Acquisition of exogenous fatty acids renders apicoplast-based biosynthesis dispensable in tachyzoites of *Toxoplasma*

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*Toxoplasma gondii* is a common protozoan parasite that infects a wide range of hosts, including livestock and humans. Previous studies have suggested that the type 2 fatty acid synthesis (FAS2) pathway, located in the apicoplast (a nonphotosynthetic plastid relict), is crucial for the parasite’s survival. Here we examined the physiological relevance of fatty acid synthesis in *T. gondii* by focusing on the pyruvate dehydrogenase complex and malonyl-CoA-[acyl carrier protein] transacylase (FabD), which are located in the apicoplast to drive *de novo* fatty acid biosynthesis. Our results disclosed unexpected metabolic resilience of *T. gondii* tachyzoites, revealing that they can tolerate CRISPR/Cas9–assisted genetic deletions of three pyruvate dehydrogenase subunits or FabD. All mutants were fully viable in prolonged cultures, albeit with impaired growth and concurrent loss of the apicoplast. Even more surprisingly, these mutants displayed normal virulence in mice, suggesting an expendable role of the FAS2 pathway in *vivo*. Metabolic labeling of the Δpdh-e1α mutant showed reduced incorporation of glucose-derived carbon into fatty acids with medium chain lengths (C14:0 and C16:0), revealing that FAS2 activity was indeed compromised. Moreover, supplementation of exogenous C14:0 or C16:0 significantly reversed the growth defect in the Δpdh-e1α mutant, indicating salvage of these fatty acids. Together, these results demonstrate that the FAS2 pathway is dispensable during the lytic cycle of *Toxoplasma* because of its remarkable flexibility in acquiring fatty acids. Our findings question the long-held assumption that targeting this pathway has significant therapeutic potential for managing *Toxoplasma* infections.

*Toxoplasma gondii* is an obligate intracellular protozoan infecting a variety of hosts, including 30% of the world’s human population and many warm-blooded animals (1). It is an extremely successful pathogen, able to survive and propagate in virtually all nucleated cells of its host (2). One fundamental question to be addressed in the field is the metabolic strategies that allow *Toxoplasma* to survive in such diverse nutritional environments. Previous studies have demonstrated flexibility of carbon metabolism in this parasite, with it being able to use glucose, glutamine, lactate, and even amino acids as carbon sources to support its bioenergetic needs (3–5). More recently, Krishnan et al. (6) have constructed a curated genome-scale metabolic model, predicting additional metabolic plasticity in *T. gondii*. On the other hand, the detailed mechanisms underlying such notable metabolic flexibility are still largely unexplored.

Fatty acids (FAs) are essential to all living cells, as they have diverse biological functions ranging from energy storage to membrane biogenesis (7). All cells have evolved sophisticated approaches to satisfy their FA demands (8). Genomic analyses suggest that *Toxoplasma* has at least three potential pathways to synthesize FAs of different chain lengths (7). First, the parasite harbors a FAS1 enzyme, like its mammalian hosts, but its biochemical activity has not yet been confirmed. Second, it encodes a FAS2 pathway located in the apicoplast, which is a vestigial plastid existing in several apicomplexan parasites, including *Plasmodium* and *Eimeria* species. Third, *Toxoplasma* contains a fatty acid elongation pathway in the endoplasmic reticulum. Although the relative contribution of these pathways to net fatty acid biosynthesis has not been fully established, the current experimental data suggest that FAS2 is involved in *de novo* synthesis of medium to long acyl chains (9), whereas the fatty acid elongation pathway produces very-long-chain and unsaturated FAs (9, 10). Recent work has also shown that *Toxoplasma* can scavenge fatty acids, likely by inducing autophagy of lipid droplets in the host cell (11).

