Identifying the major lactate transporter of *Toxoplasma gondii* tachyzoites

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*Toxoplasma gondii* and *Plasmodium falciparum* parasites both extrude L-lactate, a byproduct of glycolysis. The *P. falciparum* Formate Nitrite Transporter, *PfFNT*, mediates L-lactate transport across the plasma membrane of *P. falciparum* parasites and has been validated as a drug target. The *T. gondii* genome encodes three FNTs that have been shown to transport L-lactate, and which are proposed to be the targets of several inhibitors of *T. gondii* proliferation. Here, we show that each of the *TgFNTs* localize to the *T. gondii* plasma membrane and are capable of transporting L-lactate across it, with *TgFNT1* making the primary contribution to L-lactate transport during the disease-causing lytic cycle of the parasite. We use the *Xenopus* oocyte expression system to provide direct measurements of L-lactate transport via *TgFNT1*. We undertake a genetic analysis of the importance of the *tgfnt* genes for parasite proliferation, and demonstrate that all three *tgfnt* genes can be disrupted individually and together without affecting the lytic cycle under in vitro culture conditions. Together, our experiments identify the major lactate transporter in the disease causing stage of *T. gondii*, and reveal that this transporter is not required for parasite proliferation, indicating that *TgFNTs* are unlikely to be targets for anti-*Toxoplasma* drugs.

*Toxoplasma gondii* and *Plasmodium falciparum* are unicellular protozoan parasites that belong to the phylum Apicomplexa. *P. falciparum* is the most deadly of the *Plasmodium* species that cause malaria in humans¹. *T. gondii* infects a large proportion of the world’s population and can cause severe disease in immunocompromised individuals¹. *T. gondii* can also have devastating effects on the development of foetuses when it infects women during pregnancy². New medicines are needed urgently against both parasites.

The disease-causing tachyzoite stage of *T. gondii* parasites utilizes both glucose and glutamine as energy sources for the generation of ATP³,⁴. These parasites acquire glucose through a plasma-membrane localized Glucose Transporter (*TgGT1*)³. Glucose is metabolized via glycolysis to generate pyruvate. Pyruvate can have multiple possible fates in the parasite⁵. It can be transported into the mitochondrion, where it is further metabolized by the TCA cycle to generate ATP by oxidative phosphorylation⁶. Alternatively, pyruvate can be metabolized to form L-lactate, which is then exported from the parasite⁷,⁸. Lactate synthesis is catalyzed by lactate dehydrogenase (LDH). The *T. gondii* genome encodes two LDH enzymes (*TgLDH1* and *TgLDH2*), neither of which is required for proliferation of the disease-causing tachyzoite stage under standard in vitro culture conditions⁹,¹⁰. *T. gondii* tachyzoites that are rendered defective in glucose import or some aspects of glycolysis can still grow in vitro and in vivo, catabolizing glutamine via the TCA cycle to produce ATP and generating essential glycolytic intermediates via gluconeogenesis⁷,¹¹,¹². When glucose and glutamine are both available, both are used for ATP generation⁷.

By contrast, the disease-causing blood stages of *P. falciparum* parasites rely almost exclusively on glycolysis to generate ATP¹³. Here, L-lactate is the major product of glycolysis and must be exported from the parasite in symport with H⁺ by the *P. falciparum* Formate Nitrite Transporter (*PfFNT*) present on the parasite’s plasma membrane¹⁴. Members of the FNT family transport a variety of monocarboxylates and other anions¹⁵. They are found in numerous prokaryotic and eukaryotic microorganisms, but are not present in mammalian cells¹⁶. Compounds that kill *P. falciparum* parasites via inhibition of *PfFNT* have recently been identified, thereby validating *PfFNT* as a novel antimalarial drug target²⁰,²¹.

A recent study provided evidence that the three homologues of *PfFNT* encoded in the *T. gondii* genome are L-lactate transporters that localize to the plasma membrane when expressed from a non-native promoter²².

