The Role of Type II Fatty Acid Synthesis Enzymes FabZ, ODSCI, and ODSCII in the Pathogenesis of Toxoplasma gondii Infection

Xiao-Pei Xu1,2, Hany M. Elsheikha3, Wen-Ge Liu1, Zhi-Wei Zhang1, Li-Xiu Sun1, Qin-Li Liang1, Ming-Xin Song2* and Xing-Quan Zhu1,4,5*

1 State Key Laboratory of Veterinary Etiological Biology, Key Laboratory of Veterinary Parasitology of Gansu Province, Lanzhou Veterinary Research Institute, Chinese Academy of Agricultural Sciences, Lanzhou, China; 2 Heilongjiang Key Laboratory for Zoonosis, College of Veterinary Medicine, Northeast Agricultural University, Harbin, China; 3 Faculty of Medicine and Health Sciences, School of Veterinary Medicine and Science, University of Nottingham, Nottingham, United Kingdom; 4 College of Veterinary Medicine, Shanxi Agricultural University, Taigu, China; 5 Key Laboratory of Veterinary Public Health of Higher Education of Yunnan Province, College of Veterinary Medicine, Yunnan Agricultural University, Kunming, China

Toxoplasma gondii is an obligate intracellular protozoan parasite, which has a worldwide distribution and can infect a large number of warm-blooded animals and humans. T. gondii must colonize and proliferate inside the host cells in order to maintain its own survival by securing essential nutrients for the development of the newly generated tachyzoites. The type II fatty acid biosynthesis pathway (FASII) in the apicoplast is essential for the growth and survival of T. gondii. We investigated whether deletion of genes in the FASII pathway influences the in vitro growth and in vivo virulence of T. gondii. We focused on beta-hydroxyacyl-acyl carrier protein dehydratase (FabZ) and oxidoreductase, short chain dehydrogenase/reductase family proteins ODSCI and ODSCII. We constructed T. gondii strains deficient in FabZ, ODSCI, and ODSCII using CRISPR-Cas9 gene editing technology. The results of immunofluorescence assay, plaque assay, proliferation assay and egress assay showed that in RH1 FabZ strain the apicoplast was partly lost and the growth ability of the parasite in vitro was significantly inhibited, while for RHΔODSCI and RHΔODSCII mutant strains no similar changes were detected. RHΔFabZ exhibited reduced virulence for mice compared with RHΔODSCI and RHΔODSCII, as shown by the improved survival rate. Deletion of FabZ in the PRU strain significantly decreased the brain cyst burden in mice compared with PRUΔODSCI and PRUΔODSCII. Collectively, these findings suggest that FabZ contributes to the growth and virulence of T. gondii, while ODSCI and ODSCII do not contribute to these traits.

Keywords: Toxoplasma gondii, virulence factors, FASII enzymes, FabZ, ODSCI, ODSCII
INTRODUCTION

Toxoplasma gondii is an obligate intracellular protozoan, which is distributed all over the world. This parasite can cause adverse health impacts on the immunodeficient individuals such as AIDS patients or organ transplant recipients (Xiao et al., 2010; Robert-Gangneux and Dardé, 2012; Chemoh et al., 2015). T. gondii can survive in nearly all nucleated cells (Blume et al., 2009; MacRae et al., 2012; Xia et al., 2019) and completes its intracellular reproduction inside a parasitophorous vacuole (PV) after invading the host cell. The synthesis or uptake of fatty acids and other molecules from the host cell is important for the growth and proliferation of intracellular pathogens. Fatty acids are the main components of the PV, the membranes of the apicoplast and other organelles in T. gondii, underscoring the significance of fatty acids for the survival of T. gondii (Mordue et al., 1999; Ramakrishnan et al., 2012; Botté et al., 2013).

The type II fatty acid biosynthesis pathway (FASII) in apicomplexan protozoa is located in a unique organelle known as apicoplast, which has an independent genome of prokaryotic origin with no counterpart in the human or animal host (Ralph et al., 2004), which makes apicoplast and FASII a source of novel drug targets for T. gondii (Wiesner et al., 2008; Aygün and Mutlu, 2020). FASII, via a variety of enzymes, plays an important role in fatty acid metabolism. Phosphoenolpyruvate, after import to the apicoplast, is metabolized into pyruvate and converted into acetyl-CoA (a key precursor for FASII) via the action of pyruvate dehydrogenase complex (PDH) (Frege, 2007). Acetyl-CoA can be also produced in the nucleus and cytosol via the cation of acetyl-CoA synthetase and ATP citrate lyase (Tymoshenko et al., 2015), and in the mitochondrion via the action of branched-chain α-keto acid dehydrogenase-complex (Oppenheim et al., 2014). Although cytosolic acetyl-CoA is required for the synthesis of parasite fatty acids, metabolic adaptation by the parasite compensates for any loss of mitochondrial acetyl-CoA (Kloehn et al., 2020).