Among the aforementioned pathways of fatty acid biogenesis, FAS2 is the most extensively studied so far. It involves a series of sequential reactions catalyzed by distinct enzymes (PDH, acetyl-coenzyme A carboxylase, ACP, FabD, acyl carrier protein synthase, FabH, FabB/F, FabG, FabZ, and FabI), ultimately leading to *de novo* production of fatty acids with eight carbons or more (12, 13). FAS2 begins with production of acetyl-CoA in the apicoplast, which is believed to be catalyzed by the pyruvate dehydrogenase (PDH) complex. PDH belongs to the α-keto dehydrogenase family. It consists of four subunits, E1α, E1β, E2, and E3, which, along with respective cofactors, assemble into a complex with E1, E2, and E3 domains (14). These functional domains have pyruvate dehydrogenase, dihy-
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**Results**

**The TgPDH complex in the apicoplast originated from cyanobacteria**

The parasite genome encodes five genes annotated as PDH subunits: E1α, E1β, E2, E3-I, and E3-II. Previous work using ectopic expression of epitope-tagged PDH subunits as well as sera against native proteins showed apicoplast localization of the E1β and E2 subunits (19). We started our work by confirming the subcellular location of PDH through C-terminal tagging of each endogenous protein (Fig. S1B). A spaghetti monster HA (smHA) epitope (25) was fused to the C terminus of each subunit at the endogenous locus using CRISPR-Cas9–mediated site-specific integration in the RH Δku80 strain (26). PCR confirmed the desired integration of the smHA tag in each transgenic strain. Immunofluorescent staining showed that TgPDH-E1α, TgPDH-E1β, TgPDH-E2, and TgPDH-E3-I were

droplipoyl transacetylase, and dihydrodiploipoyl dehydrogenase activity, respectively (Fig. S1A) (15). The entire enzyme complex catalyzes net conversion of pyruvate into acetyl-CoA, CO₂, and NADH.

PDH is critical for carbon homeostasis and energy production in most eukaryotic cells (16), which have mitochondrion-dwelling PDH to link cytoplasmic glycolysis and the mitochondrial TCA cycle (17). Plant cells have a second PDH complex, located in the plastid, where it catalyzes production of acetyl-CoA from pyruvate to initiate de novo synthesis of fatty acids (18). Notably, Apicomplexa parasites like *Toxoplasma* express only one PDH complex in the apicoplast (14, 19), where it is thought to convert pyruvate to acetyl-CoA to fuel FAS2. On the other hand, these parasites have repurposed a branched-chain keto-acid dehydrogenase complex to perform the catalytic function of PDH in the mitochondrion (20). Genetic ablation of branched-chain keto-acid dehydrogenase–E1α is detrimental to the virulence of *Toxoplasma* and *Plasmodium* parasites due to impaired flux of glucose-derived carbon into the TCA cycle.

Because host cells do not have an apicoplast or the FAS2 pathway, the latter is considered an excellent drug target against apicomplexan parasites (21). Several chemicals inhibiting FAS2 enzymes have been shown to have anti-malarial, antitoxoplasmosis, and/or anti-neosporosis activities (22). Genetic studies of *Plasmodium* species suggest that the importance of FAS2 differs at different stages of the life cycle (12, 23, 24). In contrast, conditional depletion of the ACP remains the only genetic evidence to infer the importance of FAS2 in *Toxoplasma*. Knockdown of ACP results in reduced lipoylation of PDH, loss of the apicoplast, growth arrest in culture, and strong attenuation of virulence, suggesting a critical role of FAS2 in *Toxoplasma* (21). This study examined the physiological importance of PDH and FabD, revealing unprecedented plasticity in fatty acid biogenesis of *T. gondii*.

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**Figure 1. Construction of TgPDH-E1α deletion and complementation strains.** A, schematic showing generation of the Δpdh-e1α and Δpdh-e1α/PDH-E1α (CompPDH-E1α) strains via CRISPR-Cas9–assisted gene engineering. PCR1–PCR5 indicate screening of clonal mutants. B, diagnostic PCRs confirming the Δpdh-e1α and CompPDH-E1α strains. C, immunofluorescent assay showing the presence or absence of PDH-E1α. Samples were stained with anti-TgPDH-E1α (which was generated from mice using a recombinant antigen corresponding to residues 158–304 of TgPDH-E1α, immunoblot checking the expression of PDH-E1α CPN60 antibodies. Scale bars = 1 μm. D, immunoblot checking the expression of TgPDH-E1α (green); TgALD (red) was included as a loading control. The two bands for PDH-E1α in the complement strain correspond to the full-length (~70 kDa) and mature (~45 kDa, after removal of the apicoplast targeting sequence) forms of the protein.
expressed in the apicoplast, as indicated by colocalization with the organelle marker TgCPN60. On the other hand, TgPDH-E3-II colocalized with the mitochondrial marker TgHSP60 (Fig. S1C).