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Compounds found to inhibit the activity of the transporters showed some inhibition of tachyzoite proliferation at micromolar concentrations, leading to the suggestion that the TgFNTs might be essential during the T. gondii lytic cycle and ‘druggable’\textsuperscript{22}. The possibility that the compounds inhibited parasite growth via alternate means (‘off-target effects’) was not excluded. Although T. gondii tachyzoites can survive without glucose, it is possible that in the presence of glucose, the lack of a l-lactate efflux mechanism would cause l-lactate to accumulate intracellularly to concentrations sufficient to jeopardize pH regulation and/or osmotic stability. However, a genome-wide screen has provided evidence that each of the TgFNTs are dispensable during the lytic cycle\textsuperscript{23}, and a TgFNT1 knockout parasite line was recently generated and found to grow normally despite secreting less lactate into the medium than wild-type parasites\textsuperscript{24}. Here, we employed genetic and physiological approaches to investigate the roles of the three TgFNTs in situ and to determine whether the activity of one or more TgFNTs is required for the proliferation of T. gondii tachyzoites.

**Results**

**TgFNTs localize to the parasite plasma membrane.** The T. gondii genome encodes three members of the FNT family, termed TgFNT1 (TGGT1_209800), TgFNT2 (TGGT1_292110) and TgFNT3 (TGGT1_229170)\textsuperscript{22}. The TgFNTs share considerable sequence similarity with each other, with PfFNT, and with the well-characterized E. coli FNT family member FocA (Supplementary Fig. S1).

To investigate the expression and localization of each TgFNT, we attempted to generate parasite lines in which a single hemagglutinin (HA) tag was fused to the 3′ end of the open reading frame of the encoding gene. This approach was successful for TgFNT1 (Supplementary Fig. S2). Western blotting with an anti-HA antibody revealed that the resultant TgFNT1-HA protein is expressed in tachyzoites and that it had the expected mass of approximately 45 kDa (Fig. 1a). Immunofluorescence assays revealed that TgFNT1-HA localizes to the T. gondii plasma membrane (Fig. 1b).

Our attempts to generate parasites expressing detectable levels of TgFNT2-HA or TgFNT3-HA with this approach were not successful. An alternate approach was therefore used to localize TgFNT2 and TgFNT3. Parasites were transiently transfected with a vector containing the open reading frame of the encoding gene fused at the 3′ end to a 3 × HA tag, under the regulation of the constitutive α-tubulin promoter. Consistent with the findings of Erler et al.\textsuperscript{25}, the ectopically expressed TgFNT2-HA\textsubscript{3} and TgFNT3-HA\textsubscript{3} proteins both localized to the T. gondii plasma membrane (Fig. 1c,d).

**TgFNTs are not required for tachyzoite proliferation in vitro.** To determine whether any of the TgFNTs are important for parasite proliferation, we used CRISPR-Cas9 targeted genome editing to generate...
parasites in which the open reading frame of the *tgfnt1*, *tgfnt2* or *tgfnt3* gene was disrupted by frameshift mutations. For *tgfnt1*, we selected a clone (‘Δ*tgfnt1*’) with a single nucleotide insertion at position 176 from the start codon of the gene. For *tgfnt2*, we selected a clone (‘Δ*tgfnt2*’) in which nucleotide 126 was deleted, and for *tgfnt3* (‘Δ*tgfnt3*’) we selected a clone with an insertion of five nucleotides from position 123 of the gene. In all cases, the resulting frameshifts led to the introduction of premature stop codons and truncation of the encoded proteins (Supplementary Fig. S4), which are likely to render the resulting proteins non-functional. We also generated a parasite line in which all three *tgfnt* genes were disrupted. This was performed by first targeting the *tgfnt1* gene in Δ*tgfnt3* parasites. We selected a clone (‘Δ*tgfnt1/3*’) that had a single nucleotide insertion at position 176 of the *tgfnt1* gene. Next, we targeted the *tgfnt2* gene in Δ*tgfnt1/3* parasites. We selected a clone (‘Δ*tgfnt1/2/3*’) with a single nucleotide insertion at position 127 of the *tgfnt2* gene. Each of the *tgfnt* genes in Δ*tgfnt1/2/3* parasites has a premature stop codon (Supplementary Fig. S4).

