. During this process, parasites sequentially form daughter parasites that are connected by a residual body, resulting in the formation of a flower shaped “rosette” of daughter parasites. We previously showed that deletion of TgASH4 results in disordered rosettes in which cell division is disrupted and daughter pairs have individual residual bodies with altered parasite morphologies. We analyzed the effect of the ASH inhibitors from our library screen on the formation of rosettes inside the host cell and found that treatment of parasites with a 10 μM dose of the TgASH2-4 inhibitors resulted in a loss of normal rosettes and formation of the same type of disordered rosettes that we observed in the TgASH4 knockout parasites (Figures 6D and 6E). This observation suggests that TgASH2-4 inhibitors, like the general inhibitor of lipid metabolism, orlistat, produce a phenotype that we previously observed for the TgASH4 deletion mutant and which mimics the effects of excess cholesterol or oleic acid (Foe et al., 2018).

**DISCUSSION**

Although it is possible to predict protein function by assessment of sequence similarity to proteins with known functions (Dolinski and Botstein, 2007; Train et al., 2017), it is difficult to identify proteins with similar function if they lack sequence similarities. The α/β-hydrolase fold superfamily of enzymes comprises enzymes that contain a conserved catalytic triad (a serine nucleophile, catalytic acid, and histidine) and common structural fold. Despite conserved active sites and structural features, α/β-hydrolases perform diverse biological functions (Carr and Ollis, 2009; Ollis et al., 1992). Thus, the α/β-hydrolase superfamily is an excellent model to begin to understand the structure-function relationships of a common structural protein fold (Long and Cravatt, 2011). In this work, we employed bioinformatic, biochemical, cell biological, and pharmacological approaches to define the functions of genetically related serine hydrolases.
hydrolyase enzymes in the important human parasite pathogen *T. gondii*.

Sequence and structural analyses identified protein domains conserved among all four enzymes. It also highlighted some key differences, including a conserved calcium-independent phospholipase motif (Jenkins et al., 2004; Rajakumari and Daum, 2010), which is lacking in TgPPT1, and the presence of putative acyl transferase motif H(x)4D (Rajakumari and Daum, 2010) in TgASH3 and TgASH4. Although all of the ASH proteins aligned with human proteins annotated as depalmitoylases, our biochemical analyses indicate that only TgPPT1 is likely a bona fide depalmitoylase, while TgASH2, TgASH3, and TgASH4 exhibit only weak or no activity toward long-chain thioesters. Together, these biochemical results suggest that ASH proteins should be classified into three subsets based on their enzymatic activities (1) depalmitoylases (i.e., long-chain lipid thioesters)—TgPPT1; (2) medium chain lipid esterases—TgASH2, and (3) short-chain lipid esterases/de-acylases—TgASH3 and TgASH4. These classifications based on substrate specificity also fit with the alignment of the enzymes based on sequence homology (Figure 1B) and suggest they likely have distinct functions in vivo. While conserved positions in α/β-hydrolases define properties that are shared within the entire family (for example, the catalytic triad), they do not fully explain functional diversity or biological function (Long and Cravatt, 2011). These findings emphasize the complementary nature of bioinformatic and biochemical investigations. Whereas we were unable to predict substrate preference based on sequence similarity and homology modeling alone, clustering analysis suggested that the enzymes differed with respect to conserved domains, prompting our initial biochemical hypotheses.