The acyl carrier protein (ACP) plays a role in FASII. Deletion of ACP causes growth defect and decreases the virulence of T. gondii (Zumudar et al., 2006). The synthesis of FASII in the apicoplast of T. gondii plays a key role in the survival of T. gondii, and the β-hydroxyacyl carrier protein dehydratase (FabZ; TGME49_321570) is essential for FASII metabolism in T. gondii (Wu et al., 2018; Krishnan et al., 2020). The protein coding genes ODSCI (TGGT1_269400) and ODSCII (TGGT1_313050) (ToxoDB database) belong to oxidoreductase, short chain dehydrogenase/reductase family (Kowalk et al., 2009). According to KEGG analysis, ODSCI and ODSCII are involved in the FASII pathway. According to Enzyme Commission classification, they also belong to 3-oxoacyl-[acyl-carrier-protein] reductase, which is involved in type II fatty acid biosynthesis.

The knockout of FabZ led to the loss of apicoplast of T. gondii tachyzoites, and decreased the level of myristate (C14: 0) and palmitate (C16: 0), the synthetic products of FASII, which resulted in the inhibition of the growth of T. gondii (Krishnan et al., 2020). In the present study, we hypothesized that mutation in T. gondii genes FabZ, ODSCI, and ODSCII involved in fatty acid biosynthesis and metabolism may render the mutant parasite strains less virulent and incapable of establishing an infection. We constructed knockout strains using CRISPR-Cas9 technology, and evaluated the effects of deletion of FabZ, ODSCI, and ODSCII on the virulence of the mutant strains in mice and on the in vitro growth using plaque assay, intracellular proliferation assay and egress assay.

MATERIALS AND METHODS

Toxoplasma gondii Cultures

T. gondii tachyzoites of the virulent RH (Genotype I) and less virulent PRU (Genotype II) strains were maintained in vitro in human foreskin fibroblasts (HFFs, ATCC®, SCRC-1041™) as described previously (Zhang et al., 2021). The HFF cells were grown in Dulbecco’s Modified Eagle medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 10 mM HEPES (pH 7.2), 100 µg/ml streptomycin and 100 U/ml penicillin, and maintained at 37°C with 5% CO₂.

Construction of Knockout Strains

FabZ, ODSCI, and ODSCII knockout strains were constructed using CRISPR-Cas9 technology as described previously (Wang et al., 2020). Briefly, the single-guide RNA (sgRNA) of each gene was inserted into the template pSAG1-Cas9-SgUPRT plasmid using Q5 mutation kit (NEB) to replace the target RNA. The plasmid was sequenced and the positive plasmid was extracted using Endo-Free Plasmid DNA Mini Kit (Omega, Georgia, United States). The 5’ and 3’ homologous fragments of each gene were amplified from the genomic DNA of T. gondii RH strain using the designed 5’ and 3’ homologous arm specific primers. All the primers used in this study are listed in Supplementary Table 1. The DHFR resistant fragments were amplified using the pUPRT-DHFR-D plasmid as a template, and the three fragments (5’ and 3’ homologous arm and DHFR fragment) were cloned into the plasmid pUC19 employing the multi-fragment cloning approach using the CloneExpress II one-step cloning kit (Vazyme Biotech, Nanjing, China). The 5’-HR-DHR-3HR fragment was amplified using the positive plasmid as a template to produce a homologous template, which was purified by agarose gel electrophoresis. About 40 µg of the CRISPR-Cas9 plasmid and 15 µg of the purified 5’HR-DHR-3HR fragment were co-transfected into freshly egressed T. gondii tachyzoites by electroporation. Finally, the clonal strain was obtained using a limiting dilution assay and 96-well plastic culture plates containing 3 µM pyrimethamine. The knockout strains were verified by PCR analysis. PCR 1 and PCR 3 were amplified from 5’ to 3’ of FabZ, ODSCI, and ODSCII by PCR1F/R and PCR3F/R, respectively, to verify whether the DHFR resistant fragment was successfully inserted into the corresponding knockout strain. PCR2F/R amplified product was about 700 bp in length, the knockout primers were designed at the upstream and downstream of FabZ, ODSCI, and ODSCII sgRNA-F to verify the successful knockout of...
the three genes (PCR2), and the wild-type (WT) strain was used as the control. The PCR reaction conditions were as follows: initial denaturation at 95°C for 4 min, followed by 35 amplification cycles (denaturation at 95°C for 30 s, annealing at 56°C for 30 s and extension at 72°C for 1 min), and a final extension step at 72°C for 10 min. The amplified products were examined on a 1.5% agarose gel with TAE buffer containing ethidium bromide.