We next examined the evolutionary origin of all PDH subunits by phylogenetic analyses of their peptide sequences (Fig. S2). Phylograms constructed for individual TgPDH subunit revealed that E1α, E1β, E2, and E3-I segregated with corresponding plastid and cyanobacterial PDH counterparts, whereas E3-II showed a distinct cladding pattern with mitochondrial and bacterial PDH-E3 (Fig. S2). These results suggest that the four former subunits are derived from cyanobacteria via secondary symbiosis, similar to Plasmodium PDH subunits (14), and the hosting organelle apicoplast itself (27). Our phylogenetic analyses in conjunction with the aforementioned localization studies also suggest that the E1α, E1β, E2, and E3-I subunits form a functional PDH complex located in the apicoplast, whereas E3-II is likely a component of the mitochondrial dehydrogenase complex.

TgPDH-E1α supports parasite growth but is not needed for survival

To evaluate the physiological importance of TgPDH, we first knocked out the E1α subunit in the RH strain. The entire coding region of the TgPDH-E1α gene was replaced by the DHFR-TS selection marker, which confers pyrimethamine resistance (Fig. 1A). Diagnostic PCRs were used to screen single clones lacking TgPDH-E1α (Fig. 1B). Immunostaining confirmed disruption of TgPDH-E1α in the deletion mutants (Fig. 1, C and D). Unlike the parental strain, TgPDH-E1α was not detectable in the Δpdh-e1α strain. The Δpdh-e1α mutant grew slower in routine culture. Indeed, in plaque assays, which indicate the overall fitness of the parasites, the Δpdh-e1α mutant produced a distinct cladding pattern with mitochondrial and bacterial PDH-E3 (Fig. S2). These results suggest that the four former subunits are derived from cyanobacteria via secondary symbiosis, similar to Plasmodium PDH subunits (14), and the hosting organelle apicoplast itself (27). Our phylogenetic analyses in conjunction with the aforementioned localization studies also suggest that the E1α, E1β, E2, and E3-I subunits form a functional PDH complex located in the apicoplast, whereas E3-II is likely a component of the mitochondrial dehydrogenase complex.

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Taken together, these results reveal that the E1α subunit of TgPDH is required for optimal growth of T. gondii tachyzoites in vitro, but it is not essential for parasite survival.

To confirm whether the growth defect observed in the Δpdh-e1α mutant was a direct consequence of PDH-E1α inactivation, we complemented the Δpdh-e1α strain with C-terminally Ty-tagged TgPDH-E1α expressed from the HXGPRT locus (Fig. 1A). Screening PCRs confirmed the desired integration of the complementing construct (Fig. 1B). Immunostaining demonstrated that the ectopic PDH-E1α was successfully expressed and localized to the apicoplast, just like the endogenous protein (Fig. 1, C and D). Phenotypic analysis of the PDH-E1α–complemented strain showed restoration of the observed defects in the Δpdh-e1α mutant (Fig. 2), confirming that PDH-E1α is indeed required for robust growth of the parasites.