Deletion of TgASH4 yields parasites that form smaller plaques, disordered rosettes, and dispersion defects compared with wild-type parasites (Foe et al., 2018). In addition, the balance between DAG and PA is important for *T. gondii* growth and replication. Furthermore, inhibition of enzymes involved in PA and DAG formation results in defects in rosette formation (Bullen et al., 2016). Our data support a model in which these defects are caused by disruption of lipid metabolism. The lipid profiles of parasites lacking TgASH4, or ΔTgASH4 rescued with catalytic dead TgASH4 accumulate both phospholipids and neutral lipids. Specifically, parasites lacking TgASH4 have lower levels of CDP-DAG and an increase in PAs and several sterols. CDP-DAG and PA are essential lipid intermediates that are involved in *de novo* synthesis of all phospholipids in both eukaryotes and prokaryotes. CDP-DAG synthase (CDS) produces CDP-DAG from PA and cytidine triphosphate. *T. gondii* contains two phylogenetically divergent CDS enzymes (Cds1 and Cds2) that reside in the ER and the apicoplast, respectively (Kong et al., 2017). Both TgCds1 and TgCds2 are essential and conditional knockdown results in parasites with reduced growth rates that could not be restored by addition of exogenous lipids, suggest that *de novo* synthesis of lipid intermediates is essential to parasite growth (Kong et al., 2017). These observations highlight the critical link between coordinated regulation of lipid flux and cell division.

Our previous studies showed that ΔTgASH4 parasites tend to have distended residual bodies that are derived from a single parasite and multiple cell division defects. These phenotypes include incomplete cytokinesis at the apical end after division has initiated, incorrect initiation of division from the basal side, defects in endodyogeny, and defects in cytokinesis (Foe et al., 2018). Our current pharmacological experiments yielded complementary observations, wherein parasites treated with the TgASH2-4 inhibitors exhibited growth and replication defects. Importantly, these phenotypes are similar to those observed for parasites grown under excess cholesterol or oleic acid. In addition, treatment of parasites with orlistat, a drug that is known to target lipid pathways (Ross and Fidock, 2019; Yoo et al., 2020), also results in disordered rosettes. These results suggest that TgASH2-4 are generally involved in parasite lipid metabolism.

In summary, we have characterized a set of serine hydrolases that were previously thought to be depalmitoylases based on sequence alignment, but our data suggest that they are in fact esterases involved in parasite lipid metabolism. We show that parasites lacking TgASH4 have dysregulated lipid metabolism resulting in a slight increase in PA and a substantial decrease in CDP-DAG, two key intermediates in the generation of phospholipid biogenesis that supply the entire endomembrane system with phospholipids (Carman and Han, 2019). In addition, fatty acids, cholesterol esters, and sphingolipids are increased in the absence of TgASH4. Although we cannot yet identify the particular metabolic pathway that links TgASH4 activity to lipid disruption, our data support a model in which TgASH4 is either directly or indirectly involved in CDP-DAG synthesis. A balance of PA/DAG is important for lipid metabolism and cell physiology since both act as precursors for other phospholipids and as signaling molecules. An abnormal increase in lipid levels results in cells susceptible to fatty acid-induced lipotoxicity, defects in vacuolar fusion and acidification, autophagy, and cell wall integrity. In the future, we plan to perform further analysis of higher-order combinations of deletions of the TgASH proteins to more specifically characterize the lipid metabolism pathways in which they operate.

**SIGNIFICANCE**

The intracellular protozoan parasite *Toxoplasma gondii* must scavenge cholesterol and other lipids from the host to facilitate its intracellular growth and replication, while avoiding excess accumulation and lipotoxicity. However, when and how parasites mobilize scavenged lipids required for growth is still an open question. We have bioinformatically and biochemically characterized several *T. gondii* serine hydrolases that are conserved across multiple phyla with no known function. We have shown that these enzymes, although initially thought to be depalmitoylases, are rather metabolic enzymes important for lipid regulation within the parasite during growth and development. The lipidomic and pharmacological studies presented here also provide further support for the essential role of lipid metabolism in *T. gondii* and suggest that targeting the TgASH proteins could be a viable strategy for treatment of *T. gondii* infections.
The authors declare no competing interests.

AUTHOR CONTRIBUTIONS

Detailed methods are provided in the online version of this paper and include the following:

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.chembiol.2021.05.001.

ACKNOWLEDGMENTS

We thank past and present Bogyo, Boothroyd, Egan, and Yeh laboratory members for their input and suggestions. This work was funded by NIH grant R01GM111703 (to M.B.), B.M.B. was supported by the A.P. Giannini Foundation, M.L. was supported by Deutsche Forschungsgemeinschaft (DFG) for funding under the Walter Benjamin Program.