C-Terminal Tagging
The C-Terminal HA epitope tagging was used to establish the subcellular localization of FabZ, ODSCI, and ODSCII. CRISPR-Cas9 plasmids targeting the 3’ region of each gene were constructed as previously described (Cao et al., 2019). Using pLIC-3 × HA-DHFR plasmid as a template, PCR products containing about 1.5 kb (except stop codon) and 3 × HA and DHFR fragments at the 3’ of the target gene were amplified using specific primers and purified using agarose gel electrophoresis. The purified fragment and the constructed C-terminal (3’ region) specific CRISPR-Cas9 plasmid were electro-transfected into freshly egressed tachyzoites. To identify single positive clones and verify whether the HA fragment was successfully inserted into T. gondii tachyzoites, we used PCR and specific primers to amplify the FabZ gene. We also used immunofluorescence assay (IFA) to localize FabZ and acyl carrier proteins (ACP). Briefly, HFFs were infected with 1 × 10⁶ tachyzoites of RHΔFabZ strain for 24 h. The cells were fixed with 4% paraformaldehyde (PFA) for 1 h, permealized for 30 min using 0.2% Triton X-100/phosphate-buffered saline (PBS), and blocked using 5% BSA for 30 min. Cells were incubated with primary antibodies (1:1,000), including mouse anti-SAG1 antibody to stain the tachyzoite's plasma membrane or anti-ACP antibody to stain the apicoplast, and rabbit anti-HA antibody, overnight at 4°C; followed by incubation with the secondary antibody (Alexa Fluor 594 goat anti-mouse IgG; 1:1,000) to stain SAG1 and ACP, or the secondary antibody (Alexa Fluor 488 anti-rabbit IgG; 1:1,000) to stain HA, for 1 h. Image analysis was performed using an TCS SP8 confocal scanning microscope (Leica, Germany).

Immunofluorescence Localization of Apicoplast
HFFs were infected with 1 × 10⁶ tachyzoites of each of the knockout strains (RHΔFabZ, RHΔODSCI, RHΔODSCII) and the WT RH strain and cultured at 37°C for 24 h. Subsequently, the cells were fixed with 4% paraformaldehyde (PFA) for 1 h, permealized for 30 min using 0.2% Triton X-100/PBS, and blocked using 5% bovine serum albumin (BSA) for 30 min. Cells were incubated with the primary antibodies (1:1,000), including rabbit anti-IMC1 (inner membrane complex) mouse antibody to stain the tachyzoites and anti-acyl carrier protein (ACP) antibody to stain the apicoplast, overnight at 4°C, followed by incubation with the secondary antibody (Alexa Fluor 488 anti-rabbit IgG; 1:1,000) to stain IMC1 or (Alexa Fluor 594 goat anti-mouse IgG; 1:1,000) to stain the ACP 37°C for 1 h. Image analysis was performed using an TCS SP8 confocal scanning microscope (Leica, Germany).

Plaque Assay
Approximately 500 tachyzoites of the three mutant RH strains and the WT strain were inoculated into 12-well culture plates containing HFF cell monolayers. Infected plates were incubated at 37°C and 5% CO₂ for 7 days. After removing the culture medium, the infected cells were fixed with 4% PFA for 1 h and stained with 0.5% crystal violet solution for 30 min. The number and size of plaques were counted and analyzed as described previously (Wang et al., 2020).

Intracellular Proliferation Assay
Tachyzoites (1 × 10⁵) of the three mutant strains and the WT strain were inoculated into cell culture dishes containing HFF cell monolayers. After 1 h, the medium was removed and the culture dishes were washed 3 times to remove any remaining extracellular tachyzoites. Fresh medium was then added to the infected cells and incubated at 37°C and 5% CO₂ for 24 h. The cells were fixed with 4% PFA for 1 h, and then stained with mouse anti-SAG1 and Alexa Fluor 594 goat-anti mouse IgG. Parasitophorous vacuoles (PVs) were randomly examined under a fluorescence microscope, and the number of tachyzoites inside the PVs was determined.