Fatty acid synthesis is impaired in the Δpdh-e1α mutant

To determine the underlying basis of compromised growth in the Δpdh-e1α strain, we estimated de novo fatty acid synthesis by testing the flux of [13C6]glucose-derived carbon (Fig. 3A). Intracellular parasites of the parental and mutant strains were labeled with [13C6]glucose for 48 h, and subsequent incorporation of sugar-derived carbon (13C) into fatty acids was assessed by GC-MS. Our choice of labeling intracellular parasites was based on earlier work, which demonstrated that fatty acid synthesis is negligible in the extracellular stage (4). We found that the Δpdh-e1α mutant showed reduced 13C-labeling of fatty acids compared with the parental strain. The extent of tracer incorporation differed the most for myristic (C14:0) and palmitic (C16:0) acids. PDH-E1α deletion led to decreased incorporation of 13C into C14:0 and C16:0 (Fig. 3, A–C). Isotopologue profiling of myristic acid showed a significant amount of m12, m13, and m14 isotopomers (corresponding to 12, 13, and 14 carbons labeled with 13C) in the parental strain, whereas the majority of them were in the m0 fraction (i.e. no 13C labeling) in the Δpdh-e1α mutant (Fig. 3B). A similar pattern of isotopologue abundance was also found in palmitic acid (Fig. 3C), suggesting a contribution of PDH to flux of glucose-derived carbon into medium-chain fatty acids. Because these two fatty acids are believed to be produced mainly in the apicoplast by FAS2 (9), our data indicate a critical role of PDH in de novo fatty acid
synthesis. We also performed [13C6]glucose labeling in extracellular parasites (4 h) to examine carbon flux into glycolysis, the TCA cycle, and amino acids. None of these pathways were affected by PDH-E1α/H9251 disruption (Fig. 3D), as expected.

A somewhat selective reduction in synthesis of C14:0 and C16:0 in the Δpdh-e1α strain prompted us to restore the mutant’s growth by addition of exogenous C14:0 and C16:0.

Plaque and replication assays were performed in the absence or presence of 100 μM myristic acid, palmitic acid, or a mixture of both (50 μM each). Supplementation indeed improved growth of the Δpdh-e1α mutant, as shown by an almost 50% increase in plaque size with either fatty acid (Fig. 4A and B). This finding was further supported by replication rates that were significantly increased in the mutant upon addition of fatty acids (Fig. 4C). An additive effect on the mutant’s growth was also apparent when both acyl chains were provided simultaneously. Collectively, these experiments demonstrate the functional importance of PDH-E1α for fatty acid biogenesis, likely in the apicoplast.

The E2 and E3-I subunits of the PDH complex are dispensable

To consolidate the above results, we constructed two additional mutants lacking the E2 or E3-I subunits of the PDH complex, utilizing a similar homologous gene replacement strategy as shown in Fig. 1A. Genes encoding the E2 or E3-I subunits

Figure 3. The Δpdh-e1α strain exhibits reduced fatty acid synthesis. A–C, intracellular tachyzoites of the parental and Δpdh-e1α strains were labeled with [13C6]glucose for 48 h. Subsequently, fatty acids were extracted from host-free parasites, and the relative abundance of isotopologues was assessed by GC-MS. Means ± S.E. from three experiments (*, p < 0.05; **, p < 0.01; Student’s t test). A, overall fraction of 13C-labeled fatty acids regardless of the number of carbons that were labeled. B and C, 13C inclusion in isotopologues of myristic acid (C14:0) and palmitic acid (C16:0). D, flux of [13C6]glucose-derived tracer carbon into glycolysis and TCA cycle metabolites and amino acids. Extracellular parasites were labeled with [13C6]glucose for 4 h, followed by metabolite extraction and GC-MS analysis. W, RH; Δ, Δpdh-e1α; Pyr, pyruvate; Lac, lactate; Cit, citrate; α-Kg, α-ketoglutarate; Suc, succinate; Fum, fumarate; Mal, malate.

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were completely replaced by the DHFR-TS* selection marker. Screening PCRs confirmed successful deletion of individual genes in corresponding mutants (Fig. 5, A and B). The Δpdh-e2 mutant was further ascertained by immunostaining and immunoblot analyses (Fig. 5, C and D). The Δpdh-e2 and Δpdh-e3-I strains displayed reduced growth in plaque and proliferation assays (Fig. 6, A and B), phenocopying the Δpdh-e1α mutant (Fig. 2). Nonetheless, the Δpdh-e2 and Δpdh-e3-I mutants were viable in prolonged cultures, suggesting that neither subunit is essential for tachyzoites.