We tested whether any of the *TgFNT* isoforms were important for parasite proliferation. We first assessed this using plaque assays. Plaques (zones of clearance) in a host cell monolayer result from multiple cycles of host cell invasion, replication and lysis, with the size of the plaques providing an indication of parasite growth rate. Δ*tgfnt1*, Δ*tgfnt2*, Δ*tgfnt3*, and Δ*tgfnt1/2/3* parasites all produced plaques comparable in size to those of their parents (Supplementary Fig. S5), indicating no significant growth defect in any of the four knockout lines.

We also compared the growth rates of the different parasites using a fluorescence assay (described previously25), taking advantage of the fact that all the parasites express a tandem dimeric Tomato (tdTomato) red fluorescent protein. In each experiment, a sigmoidal growth curve was generated for each parasite line, with a lag phase, exponential growth phase and plateau (Fig. 2a). Using the maximum slope of the growth curve as a read-out for parasite proliferation, we found no differences in growth rates among Δ*tgfnt1*, Δ*tgfnt2*, Δ*tgfnt3*, and Δ*tgfnt1/2/3* parasites all produced plaques comparable in size to those of their parents (Supplementary Fig. S5), indicating no significant growth defect in any of the four knockout lines.

We compared the growth rates of the different parasites using a fluorescence assay (described previously25), taking advantage of the fact that all the parasites express a tandem dimeric Tomato (tdTomato) red fluorescent protein. In each experiment, a sigmoidal growth curve was generated for each parasite line, with a lag phase, exponential growth phase and plateau (Fig. 2a). Using the maximum slope of the growth curve as a read-out for parasite proliferation, we found no differences in growth rates among Δ*tgfnt1*, Δ*tgfnt2*, Δ*tgfnt3*, Δ*tgfnt1/2/3*, and wild-type parental parasites (Fig. 2b). We conclude that none of the *TgFNT* isoforms, individually or collectively, are required for the normal proliferation of tachyzoites under in vitro culture conditions.

*TgFNTs* transport l-lactate, with *TgFNT1* making the primary contribution to l-lactate transport across the plasma membrane in extracellular tachyzoites. We used the *tgfnt* knockout parasites to investigate whether one or more of the *TgFNTs* provide a route for l-lactate transport across the plasma membrane in *T. gondii*. The parasite experiments described in this section were all performed with extracellular tachyzoites that had either recently emerged from their host cells or that had been manually released from their host cells by passage of the cell culture through a 26G needle. First, we investigated the effect on parasite cytosolic pH of adding a high concentration of l-lactate to the extracellular solution. This approach has been used previously with *P. falciparum* parasites to investigate PfFNT, which transports l-lactate in symport with H+ ions25,26.
Although the physiological role of PfFNT is to remove l-lactate from the parasite cytosol, it is bidirectional. Thus, when a high concentration of l-lactate is added to the extracellular solution to impose an inward l-lactate concentration gradient, a decrease in cytosolic pH resulting from l-lactate:H+ entry can be detected15,21. We measured cytosolic pH in extracellular T. gondii tachyzoites by loading them with the pH-sensitive dye BCECF. The measurements were performed at 4 °C to reduce the rate at which pH-regulatory mechanism(s) (including the H+-extruding V-type H+ ATPase26,27) counteract the l-lactate-mediated pH change. When the wild-type parental parasites suspended in a physiological saline solution were exposed to 10 mM l-lactate, we observed a rapid decrease in the pH of the parasite cytosol (Fig. 3a). An abrupt l-lactate-induced decrease in cytosolic pH was also seen in Δtgfnt2 (Fig. 3c) and Δtgfnt3 (Fig. 3d) parasites. In contrast, the addition of 10 mM l-lactate to Δtgfnt1 parasites had little effect on the pH of the cytosol (Fig. 3b), consistent with parasites lacking TgFNT1 having a greatly reduced capacity for l-lactate:H+ symport across the plasma membrane. The subsequent addition of the protonophore CCCP (10 µM) dissipated the H+ gradient across the plasma membrane, with the cytosolic pH decreasing to a value close to the extracellular pH (Fig. 3b), demonstrating the responsiveness of the pH-detection system in these cells.