AUTHOR CONTRIBUTIONS

O.O. designed and conducted the experiments, O.O. and M.B. conceptualized the work, analyzed data, and wrote the paper. O.O., B.M.B., and M.B. analyzed lipidomics data. M.J.N., K.M.L., P.C., and O.O. analyzed sequence, phylogenetic, and homology modeling. S.M.T. and J.Z.L. performed lipidomics analysis by LC-MS. I.T.F. generated recombinant plasmid constructs. N.A. synthesized QSE and QSE substrates.

DECLARATION OF INTERESTS

The authors declare no competing interests.

REFERENCES


## STAR★METHODS

### KEY RESOURCES TABLE

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RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Matthew Bogyo (mbogyo@stanford.edu).

Material availability
Plasmids and chemical reagents generated in this study are available by request to the lead contact. We may require a payment and/or a completed Materials Transfer Agreement if there is potential for commercial application.

Data and code availability
The lipidomics datasets generated during this study are available at Dryad: https://doi.org/10.5061/dryad.gf1vhhmpq.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Parasite culture
*T. gondii* strains Ku80, ASH4, ASH4 [ASH4], and ASH4 [ASH4 S14A] were grown in human foreskin fibroblasts (HFFs) using a mixture of Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% FetalPlex animal serum complex (Gemini Biotech, catalog no. 100602), 2mM L-glutamine, and a cocktail of 100μg/ml penicillin, and 100 μg/ml streptomycin. Parasites were cultured at 37 °C in 5% CO₂.

Bacterial culture for protein expression
HIS6-PPT1 (rPPT1), HIS6-ASH2 (rASH2), HIS6-ASH3 (rASH3), HIS6-ASH4 (rASH4) and with their mutant counterparts HIS6-PPT1S128A (rPPT1S128A), HIS6-ASH2S192A (rASH2S192A), HIS6-ASH3S277A (rASH3S277A), and HIS6-ASH4S124A (rASH4S124A) were expressed from the pIF22 and pIF22-S124A plasmids in BL21-CodonPlus (DE3)-RIL Escherichia coli (Agilent Technologies, catalog no. 230245). Expression was induced with 0.5 mM IPTG (isopropyl-B-D-thiogalactopyranoside) for 16 h at 19 °C.

METHOD DETAILS

Plasmid construction
Bacterial expression construct was created by PCR amplification of the ASH2-4 open reading frame (ORF) from the strain RH cDNA using primers. The PCR product was digested with Nde1 and BamH1 and ligated into pET-28A, resulting in 6XHis- appended to the N-terminus of *Tg* PPT1, *Tg* ASH2, *Tg* ASH3 and *Tg* ASH4.

Phylogenetic analysis
Sequence alignments were performed using Clustal omega algorithm (Sievers and Higgins, 2018). For modeling of the proteins, we searched for the best structural template by performing sequence alignment of a target sequence with the sequences of proteins, for which experimental structure is known. After which we used MODELER program (Ollis et al., 1992), this program builds three-dimensional model that satisfy various spatial restraints of the template structure. FFAS program (Jaroszewski et al., 2005) was then used to find the best templates (Jaroszewski et al., 2011). Building the final structural models has been done by MODELER program (Sali and Blundell, 1993). For each protein two structures were build, and only a single final model that satisfy/model best the geometries of the catalytic triad SDH in the active site of the enzymes and its FFAS score is the lowest has been selected for further analysis for each enzyme. The information about templates two structures were used for model building, including sequence identity, FFAS scores, positions of catalytic residues in the model and brief characteristics of the models used for final selection of the models is summarized in Table S1. The final models selected for structural analysis have been built using the following templates from PDB: for ASH1 – 5syn_A, ASH2 – 6imp_A, ASH3 – 5g59_A and for ASH4 – 5g59_A. The best models for ASH2-4 were aligned (PyMOL “super”) to that of ASH1 (*Tg*PPT1). The region of the active site cavity was visualized in using the “Cavity and Pockets” surfaces generated by PyMOL, taking into consideration the contiguity of the surfaces as well as the direction of the serine nucleophile. HOLLOW¹ (1.1) was used to generate a cast of the protein surface in a 15.0 Å radius centered on the active site serine oxygen (OG), with interior probe and surface probe radii set to 1.4 and 4.2 Å respectively (Ho and Gruswitz, 2008). The resulting casts were manually trimmed to coincide with the region of interest observed in PyMOL.