**PDH is dispensable for the parasite virulence**

Our results strongly assert that the parasite can tolerate the absence of the PDH complex, albeit with impaired reproduction in vitro. To examine the in vivo relevance, we infected ICR mice with the abovementioned mutants lacking individual PDH subunits and subsequently scored the survival of infected animals. Surprisingly, although all three PDH-deficient mutants had a variable degree of growth defects in culture, none of them showed evidence of attenuated virulence in mice (Fig. 6C). In all cases, parasitized animals succumbed to death within 10–12 days of infection, suggesting complete dispensability of the PDH complex for parasite virulence while questioning the significance of the FAS2 pathway during acute infection with *T. gondii*.

**PDH mutants show loss of the apicoplast correlated with their growth phenotypes**

Interruption of metabolic processes in the apicoplast is typically associated with its loss over time, as reported for the ACP depletion mutant (21). We therefore examined the integrity of the apicoplast in the above PDH mutants. The Δpdh-e1α, Δpdh-e2, and Δpdh-e3-I strains were subjected to immunostaining of the organelle marker TgCPN60. In the parental strain, nearly all parasites harbored an apicoplast, as indicated by the bright TgCPN60 signal (Fig. 7A). In the PDH deletion mutants, however, the integrity of the apicoplast varied. The Δpdh-e2 mutant did not seem to show significant defects in apicoplast maintenance, as judged by strong TgCPN60 staining in most tachyzoites (Fig. 7, A and B). The Δpdh-e1α mutant experienced modest apicoplast loss, in which about 15% parasites did not have detectable TgCPN60. The Δpdh-e3-I mutant displayed the most dramatic phenotype, with almost 50% of parasites devoid of TgCPN60 signal. The degree of apicoplast loss in individual mutants correlated well with the growth defects in plaque assays.

The apicoplast integrity in the PDH mutants was also assessed using an anti-TgPYK2 antibody. In the parental (RH) strain, TgPYK2 was mainly localized in the apicoplast, although it also displayed weak staining in vesicle-like structures. In contrast, in the PDH-E1α and PDH-E2 knockout mutants, TgPYK2 signal in the apicoplast was reduced, and increased staining of vesicle-like structures was observed (Fig. S4). Such a shift of the PYK2 staining signal was even more evident in the Δpdh-e3-I mutant, half of which had TgPYK2 detectable only in vesicle-like structures but not in the apicoplast (Fig. S4). These results confirmed our aforementioned observations with TgCPN60 staining. Although the basis of such differential apicoplast loss in different PDH mutants is inexplicable, our results suggest a role of PDH in maintenance of this organelle.
Deletion of FabD confirms nonessentiality of the FAS2 pathway

The genetic disruption of three PDH subunits indicated that FAS2 is dispensable, which is in contrast to a previous report of conditional depletion of ACP (21). We therefore revisited the physiological significance of FAS2 by targeting the FabD enzyme, which catalyzes transfer of the malonyl group from malonyl-CoA to ACP to produce malonyl-ACP. The latter serves as the carbon donor for the elongation cycles of FAS2. Interestingly, FabD was readily knocked out in the type 1 RH strain (Fig. 8A). Plaque and replication assays demonstrated that \( \Delta \text{pdh-e2} \) mutants also exhibited noticeable growth defects (Fig. 8, B and D), which could be restored by complementation. Moreover, although some \( \Delta \text{fabD} \)-infected animals survived slightly longer than the control group, the virulence of the \( \Delta \text{fabD} \) mutant was not significantly different from that of the parental strain (Fig. 8E). These phenotypes resonate well with that of the PDH-deficient mutants. Hence, these results together show that the FAS2 pathway is indeed not essential in tachyzoites.