To confirm that the reduction in l-lactate:H+ transport in Δtgfnt1 parasites was a consequence of the absence of TgFNT1 expression, we complemented Δtgfnt1 parasites with an ectopically-expressed copy of tgfnt1 (under the control of the constitutive a-tubulin promoter). In these parasites, the addition of 10 mM l-lactate to the extracellular solution caused a pronounced decrease in cytosolic pH (Fig. 3e). These data are consistent with TgFNT1 serving as the primary L-lactate:H+ transporter on the plasma membrane in extracellular tachyzoites.

We also investigated whether TgFNT2 or TgFNT3 could restore l-lactate:H+ transport in Δtgfnt1 parasites when constitutively expressed in these parasites. We found that Δtgfnt1 parasites complemented with either tgfnt2 or tgfnt3 under the control of the a-tubulin promoter underwent a decrease in cytosolic pH on addition of 10 mM l-lactate to the extracellular solution (Fig. 3f,g). The rate of the l-lactate-induced cytosolic pH decrease was lower in Δtgfnt1 parasites complemented with tgfnt2 (Fig. 3f) than in parasites complemented with tgfnt1 (Fig. 3e) or tgfnt3 (Fig. 3g). The initial rates (ΔpH/min, mean ± SEM, n = 3; determined by fitting lines to the initial linear portions of the traces) were 0.82 ± 0.19, 0.18 ± 0.09, and 0.57 ± 0.10 for Δtgfnt1 parasites complemented with tgfnt1, tgfnt2 and tgfnt3, respectively. There was a significant difference between the rates of pH decrease for Δtgfnt1 parasites complemented with tgfnt2 and Δtgfnt1 parasites complemented with either tgfnt1 or tgfnt3 (P < 0.05, unpaired t-tests). This could result from a difference between TgFNT2 and the other TgFNTs with respect to transporter function, regulation, and/or plasma membrane expression. Together, our data show

Figure 3. Measurements of the effect of l-lactate on cytosolic pH (pHcyt) reveal a role for TgFNT1 in lactate transport across the parasite plasma membrane. Extracellular tachyzoites were loaded with BCECF and suspended in Saline Solution at 4 °C. The grey arrows denote the addition of 10 mM l-lactate. The red arrow denotes the addition of the protonophore CCCP (10 µM). The top panels show data for Δtgfnt1 (b), Δtgfnt2 (c), Δtgfnt3 (d) parasites and their wild-type parental parasites (a). The bottom panels show data for Δtgfnt1 parasites complemented with tgfnt1 (e), tgfnt2 (f) or tgfnt3 (g). The traces are from a single experiment for each parasite line, and are representative of those obtained in at least three independent experiments. The time taken to reach a minimum pHcyt value after the addition of l-lactate was >400 s in all experiments for Δtgfnt1 parasites complemented with tgfnt2, compared to <150 s in all experiments for WT parasites, Δtgfnt2 parasites, Δtgfnt3 parasites, and Δtgfnt1 parasites complemented with tgfnt1 or tgfnt3.
that knockout of TgFNT1 prevents l-lactate -induced pH changes in the parasite cytosol, which can be rescued by ectopic expression of TgFNT1, TgFNT2 and TgFNT3. Our data also indicate that TgFNT2 and TgFNT3 make minimal contribution to l-lactate:H+ transport in wild-type tachyzoites. It is possible that the native tgifnt2 and tgifnt3 genes are not expressed at significant levels during this stage of the parasite life cycle.