Protein purification
Recombinant proteins were purified as previously described (Child et al., 2013; Foe et al., 2015, 2018; Yoo et al., 2020) Protein concentrations were quantitated by bicinchoninic acid (BCA) assay.

In vitro biochemical assays
4MU substrate assays were performed as previously described (Foe et al., 2018). Substrates were used at a final concentration of 10 μM. Enzymes were used at a 50nM final concentration for the experiments represented in Figure 1A. Enzymes were used at a final
concentration of 50 nM for the experiments represented in Figure S1B in the supplemental information. Fluorescence (Ex = 335 nm and Em = 450 nm) was measured every minute on a Cytation 3 imaging reader (BioTek, Winooski, VT, USA) for 60 min. 4-Nitrophenol acetate (4NPA) and 4-nitrophenol thiol-acetate (4-S-NPA) assays were performed as previously described for 4-nitrophenol octanoate (4NPO) (Foe et al., 2018). Enzymes were used at a final concentration of 150 nM. Absorbance was monitored on a Cytation 3 imaging reader (BioTek, Winooski, VT, USA).

In vitro depalmitoylase activity assays
QStE and QSE assays were performed as previously described method (Amara et al., 2019; Foe et al., 2018). Purified enzymes were diluted in reaction buffer (1X PBS, 0.05% Tx-100, pH 7.4) and pipetted into a black plate. Fluorogenic substrates were then added and incubated at 37°C for 60 minutes. Reactions were monitored by a fluorescence reader (BioTek, Winooski, VT, USA). Reaction rates were determined by linear regression analysis of the initial linear phase of hydrolysis. Steady-state kinetics
Parameters of steady state kinetics were determined by measuring the increase of fluorescence of hydrolyzed fluorogenic substrates at 384-well black plate. Recombinant enzymes were then added to a final concentration of 50 nM. Reaction progress was recorded (Ex = 335 nm, Em = 450 nm) over a period of 30-60 minutes at 37°C on a Cytation-3 imaging reader (BioTek, Winooski, VT, USA). Concentrations of hydrolyzed products were determined using a standard concentration curve (unconjugated 4MU). Reaction rates were determined by linear regression analysis of the initial velocity (V0) of hydrolysis. Steady-state kinetics
Steady-state kinetics
Parameters of steady state kinetics were determined by measuring the increase of fluorescence of hydrolyzed fluorogenic substrates (4MU (Foe et al., 2018)). Stock solutions of fluorogenic substrates were diluted in reaction buffer (20 mM HEPES, 150 mM NaCl, 10 mM CHAPS, pH 7.4) and pipetted into a black plate. Fluorogenic substrates were then added and incubated at 37°C for 60 minutes. Concentrations of hydrolyzed products were determined using a standard concentration curve (unconjugated 4MU). Reaction rates were determined by linear regression analysis of the initial velocity (V0) to these data using GraphPad Prism. Catalytic efficiencies were calculated from the product of $K_{cat}/K_m$.

Generation of parasite lysates
T. gondii parasites were harvested by syringe lysis of infected HFF monolayers and lysed in PBS containing 5% NP40 and 0.1% SDS. in phosphate buffered saline (PBS). Lysates were clarified at 13000RPM in a table top centrifuge for 30 minutes, and the protein concentration was quantified with a BCA assay (Thermo Scientific).