Discussion

PDH is a central enzyme complex in carbon metabolism that converts glycolysis-derived pyruvate into acetyl-CoA (16). In apicomplexan parasites, there is only one PDH complex, which is localized in the apicoplast instead of the mitochondrion (12). The apicoplast is a unique organelle in most apicomplexan parasites that houses many important metabolic pathways, including fatty acid synthesis (27). The PDH complex is believed to produce acetyl-CoA, which is required by the FAS2 pathway in the apicoplast of \( T. gondii \) (14). According to a previous report (21), depletion of ACP (a core component of FAS2) severely reduced parasite growth in plaque assays and attenuated virulence in mice, implying an essential nature of FAS2 in the apicoplast. Accordingly, the PDH complex was also expected to be vital for the lytic cycle. Unexpectedly, however, we discovered that, despite a slower growth phenotype \textit{in vitro}, mutants lacking the E1α, E2, and E3-I subunits of \( Tg \)PDH are viable in prolonged cultures and fully virulent in mice, demonstrating dispensability of PDH for parasite survival and virulence. The viability of PDH-deficient mutants also suggests nonessential-
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Figure 7. Assessment of apicoplast integrity in PDH mutants. A, immunofluorescent staining of the apicoplast-specific marker TgCPN60. Yellow arrows indicate parasites with no evident apicoplast staining. Scale bars = 2 μm. B, quantification of apicoplast from A. The percentage of parasites with apicoplasts in parasitophorous vacuoles (PVs) of different sizes was determined by TgCPN60 and TgMIC2 staining (means ± S.E. of 150 vacuoles, n = 3 assays; *, p < 0.05; **, p < 0.01; Student’s t test).

Figure 8. FabD deletion mutants are viable in culture and virulent in mice. A, screening PCR confirmed successful replacement of FabD by the selection marker in ΔfabD. B, plaque assay of the indicated strains. C, relative size of plaques in B (means ± S.E.; n = 2 assays, each with 3 replicates; ***, p < 0.001; one-way ANOVA with Bonferroni’s post-test). D, intracellular replication (24 h) assay (***, p < 0.001; two-way ANOVA with Bonferroni’s post-test). E, survival of mice infected with the indicated strains.

ity of FAS2. These results were confirmed by the ΔfabD mutant, which was also viable in sustained culture and fully virulent in mice.

In further work, we demonstrated a role of the PDH complex in de novo fatty acid synthesis. Among different subunits of the PDH complex, E1α and E1β form a heterotetrameric E1 domain that catalyzes decarboxylation of pyruvate to initiate the whole reaction catalyzed by PDH (15). We characterized the Δpdh-e1α mutant to decipher the function of PDH in T. gondii. When labeled with [13C6]glucose, the flux of sugar-derived carbon into myristic acid (C14:0) and palmitic acid (C16:0) declined significantly in the Δpdh-e1α strain. C14:0 and C16:0 are primarily produced by FAS2 (7). Therefore it can be argued that TgPDH activity supports acyl chain synthesis in the apicoplast and that reduced FAS2 activity upon deletion of PDH-E1α underlies the growth defect. Conversely, growth of the Δpdh-e1α mutant was improved by supply of exogenous fatty acids, suggesting that T. gondii can salvage fatty acids from the environment, although the detailed mechanisms warrant further investigation. The salvage and de novo synthesis pathways may cooperate to ensure fatty acid biogenesis under diverse conditions.

Notably, deletion of PDH-E1α did not completely abolish the flux of [13C6]glucose-derived carbon into C14:0 and C16:0 acyl
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chains (Fig. 3, A–C), which can be explained as follows. Because metabolic labeling was performed with parasites replicating in host cells, there may have been a contribution of host components in our parasite samples. Moreover, salvage of host-derived fatty acids may be induced upon deletion of PDH-E1α. Likewise, we cannot rule out a contribution of the FAS1 pathway in the parasite, although its activity has not yet been reported. Last but not least, PDH may not be the only source of acetyl-CoA in the apicoplast. There are acetyl-CoA pools in the parasite cytosol and mitochondrion that may fuel the apicoplast with acetyl-CoA, although there is no reported apicoplast-localized acetyl-CoA transporter. Nonetheless, considering such additional sources of acetyl-CoA, we deleted FabD to examine the actual significance of FAS2. Because the PDH and FabD deletion mutants phenocopied each other, we believe that even if there is yet another source of acetyl-CoA for the apicoplast, its contribution to the FAS2 pathway is probably rather minor.