Our pH data implicate TgFNT1 as the primary l-lactate transporter in tachyzoites. Erler et al.22 provided evidence for l-lactate transport by TgFNT1-3 expressed in yeast, however truncation of the proteins was required to achieve their functional expression. To test whether full-length TgFNT1 is a l-lactate transporter, we expressed it in Xenopus oocytes. We have previously expressed PfFNT in this system and shown that it transports l-[14C]lactate15 and is inhibited by the antiplasmodial compound MMV00783921. We measured the uptake of l-[14C]lactate into Xenopus oocytes expressing full-length TgFNT1 (with a C-terminal HA tag) at 27.5 °C over 10 min at pH 6.4. The uptake of l-[14C]lactate by oocytes injected with 30 ng of crRNA encoding TgFNT1 was higher than that of non-injected oocytes and of oocytes injected with 30 ng of crRNA encoding PfFNT (P < 0.001; one-way ANOVA; Fig. 4). Consistent with previous results21, l-[14C]lactate uptake by PfFNT-expressing oocytes was inhibited by MMV007839 (4 µM; Fig. 4). In contrast, 4 µM MMV007839 had no effect on the uptake of l-[14C]lactate by TgFNT1-expressing oocytes (Fig. 4). These data provide direct evidence that TgFNT1 is a l-lactate transporter. The finding that TgFNT1 is less sensitive than PfFNT to inhibition by MMV007839 is consistent with the results of Erler et al.22, who found that 29 µM MMV007839 was required for half-maximal inhibition of l-lactate transport by a C-terminally truncated form of TgFNT1, whereas half-maximal inhibition of l-lactate transport by PfFNT is achieved at sub-micromolar concentrations20,21.

As a final test of the role of TgFNT1 in l-lactate transport across the plasma membrane in extracellular tachyzoites, we measured the uptake of l-[14C]lactate by extracellular Δtgfnt1, Δtgfnt1/2/3, and wild-type parental parasites. As noted above, FNTs mediate bidirectional transport, and measuring the movement of l-[14C]lactate into parasites provides a convenient means of assessing transport activity. Similar to previous experiments with P. falciparum15,21, the experiments with T. gondii presented here were conducted at a low pH (6.1) to increase (H+-coupled) l-lactate uptake by the parasites, and at a low temperature (4 °C) to slow the transport process. We found that the uptake of lactate, as measured over 10 min, was greatly reduced in Δtgfnt1 and Δtgfnt1/2/3 parasites relative to wild-type parental parasites, with little difference observed between Δtgfnt1 and Δtgfnt1/2/3 parasites (Fig. 5a). We estimated the initial rate of l-lactate uptake for each parasite line. The initial rate of l-lactate uptake was > 7.5-fold greater in wild-type parental parasites than in Δtgfnt1 or Δtgfnt1/2/3 parasites, and there was no significant difference in the initial rate between Δtgfnt1 and Δtgfnt1/2/3 parasites (Fig. 5b). Consistent with the pH data, these data indicate that TgFNT1 plays the primary role in l-lactate transport across the plasma membrane in wild-type extracellular parasites.

To investigate the possibility that disruption of the tgifnt1 gene could give rise to a general defect in solute uptake, we measured the uptake of two additional radiolabelled compounds: the glucose analogue [14C]2-deoxyglucose ([14C]2-DOG) and the amino acid l-[14C]arginine. The time courses for the uptake of
Figure 5. TgFNT1 is critical for l-lactate transport across the parasite plasma membrane. (a,c,e) Time courses for the uptake of l-lactate (a), 2-DOG (c) and L-arginine (e) by wild-type parasites (blue symbols), Δtgfnt1 parasites (white symbols) and Δtgfnt1/2/3 parasites (dark red symbols). The data shown are the mean ± SD from three to four independent experiments for each line. For clarity, only positive or negative error bars are shown for the data for certain parasite lines. Where not shown, error bars fall within the symbols. (b,d,f) The initial rates of uptake for l-lactate (b), 2-DOG (d), and L-arginine (f) estimated from the same set of experiments. The bars show the mean ± SD obtained from three or four independent experiments for each line, and the symbols show the results obtained in individual experiments. For panels b,d and f, the data for each parasite line were compared with the data for every other parasite line (one-way ANOVA, followed by post hoc Tukey tests where significant differences were present). ***P < 0.001; other comparisons (those in d and f) did not reveal significant differences.
tgfnt1 was greatly reduced in Δ in situ. We measured the rate of l-lactate transport across the parasite plasma membrane, and found that it Tg and physiological approaches we confirmed that TgFNT1 functions as a plasma membrane l-lactate transporter by a large inward l-lactate concentration gradient was not observed in Δ cytosolic pH of adding 10 mM l-lactate to the external medium. The decrease in cytosolic pH normally induced by a large inward l-lactate concentration gradient was not observed in Δtgfnt1 parasites, consistent with these parasites having a defect in l-lactate:H⁺ transport. Complementing Δtgfnt1 parasites with an ectopic copy of the tgfnt1 gene restored the l-lactate-induced pH change.