Egress Assay
Tachyzoites (1 × 10⁵) of the three mutant strains and the WT strain were inoculated into cell culture dishes containing HFF cell monolayers. After incubation at 37°C with 5% CO₂ for 1 h, cultures were washed 3 times with PBS to remove any extracellular tachyzoites. Fresh medium was then added and culture was future incubated for 36 h, and infected cells were treated with 3 μM calcium ionophore A23187 diluted in DMEM as described previously (Zhang et al., 2021). Live cell microscopy was used to monitor and image the tachyzoite egress from the host cell.

Quantitative Polymerase Chain Reaction (qPCR) Assay
Quantitative PCR (qPCR) was performed to quantify the expression of uracil phosphoribosyltransferase ([UPRT] [TGGT1_312480]) gene and apicoplast gene (TGGT1_302050) between WT strain and ΔFabZ strain. The 18 s rRNA gene was used as an internal reference for normalization. The qPCR assay was performed as described previously (Wu et al., 2009; Yeh and DeKisi, 2011; Reiff et al., 2012; Bansal et al., 2017). Genomic DNA was isolated from freshly egressed tachyzoites using Tiangen DNA extraction kit according to the manufacturer's directions. The apicoplast and UPRT-specific gene sequences were amplified using Max Super-Fidelity DNA Polymerase (Vazyme). The amplification reaction mixture included 10 μl of 2 × SYBR Green pro Tág HS Premix (final concentration 1 ×) (Accurate Biology), 0.4 μl of 10 μM of each primer, 40 ng of the extracted T. gondii DNA, 0.4 μl of ROX Reference Dye, and sterile water to a final volume of 20 μl. qPCR was performed using an
initial denaturation at 95°C for 30 s; followed by 40 cycles of amplification at 95°C for 10 s, 56°C for 20 s, and 72°C for 30 s.

Tachyzoite-Bradyzoite Transformation

Tachyzoites of the knockout strains (PRUΔFabZ, PRUΔODSCI, and PRUΔODSCII) and the WT PRU strain were used to infect the HFF cells cultured in alkaline RPMI-1640 medium (pH 8.2). After 3 days, the cells were fixed with 4% PFA for 1 h, subjected to permeabilization with 0.2% Triton X-100/PBS for 30 min at room temperature. Cells were blocked with PBS plus 5% BSA for 30 min and then incubated with the primary antibody (rabbit anti-IMC1) (1:1,000) overnight at 4°C, washed, and then incubated with goat anti-rabbit IgG Alexa Fluor 594 (1:1,000, Invitrogen, California, United States) 37°C for 1 h. Fluorescein isothiocyanate (FITC)-conjugated Dolichos biflorus lectin (DBL) was used to show the cyst wall (green). Slides were analyzed using a TCS SP8 confocal scanning microscope (Leica, Germany). We also quantified the rate of tachyzoite-bradyzoite transformation by subjecting *T. gondii*-infected HFF cells to immunostaining using DBL (a bradyzoite-specific marker) and IMC1 (stain of total parasites) as described previously (Xia et al., 2018). At least 100 PVs were examined in each parasite strain, and the experiment was repeated at least 3 independent times. The transformation rate of bradyzoites was calculated by dividing the number of DBL positive PVs by the number of IMC1 positive PVs.

Characterization of RHΔFabZ, RHΔODSCI, RHΔODSCII in Mice

Female specific-pathogen-free (SPF) Kunming mice (7-weeks-old) were purchased from the Center of Laboratory Animals, Lanzhou Veterinary Research Institute, Chinese Academy of Agricultural Sciences. Mice (8 mice/cage/group) were housed in an enriched environment with 12-h light/dark cycle. The food and water were provided *ad libitum*. Mice were infected intraperitoneally (i.p.) with freshly egressed tachyzoites of the respective mutant strains or WT strains. The number of tachyzoites of RH or PRU strain used in the inoculation was 100 and 5 × 10⁴ per mouse, respectively. The level of disease severity in RH and PRU-infected mouse groups was carefully monitored twice daily. Well-defined humane endpoints were used to identify mice that must be euthanized to avoid unnecessary suffering. The number of cysts in the brain homogenates of the PRU-infected mice that survived until 30 days after infection were recorded as described previously (Wang et al., 2018).