Hit compound competition labeling assay with Tg WT lysate
Wild-type parasite lysate was resuspended in PBS buffer with 0.5% NP40 and 0.1% SDS to a total volume of 50 μL. Each hit compound (10μM) was pre-incubated with the lysate at 37°C for 30 minutes. After pre-incubation, FP-TAMRA was added to a final concentration of 1 μM and incubated for 30 minutes at 37°C. Reactions were quenched with 1× SDS-PAGE loading buffer, boiled for 10 minutes, and resolved by SDS-PAGE. Equal protein loading was assessed using coomassie stain.

FP-TAMRA competition assays
For competition assays, 300 ng of purified enzyme was incubated with DMSO or inhibitors at 10 mM for 30 minutes on ice, then 1 mM FP- TAMRA was added and samples were incubated for additional 30 at 37°C. Reactions were quenched with reducing SDS sample buffer, and the entire sample was resolved by SDS-PAGE. A typhoon flat-bed scanner was used to scan the gel (532-nm laser, 610-nm filter, PMT800).

Plaque assays
Confluent monolayers of HFFs grown in 6 well plates were infected with freshly egressed tachyzoites isolated from host cells by syringe lysis and filtered with a 5-μm filter to remove host cell debris. The number of parasites per microliter was counted on hemocytometer. A total of 200 parasites were added to confluent HFFs in 6-well dishes. Parasites were grown for 7 days, fixed with cold methanol, and stained with crystal violet.

Immunofluorescence microscopy
Parasites were allowed to infect confluent HFFs on coverslips for 24 h, after which coverslips were fixed 4% paraformaldehyde. Cover slips were permeabilized with 0.2% Triton X-100 in 1X phosphate-buffered saline (PBS) and blocked in 3% bovine serum albumin (BSA) in 1X PBS for 30 min. Parasites were stained with anti-Toxo-fluorescein isothiocyanate (anti-Toxo-FITC) (Thermo Fisher, catalog no. PA1-7253) antibody by incubating overnight at 4°C in 3% BSA in 1X PBS at 1:1000 dilution. After incubation with primary antibodies, coverslips were washed 3 times with 1 ml 1X PBS. Mounting medium was used with DAPI (4′,6-diamidino-2-phenylindole; Vector Laboratories Inc., catalog no. H-1200) to mount coverslips to slides. Slides were imaged by confocal microscopy on an LSM 700 laser scanning confocal microscope. The intensity levels of the images were adjusted such that no data were removed from images.
Intracellular growth assays
HFFs grown on coverslips in 24 well plates were infected with freshly egressed tachyzoites pre-treated with either DMSO or 10μM hit compounds, and grown in confluent HFF coverslips for 24 h. Coverslips were fixed and stained with the anti-Toxoplasma-FITC antibody as described above. Slides were imaged on an LSM 700 laser scanning confocal microscope. Cover slips were counted, and at least 150 vacuoles with 8 parasites/vacuole were counted.

Extraction of lipids from parasites and TLC analysis
For total lipids collection, parasites were grown in confluent HFF for 24 h. Parasites were collected by centrifugation at 1000g and washed twice with 1X PBS. Equal number of parasites were collected, extracted with 1.5 ml of 2:1 chloroform/methanol by vortexing at room temperature for 1hr. 0.3 ml of ddH2O was added, the samples were vortexed on high for 1 minute, and phases were separated by centrifugation at 1000 rpm in a clinical centrifuge at room temperature. The upper phase was removed by aspiration and the organic phase washed with 0.25 ml 1:1 methanol/water. After phase separation, the lower organic phase was transferred to a new borosilicate tube and dried down under a stream of liquid nitrogen. Chloroform-resuspended samples were loaded onto silica gel TLC plates and resolved in 1D twice, first using chloroform/methanol/H2O (65:25:4) to mobilize the polar lipids and secondly using Hexane/acetone (100:1) to mobilize the polar lipids. For visualization of the separated lipids, the developed TLC plates were air dried.