Consistent with Erler et al., we found that TgFNT2 and TgFNT3 localize to the plasma membrane when expressed ectopically under the control of a non-native promoter. Furthermore, in line with previous evidence that truncated forms of TgFNT1-3 transport l-lactate in yeast, our data suggest that (full-length) TgFNT2 and TgFNT3 can transport l-lactate. Complementing Δtgfnt1 parasites with constitutively-expressed tgfnt2 or tgfnt3 resulted in the restoration of l-lactate:H⁺ transport, as observed in measurements of cytosolic pH.

Our data indicate that, despite being capable of transporting l-lactate, TgFNT2 and TgFNT3 make little contribution to l-lactate transport across the plasma membrane in wild-type extracellular tachyzoites. Disrupting tgfnt1 alone was sufficient to abolish the l-lactate-induced acidification observed in parental parasites. Furthermore, there was no difference in the rate of l-[14C]lactate transport across the plasma membrane in Δtgfnt1 parasites and Δtgfnt1/2/3 parasites. Thus, expression of the endogenous tgfnt2 or tgfnt3 genes in Δtgfnt1 parasites does not appear to compensate for the l-lactate transport defect observed when TgFNT1 is not expressed. It is possible that TgFNT2 and TgFNT3 are not expressed at appreciable levels during the lytic cycle in the T. gondii strain used in this study. This would also explain our inability to localize these proteins by tagging the 3’ ends of the endogenous genes. Our data are partially supported by a recent proteome-wide localization study of tachyzoites, which predicted that TgFNT1 localizes to the plasma membrane whereas TgFNT3 was not detectable in the proteome. This study predicted that TgFNT2 may localize to micronemes, which is inconsistent with our localization studies.

A key finding in our study was that the lack of expression of all three TgFNT proteins does not affect the proliferation of T. gondii tachyzoites under standard in vitro culture conditions. This contrasts with the situation in P. falciparum, where PfFNT has been validated as a drug target and must therefore be important for parasite proliferation. However, it is consistent with previous studies highlighting the metabolic flexibility of T. gondii parasites, including recent evidence that the LDH enzymes required to convert pyruvate to lactate are not essential for T. gondii proliferation under standard in vitro culture conditions. Our data are also consistent with a genome-wide screen that provided evidence that each of the TgFNTs is dispensable during the lytic cycle (the ‘phenotype scores’ for TgFNT1, TgFNT4 and TgFNT3 were 1.10, 1.31 and -0.23, respectively), and with a very recent study in which TgFNT1 was knocked out. Kloehn et al. found that TgFNT1 knockout parasites grew normally in vitro, but that they secreted less lactate into the medium than wild-type parasites. Our finding that all three TgFNTs can be knocked out simultaneously call into question the view that the TgFNTs are suitable drug targets, and suggest that the compounds reported to inhibit the activity of TgFNT proteins by Erler et al. are unlikely to exert their effects on T. gondii growth via inhibition of any of the three transporters.

It should be noted that the disruption of certain glycolytic enzymes that are not essential for the in vitro proliferation of T. gondii parasites has been associated with virulence defects in mice and/or reductions in the formation of bradyzoite cysts in the brain. Parasites lacking hexokinase had a greatly reduced ability to form mature bradyzoite cysts in the brain in mice compared to parental parasites. Furthermore, while tachyzoites in which the genes encoding TgLDH1 and TgLDH2 were simultaneously disrupted proliferated normally under standard conditions, their growth was impaired in low-oxygen conditions in vitro and in mice, and the numbers of bradyzoite cysts formed in the brain were greatly reduced. Thus, it remains possible that the expression of one or more TgFNT proteins is important in vivo and/or in different stages of the T. gondii life cycle.