The phenotypes of the PDH/FabD deletion mutants are not fully consistent with that of the mutant depleted with ACP (21). Although these strains showed impaired growth in vitro, the most obvious difference is their virulence. The ACP knockdown strain is highly attenuated in mice whereas PDH/FabD-null mutants are not. Such discrepancies may be explained by the following. TgACP may have other crucial roles beyond its core function as an acyl carrier protein. In this regard, PfACP has already been shown to have acylation and malonyl-transf erase activities, which may contribute to the apicoplast metabolism beyond FAS2 (29, 30). Given the conservation, TgACP could exhibit similar activities. The ACP depletion mutant was engineered in the TATi strain, which itself is highly attenuated in mice for unknown reasons. Therefore, it is difficult to draw solid conclusions regarding the effect of ACP knockdown on parasite virulence using this strain. On the other hand, using clean genetic knockouts, our results have clearly shown that PDH and FabD are important but dispensable for parasite virulence. The recent work of Krishnan et al. (6) regarding FabZ is consistent with the data reported here.

Since its identification, FAS2 has been studied extensively in malaria parasites as a potential drug target. However, more recent work using gene deletion mutants (FabI, FabB/F, and FabZ, etc.) demonstrated that FAS2 is dispensable for blood-stage parasites but critical for mosquito or liver-stage developments (12, 23, 24). The dispensability of FAS2 for blood-stage Plasmodium is explained by their ability to scavenge fatty acids from red blood cells (24). Consistently, Plasmodium species lacking PDH subunits show normal erythrocytic development but growth defects during the hepatic or mosquito stages (23, 31, 32). Similarly, FAS2 in Toxoplasma may have more important roles in other stages, such as during sporulation, when the parasite undergoes extensive membrane synthesis but does not have access to host-derived resources. At least the tachyzoite stage has multiple routes to acquire fatty acids, which render de novo synthesis dispensable, questioning the long-believed potential of FAS2 as an antiparasitic drug target.

Experimental procedures

Biological resources and reagents

Type I RH and its derivative strains as well as the PDH and FabD deletion mutants generated here were propagated in human foreskin fibroblast cells (ATCC), as described elsewhere (33). Anti-Ty (mouse), anti-TgSAG1 (mouse), anti-TgMIC2 (mouse), and anti-TgALD (rabbit) antibodies were provided by David Sibley (Washington University, St. Louis, MO). Anti-TgCPN60 (rabbit) was provided by Honglin Jia (Harbin Veterinary Research Institute). Anti-TgPDH-E2 (mouse) was donated by Wolfgang Bohne (University of Göttingen, Göttingen, Germany). Anti-HA (rabbit) was purchased from Medical & Biological Laboratories Co.

Plasmid construction

All primers used in this study are listed in Table S1. Locus-specific CRISPR constructs were generated by replacing the UPRT-targeting guide RNA in pSAG1-Cas9-U6-sgUPRT with gene-specific guide RNAs, following a protocol described elsewhere (33, 34). Homology templates used for gene replacements (pPDH-E1α::DHFR-TS*, pPDH-E2::DHFR-TS*, pPDH-E3::DHFR-TS*, and pFabD::DHFR-TS*) were constructed by multifragment cloning using the ClonExpress MultiS Cloning Kit (Vazyme Biotech, Nanjing, China). Briefly, the 5′ and 3′ homology arms of each gene (~1 kb) were amplified from the genomic DNA of the RH strain and cloned into the pUC19 vector along with the pyrimethamine selection cassette DHFR-TS*. The latter was amplified from pUPRT-DHFR-D (34). To make the complementation constructs pComp-PDH-E1α and pComp-FabD, TgLDH1 was replaced in pCom-LDH1 (35) with the epitope-tagged coding sequence of TgPDH-E1α or FabD, respectively.

Generation of transgenic parasites

Endogenous tagging was performed as described previously (36). The 3′ genomic tagging cassette (smHA epitope and DHFR-TS* selection marker) flanked by gene-specific homology arms (50 bp) was amplified from the pSL24m-Linker-smFP-DHFR-LoxP-T7 plasmid (25) using the indicated primers (Table S1). Amplicons were then cotransfected into RH Δku80 along with a CRISPR plasmid targeting the 3′ UTR of the gene of interest (26). Transfected cells were selected with 1 μM pyrimethamine (Sigma-Aldrich), followed by PCR screening and immunostaining.