Bioinformatics Analysis

Information on the genomic characteristics of each gene, *FabZ*, *ODSCI*, and *ODSCII*, including the number of signal peptides, exons and transmembrane domains, and the temporal expression during the parasite cell cycle stages and in different *T. gondii* genotypes were obtained from ToxoDB.¹

Data Analysis

The data were analyzed by GraphPad Prism 7 (GraphPad Software, La Jolla, CA, United States). All the experiments were repeated 3 times and data are presented as means ± standard errors of the means (SEMs). The *F*-test was used for equal variance detection, and the means were compared using Student’s *t*-test or the Wilcoxon test.

RESULTS AND DISCUSSION

In this study we investigated whether deletion of *FabZ*, *ODSCI*, and *ODSCII* genes involved in the FASII pathway, using CRISPR-Cas9 system, influences the *in vitro* growth and *in vivo* virulence of *T. gondii*.

Localization of *FabZ*, *ODSCI*, and *ODSCII*

The *FabZ*, *ODSCI*, and *ODSCII* were tagged with three HA epitopes at the C terminus to characterize their cellular localization. The results of IFA analysis showed that for *FabZ*, a fluorescence signal can be observed in the apicoplast, indicating that *Tg*-FabZ is expressed in the apicoplast in the WT strain (Figures 1A,B). In contrast, no fluorescence signal was detected in the apicoplast in case of *Tg*-ODSCI and *Tg*-ODSCII (data not shown), possibly due to their low expression in the tachyzoite stage.

Construction of ΔFabZ, ΔODSCI, and ΔODSCII Strains

To study the biological function of *FabZ*, *ODSCI*, and *ODSCII* in *T. gondii*, we deleted the corresponding encoding genes in RH and PRU strains using CRISPR-Cas9 system. The coding regions of *FabZ*, *ODSCI*, and *ODSCII* were replaced by 5HR-DHFR-3HR fragments using homologous recombination (Figure 2A). Then, the constructed ΔFabZ, ΔODSCI, and ΔODSCII strains were confirmed by PCR analysis. PCR1 and PCR3 were used to establish whether DHFR was successfully

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¹http://toxodb.org

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**Figure 1** | Colocalization of *FabZ* and apicoplast in *T. gondii* as determined by indirect immunofluorescence microscopy. (A) Colocalization of acyl carrier protein (ACP) in the apicoplast (red) and *TgFabZ*-HA (green). (B) Colocalization of the plasma membrane protein SAG1 of *T. gondii* tachyzoites (red) and *TgFabZ*-HA (green). The results indicate that *FabZ* is located in the apicoplast. Scale bars, 10 µm.
inserted into the target gene sequence. The PCR products were detected in the clonal mutant strains, but not in the parental RH or PRU strain. PCR2 was used to verify that the target gene is successfully deleted, where the amplified fragments were detected in the parental strain but not in the knockout strain (Figures 2B,C). These results showed that mutant strains, ΔFabZ, ΔODSCI, and ΔODSCII, were successfully constructed.

The Effects on the in vitro Growth of T. gondii

In the present study, we determined whether FabZ, ODSCI, and ODSCII play a role in the growth of T. gondii by using the plaque assay, replication assay and egress assay. Significantly smaller and fewer plaques were formed by ΔFabZ strain compared to that produced by WT RH strain. There was no difference in the size and number of plaques formed by ΔODSCI or ΔODSCII when compared with WT RH strain (Figures 3A,B). These results show that lack of FabZ gene causes a considerable reduction in the growth of the RH strain; however, deletion of ODSCI and ODSCII did not have any impact on the rate of the parasite’s growth.

The results of intracellular proliferation assay showed that there were mainly 2 and 4 parasites in the PVs of ΔFabZ strain, and the number of PVs with 8 and 16 tachyzoites were significantly less than those detected in the PVs produced by the WT strain (Figure 3C), indicating that FabZ disruption significantly inhibited the replication of T. gondii. In contrast, ODSCI and ODSCII do not contribute to the parasite growth in vitro (Figure 3C). A similar finding was detected in P. yoelii where deletion of FabZ, a key enzyme in the FASII pathway, was found to prevent P. yoelii from forming merozoites during the liver infection stage (Vaughan et al., 2009).