Sample preparation for lipidomics
For lipid analysis, 5 T25 flask of confluent monolayer of HFF were infected with parasites and grown for 24 hrs. Parasites were subsequently collected and resuspended in 1.5 ml of 1XPBS buffer and transferred into glass vials pre-loaded with 2:1 chloroform/methanol/1XPBS to a final ratio of 2:1:1 chloroform/methanol/1XPBS. Samples were vigorously shaken and lipid extraction was performed explained above. Five vials of lipids extracted from each strain (wild type and ΔTgASH4 parasites) were dried under nitrogen gas flow. Lipids were temporarily stored on dry ice before being transferred to a -80 freezer awaiting analysis on Q-TOF.

Lipid analysis using high-performance liquid chromatography-mass spectrometry
Mass spectrometry analysis was performed with an electrospray ionization source on an Agilent 6545 Q-TOF LC/MS in positive and negative ionization modes. For Q-TOF acquisition parameters, the mass range was set from 100 to 1200 m/z with an acquisition rate of 10 spectra/second and time of 100 ms/spectrum. For Dual AJS ESI source parameters, the drying gas temperature was set to 250°C with a flow rate of 12 l/min, and the nebulizer pressure was 20 psi. The sheath gas temperature was set to 300°C with a flow rate of 12 l/min. The capillary voltage was set to 3500 V and the fragmentor voltage was set to 100 V. For separation of nonpolar metabolites, reversed-phase chromatography was performed with a Luna 5 μm C5 100 Å LC column (Phenomenex cat # 00B-4043-E0). Samples were injected at 20 μl each. Mobile phases for positive ionization mode acquisition were as follows: Buffer A, 95:5 water/methanol with 0.1% formic acid; Buffer B, 60:35:5 isopropanol/methanol/water with 0.1% formic acid. Mobile phases for negative ionization mode acquisition were as follows: Buffer A, 95:5 water/methanol with 0.1% ammonium hydroxide; Buffer B, 60:35:5 isopropanol/methanol/water with 0.1% ammonium hydroxide. All solvents were HPLC-grade. The flow rate for each run started with 0.5 minutes 95% A / 5% B at 0.6 ml/min, followed by a gradient starting at 95% A / 5% B changing linearly to 5% A / 95% B at 0.6 ml/min over the course of 19.5 minutes, followed by a hold at 5% A / 95% B at 0.6 ml/min for 8 minutes and a final 2 minute at 95% A / 5% B at 0.6 ml/min. Raw files were converted to mzXML format with MSConvert (ProteoWizard) using the Peak Picking Vendor algorithm. Files were analyzed using the web-based XCMS platform (Tautenhahn et al., 2012) with the following settings: signal to noise threshold, 6; maximum tolerated m/z deviation, 30 ppm; frame width for overlapping peaks, 0.01; and peak width, 10-60 s. Integrated peak intensities were normalized between conditions by median fold change. Ions were matched to the METLIN database with a 10 ppm tolerance for mass error. The tables containing potential metabolite annotations for each mass feature were downloaded from the XCMS platform. In order to analyze the metabolites according to their chemical classes, metabolite names were transformed into universally readable chemical identifiers: Using python 3 and the pubchempy wrapper for the PubChem PUG REST API (Fahy et al., 2007) metabolite names were searched against the PubChem database (Kim et al., 2018) and the PubChem CID and SMILES data were matched to the respective mass features. If a given mass feature could correspond to numerous metabolites, all potential metabolites were considered.

Library screening with purified proteins
For each covalent inhibitor, DMSO stocks were diluted in reaction buffer (20 mM HEPES, 150 mM NaCl, 10 mM CHAPS, pH 7.4) to a final concentration of 300 mM, followed by the addition of purified recombinant depalmitoylase at a final concentration of 50 nM. Reactions were incubated at 37°C and fluorescence was measured (Ex = 410 nm, Em = 450 nm) over 30 minutes. Data points
collected from three replicates were normalized to the DMSO control and initial rates were calculated from a linear fit of the activity curves and expressed as relative fluorescence units per second (RFU/sec).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Details for each experiment, including number of replicates, can be found in the figure legends and the STAR Methods section. Quantitative analyses, including curve fitting for kinetic and inhibition constants, were performed with PRISM (GraphPad v. 8.4). Protein sequence matches were considered significant for HMMER e-values <0.01.