The Δpdh-e1α, Δpdh-e2, Δpdh-e3-I, and ΔfabD mutants were constructed by CRISPR-Cas9–assisted homologous recombination as described earlier (33, 34). Briefly, the gene-specific CRISPR plasmid was cotransfected with homologous donor amplicons into freshly purified tachyzoites and then selected with pyrimethamine as mentioned above. Subsequently, clonal mutants were isolated and identified by PCR screening and immunostaining. The PDH-E1α and FabD complement strain was generated by cotransfecting the HXGPRT-specific (for PDH-E1α complementation) or UPRT-specific (for FabD complementation) CRISPR plasmids along with the PDH-E1α- or FabD-expressing cassettes into the correspond-
ing mutants and selected with 30 μM chloramphenicol (Sigma-Aldrich).

**Phenotypic analysis of parasites**

Plaque assays were performed as described previously (34). Similarly, invasion, replication, and egress assays were executed according to established protocols (35, 37). To test virulence, 7-week-old female ICR mice (Hubei Provincial Center of Disease Control and Prevention) were infected by intraperitoneal injection of the indicated strains (100 tachyzoites/mouse). Infected animals were monitored daily for symptoms and survival over time. All animal experiments were approved by the Ethics Committee of Huazhong Agricultural University (Permit HZAUMO-2017-029).

**Metabolite measurements by GC-MS**

To analyze fatty acids, intracellularly proliferating parasites were cultured in medium containing 8 mM [13C6]glucose (Sigma Aldrich) for 48 h. Tachyzoites were purified from host cells by 3.0-μm filtration, and samples (2.5 × 107 cells) were lysed in 1 ml of ice-cold methyl alcohol (50%) and then ultra-sonicated (5 pulses of 1 min, each with a 1-min interval), followed by centrifugation (1.6 × 104 × g, 4 °C, 15 min). Subsequently, 800 μl of supernatant and 10 μl of internal standard (250 μg/ml nonadecanoic acid) were mixed and methylated overnight in Polytetrafluoroethylene screw-cap glass vials with 1 ml of 10% methanolic acetyl-chloride and 250 μl of n-hexane at room temperature. Then 5 ml of 6% potassium carbonate solution was added, and 150 μl of the n-hexane layer was transferred to a glass vial. Soon thereafter, 20 μg of anhydrous sodium sulfate was added to eliminate any trace of water, followed by transfer of a 60-μl sample for GC-MS (38).

To examine other metabolites, fresh extracellular tachyzoites (2.5 × 107) were labeled with 8 mM [13C6]glucose in glucose-free DMEM for 4 h. Parasites were pelleted and lysed in 1 ml of ice-cold methyl alcohol (50%), as mentioned above. Chlo-roform (0.8 ml) was added, and samples were vortexed for 30 s. The mixture was subjected to ultrasonication (5 cycles of 1 min, each with a 1-min interval), and samples were centrifuged (1.6 × 104 × g, 4 °C, 15 min). 700 μl of supernatant and 10 μl of internal standard (50 μg/ml 1-norleucine) were mixed, followed by evaporation to dryness under a nitrogen stream. The dry residue was reconstituted in 30 μl of 20 mg/ml methoxyamine hydrochloride in pyridine, and the resulting mixture was incubated at 37 °C for 90 min. Finally, 30 μl of N-tetra-Butyldimethylsilyl-N-methyltrifluoroacetamide (containing 1% Tert-butyldimethylsilyl) was added into the mixture and derivatized at 55 °C for 60 min before GC-MS. Instrument analysis was performed as described previously (35).

**Data analysis and statistics**

All experiments were performed at least three independent times unless specified otherwise. Data plotting and statistical analysis, as indicated in the figure legends, were done using GraphPad Prism (GraphPad Software Inc.).

**Importance of PDH and FAS2 in Toxoplasma**

**Data availability**

All data used in this work are contained within the manuscript.

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**Abbreviations**—The abbreviations used are: FA, fatty acid; PDH, pyruvate dehydrogenase; ACP, acyl carrier protein; FabD, malonyl-CoA-[acyl carrier protein] transacylase; smHA, spaghetti monster HA; TCA, tricarboxylic acid; ANOVA, analysis of variance.

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