We examined whether the deletion of FabZ, ODSCI, and ODSCII had an impact on T. gondii egress. The egress of T. gondii occurs when there are 64 or more tachyzoites inside the PV, and calcium is a key factor in mediating this process. 3 µM calcium ionophore A23187 was used to stimulate the tachyzoites release from the PV (Black et al., 2000a; Black and Boothroyd, 2000b; Arrizabalaga et al., 2004; Caldas et al., 2009). The results showed that most tachyzoites of ΔFabZ, ΔODSCI, or ΔODSCII and WT strain egressed within 2 min, indicating that the knockout of these three genes do not have any impact on the egress of T. gondii (Figure 3D).
FIGURE 3 | Characterization of the growth kinetics of the knockout strains in vitro. (A) Representative images of the plaque assay of HFFs infected with RH WT strain and mutant strains (RHΔFabZ, RHΔODSCI, and RHΔODSCII) for 7 days. (B) Plaque size presented as arbitrary units (a. u.). The relative size of RHΔFabZ plaque was significantly smaller and the number of plaque was fewer than that of RH WT strain, and there was no significant difference between RHΔODSCI or RHΔODSCII and RH WT strain. (C) The replication of T. gondii in HFFs was quantified by counting the number of T. gondii tachyzoites (1, 2, 4, 8, 16) per vacuole 24 h after infection with RH WT strain, RHΔFabZ, RHΔODSCI, or RHΔODSCII strain. RHΔFabZ replication was significantly inhibited, while RHΔODSCI and RHΔODSCII strains had similar intracellular replication kinetics to WT strain. (D) HFFs were infected with RH WT strain, RHΔFabZ, RHΔODSCI, or RHΔODSCII strain for 36 h, followed by treatment with 3 µM calcium ionophore A23187 to monitor the egress of tachyzoites at 0, 1, and 2 min after treatment. The results showed that the egressed pattern of WT strain was similar to that of all mutant strains. The data are presented as means ± SEMs (n = 3 in triplicates). ***P < 0.001; ns, non-significant by Student’s t-test and Gehan-Breslow-Wilcoxon test.
Effect of *FabZ*, *ODSCI*, and *ODSCII* Deletion on Apicoplast of *T. gondii*

The ACP fluorescent signal was hardly observed in the apicoplast of Δ*FabZ* tachyzoites, but was clearly detected in the tachyzoites of Δ*ODSCI* or Δ*ODSCII* and WT strains (Figure 4A). We further investigated the role of Δ*FabZ* in the stability of the apicoplast genome by comparing the replication of the apicoplast gene and the nuclear gene in Δ*FabZ* compared with WT by using qPCR. The results showed that compared with WT strain, the apicoplast gene expression of Δ*FabZ* was significantly decreased (Figure 4B), indicating that the presence of *FabZ* is essential for the maintenance of apicoplast genome of *T. gondii*.

In this study, the apicoplast gene (TGGT1_302050) and UPRT gene (TGGT1_312480) of *T. gondii* were successfully amplified. We also found that the apicoplast of Δ*FabZ* was lost, and the replications of the apicoplast genome, and the growth and intracellular proliferation were significantly reduced, indicating that FabZ plays a key role in the growth of *T. gondii*. These results show that the interruption of FASII pathway in the apicoplast by deleting *FabZ* may be the cause of the observed growth defects in Δ*FabZ* strain.

Effect of Deletion of *FabZ*, *ODSCI*, and *ODSCII* on Phenotypic Transformation

After incubating tachyzoites with RPMI-1640 (pH 8.2) for 3 days, the parasite cyst wall and tachyzoites were stained with DBL and IMC1, respectively. IFA results showed that *FabZ*, *ODSCI*, and *ODSCII* do not contribute to the tachyzoite-bradyzoite transformation process. No significant difference was detected in the tachyzoite-bradyzoite transformation rate between any of the examined *T. gondii* WT and mutant PRU strains (Figure 5B).

The Virulence of Δ*FabZ*, Δ*ODSCI*, and Δ*ODSCII* in Mice

We examined the impact of deletion of *FabZ*, *ODSCI*, and *ODSCII* on the pathogenicity of *T. gondii*. Approximately 100 tachyzoites of each strain, including RH, RHΔ*FabZ*, RHΔ*ODSCI*, and RHΔ*ODSCII*, were injected into Kunming mice by i.p. inoculation. The mice infected by RHΔ*FabZ* strain reached their humane endpoint up to 15 dpi, while mice infected by RHΔ*ODSCI*, RHΔ*ODSCII*, or WT strain reached their humane endpoint up to 11 dpi (Figure 6A). The survival rate of mice inoculated with wild-type PRU was 62.5%, and the survival rates of mice inoculated with knockout strains PRUΔ*FabZ*, PRUΔ*ODSCI*, and PRUΔ*ODSCII* were 50, 50, and 62.5%, respectively, indicating that *FabZ*, *ODSCI*, and *ODSCII* genes do not play any role in the virulence of *T. gondii* PRU strain in mice (Figure 6B).

We examined the effects of PRUΔ*FabZ*, PRUΔ*ODSCI*, and PRUΔ*ODSCII* on the cyst formation of *T. gondii*. Approximately 5 × 10⁴ tachyzoites of WT PRU and each of the three mutant strains were injected by the i.p. route into Kunming mice. After 30 days the number of cysts in the brain of the survived mice was counted. As shown in Figure 6C, the number of cysts in mice infected by PRUΔ*FabZ* (51.25 ± 3.198) was significantly (P < 0.01) lower than that in mice infected by WT (103.7 ± 8.686), while the number of cysts in mice infected with PRUΔ*ODSCI* and PRUΔ*ODSCII* was not significantly different from that of the WT-infected mice.

Previous studies have shown that ACP is a key enzyme in the FASII pathway, and that deletion of the corresponding gene leads to defects in the growth of parasites *in vitro*, and increases the survival time of mice infected with mutant strain compared to mice infected by WT strain (Behnke et al., 2006; Mazumdar et al., 2006). Our results showed that deletion of *FabZ* induced defects in the growth of *T. gondii* *in vitro*, improved the survival of mice infected by the RH strain, and decreased the number of brain cysts in mice infected by PRUΔ*FabZ*

FIGURE 4 | The effect of gene knockout on the apicoplast of *T. gondii* RH strain. (A) Immunofluorescence localization of the anti-acyl carrier protein (ACP) in the apicoplast using an apicoplast-specific antibody. The ACP specific fluorescence was hardly detected in the apicoplast of tachyzoites of Δ*FabZ*, but was clearly visible in the tachyzoites of Δ*ODSCI*, Δ*ODSCII*, and WT strains. Scale bars, 10 μm. (B) Quantitative PCR was performed to detect apicoplast and nuclear gene in Δ*FabZ* and WT RH strains. The expression of the apicoplast gene in Δ*FabZ* strain was significantly decreased. The data are presented as means ± SEMs (n = 3 in triplicates). *P < 0.05; ns, non-significant by Student’s t-test and Gehan-Breslow-Wilcoxon test.
FIGURE 5 | Tachyzoite-bradyzoite transformation of ΔFabZ, ΔODSCI, or ΔODSCII using immunofluorescence analysis. HFFs were infected with PRU WT, PRUΔFabZ, PRUΔODSCI, or PRUΔODSCII strains and maintained in alkaline medium (pH = 8.2) for 3 days, then the parasites were stained with anti-IMC1 (inner membrane complex) and Alexa Fluor 594 (red). The parasite cyst walls were stained with FITC-conjugated Dolichos biflorus lectin, DBL (green). (A) The deletion of FabZ, ODSCI, or ODSCII in PRU strain did not have any remarkable impact on the tachyzoite-bradyzoite transformation ability. Scale bars, 10 µm. (B) No significant difference in the parasite stage transformation was detected between any of the tested parasite strains.

FIGURE 6 | The virulence and cyst forming ability of ΔFabZ, ΔODSCI, or ΔODSCII in vivo. (A) The survival of Kunming mice inoculated by 100 tachyzoites of the RH (mutant or wild-type) strains. (B) The survival of Kunming mice infected by 5 × 10⁴ tachyzoites of the PRU (mutant and wild-type) strains. Mice infected by RH (A) or PRU (B) strain were checked twice daily by two independent observers for clinical signs and a Kaplan Meier survival curve plotted the infected mice that reached their humane endpoints. (C) The number of cysts in the mouse brain homogenates were counted at 30 days after mice were infected by 5 × 10⁴ PRU tachyzoites. The data are shown as means ± SEMs (8 mice/group). **P < 0.01; ns, non-significant by Student’s t-test.
Expression profiles of FabZ, ODSCI, and ODSCII in *T. gondii*. (A) The temporal expression profile of FabZ, ODSCI, and ODSCII during the cell cycle of *T. gondii* RH strain. (B) The expression profile of FabZ, ODSCI, and ODSCII of *T. gondii* in type I (RH and GT1), type II (PRU and ME49) and type III (CTG and VEG) strains of *T. gondii*.

### Table 1: Bioinformatic characteristics of Toxoplasma gondii genes.

| Name   | Gene ID         | Product description                                      | Exons | Phenotype value | TMHMM | Predicted signal peptide |
|--------|-----------------|----------------------------------------------------------|-------|-----------------|-------|--------------------------|
| FabZ   | TGME49_321570   | Beta-hydroxyacyl-acyl carrier protein dehydratase         | 2     | 0.67            | No    | Yes                      |
| ODSCI  | TGGT1_269400    | Oxidoreductase, short chain dehydrogenase/reductase family protein | 5     | 0.08            | No    | No                       |
| ODSCII | TGGT1_313050    | Oxidoreductase, short chain dehydrogenase/reductase family protein | 6     | 0.37            | No    | No                       |

*Prediction of transmembrane helices was performed using the TMHMM program version 2.0.*

cysts of the PRU strain, indicating the important role that FabZ plays in the growth and virulence of *T. gondii*.

### Expression Profiles of FabZ, ODSCI, and ODSCII

Many genes exhibit temporal expression patterns during the replication of *T. gondii* tachyzoites, which is related to biological processes specific to the parasite development and replication (Behnke et al., 2006). Previous studies have found that some ROP genes, such as *ROP5, ROP16,* and *ROP17,* are cell cycle-dependent (Behnke et al., 2006). However, some ROPs do not show periodic expression characteristics, such as *ROP21, ROP27,* and *ROP28* (Jones et al., 2017).

The expression profiles of FabZ, ODSCI, and ODSCII genes involved in FASII pathway were analyzed under different conditions according to the data available in ToxoDB. There was no difference in the expression of FabZ at different time points in the parasite's respective cell cycle stages, but the expression of FabZ was significantly higher than that of ODSCI and ODSCII at the same time points (Figure 7A) (Behnke et al., 2010). Likewise, there was no difference in the expression of each gene between strains of the three main *T. gondii* genotypes (Type I, Type II, and Type III); however, the expression of FabZ was higher than that of ODSCI and ODSCII in the same examined strains (Figure 7B).

Studies have shown that the biological characteristics of some genes in *T. gondii* are expressed from a single exon and contain predicted signal peptides, and most proteins enter the secretory pathway mediated by signal peptides in eukaryotic cells (Devillers-Thiery et al., 1975; Lorenzi et al., 2016). The bioinformatics features, including the exon number, signal peptide and transmembrane domain of FabZ, ODSCI, and ODSCII are summarized in Table 1. Most of the proteins that perform biological functions in the apicoplast of *T. gondii* are encoded by the nuclear genome and transported to the apicoplast. Studies have shown that the signal peptide of the malaria parasite can transport proteins to the parasite's apicoplast (Meyer et al., 2018). The present and previous studies detected specific signals of FabZ and ACP (Waller et al., 1998, 2000) in the apicoplast. These results suggest that FabZ and ACP, encoded by the nuclear genome of *T. gondii*, are transported to the apicoplast likely via signal peptides to exert their functions (Waller et al., 1998, 2000).

### Conclusion

In this study, FabZ, ODSCI, and ODSCII involved in FASII pathway of the apicoplast were deleted using CRISPR-Cas9 system, and the *in vitro* growth and *in vivo* pathogenicity of the mutant strains were studied. Deletion of FabZ caused the loss of the apicoplast and reduced the growth of *T. gondii* *in vitro*, while deletion of ODSCI and ODSCII had no effect on the parasite's growth. Deletion of FabZ reduced the virulence of the RH strain and decreased the brain cyst burden of the PRU strain.
in Kunming mice. Bioinformatics analysis showed that \textit{FabZ} is likely to be transported to the apicoplast, via its signal peptides, to participate in the FASII pathway. These findings show that \textit{FabZ} plays an important role in the pathogenesis of \textit{T. gondii} infection.

**DATA AVAILABILITY STATEMENT**

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s.

**ETHICS STATEMENT**

The animal study was reviewed and approved by the Animal Ethics Committee of Lanzhou Veterinary Research Institute, Chinese Academy of Agricultural Sciences.

**AUTHOR CONTRIBUTIONS**

X-QZ, M-XS, and HE conceived and designed this study, and critically revised the manuscript. X-PX performed the experiments, analyzed the data, and drafted the manuscript. W-GL, Z-WZ, L-XS, and Q-LL participated in the implementation of the study. All authors read and approved the final version of the manuscript.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2021.703059/full#supplementary-material

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