### Discussion

In this study we pinpoint TgFNT1 as the primary contributor to l-lactate transport across the plasma membrane of extracellular T. gondii tachyzoites. Tagging the 3’ end of the native tgfnt1 gene revealed that the encoded protein localizes to the parasite plasma membrane. Experiments in which TgFNT1 was studied in isolation in Xenopus oocytes provided direct evidence that full-length TgFNT1 mediates the transport of l-[14C]lactate. Using genetic and physiological approaches we confirmed that TgFNT1 functions as a plasma membrane l-lactate transporter in situ. We measured the rate of l-lactate transport across the parasite plasma membrane, and found that it was greatly reduced in Δtgfnt1 tachyzoites compared to wild-type parasites. We also investigated the effect on cytosolic pH of adding 10 mM l-lactate to the external medium. The decrease in cytosolic pH normally induced by a large inward l-lactate concentration gradient was not observed in Δtgfnt1 parasites, consistent with these parasites having a defect in l-lactate:H⁺ transport. Complementing Δtgfnt1 parasites with an ectopic copy of the tgfnt1 gene restored the l-lactate-induced pH change.

### Methods

**Ethics statement.** Ethical approval of the work performed with the adult female Xenopus laevis frogs was obtained from the Australian National University (ANU) Animal Experimentation Ethics Committee (Animal Ethics Protocol Number A2013/13) in accordance with the Australian Code of Practice for the Care and Use of Animals for Scientific Purposes.

**Host cell and parasite culture.** T. gondii parasites were cultured in human foreskin fibroblasts (a gift from Holger Schlüter, Peter MacCallum Cancer Centre) at 37 °C in a humidified 5% CO₂ incubator. The culture medium was Dulbecco’s modified Eagle’s medium (DMEM) containing 25 mM glucose, and supplemented with 1% v/v fetal calf serum, 200 μM L-glutamine, 50 U/mL penicillin, 50 µg/mL streptomycin, 10 µg/mL gentamicin, and 0.25 µg/mL amphotericin B. TATi/Tomato parasites were used as the parental wild-type strain for the
Δtgfnt parasite strains that we generated in this study. TATiΔku80 parasites were used as the parental wild-type strain for the 3’ replacement strains generated in this study.

**Generation of genetically modified T. gondii lines for localization studies.** To generate a 3’ HA tag replacement in the tgfnt1 locus, we amplified a 1.1 kb region of the 3’ end of the tgfnt1 open reading frame using primers p4 and p5 (Supplementary Table S1). The PCR product was digested with BglII and AvrII and ligated into the BglII and AvrII sites of the psgCHΔpsI1 vector, a variant of psgCH1 wherein the PsI site has been filled in. The resulting vector was linearized with SphI and transfected into TATi/Δku80 parasites using standard procedures, with successfully transfected parasites selected with chloramphenicol. TgFNT2 and TgFNT3 were localized by expressing C-terminally 3 × HA tagged forms of the proteins under the control of the constitutive α-tubulin promoter. The tgfnt2 open reading frame was amplified from T. gondii cDNA using primers p6 and p7 (Supplementary Table S1). The PCR product was digested with AvrII and BclI and ligated into the AvrII and BglII sites of the pUgCTH3 plasmid. The tgfnt3 open reading frame was amplified from cDNA using primers p8 and p9 (Supplementary Table S1). The PCR product was digested with AvrII and BglII and ligated into the AvrII and BglII sites of pUgCTH3. In both cases the vectors were transfected into TATi/tomato parasites.

**Generation and complementation of tgfnt knockout parasites.** We used a CRISPR/Cas9-based genome editing approach to generate parasite lines in which the open reading frames of tgfnt genes were disrupted. Using the Q5 Site-Directed Mutagenesis kit (New England Biolabs), we generated modified versions of the pSAG1::Cas9-U6::sgUPRT vector (Addgene plasmid # 5446735) to express guide RNAs targeting each tgfnt gene using a Q5 mutagenesis approach, as described by the manufacturer (New England Biolabs). The primers used for the Q5 mutagenesis reactions were p10 and p11 for tgfnt1, p12 and p11 for tgfnt2, and p13 and p11 for tgfnt3 (Supplementary Table S1). The resulting vectors co-express the guide RNA targeting the appropriate tgfnt gene and Cas9-GFP. They were transfected into TATi/tomato parasites, generating Δtgfnt1, Δtgfnt2, and Δtgfnt3 parasites (Supplementary Fig. S4). Parasites in which two and three tgfnt genes were disrupted were then made by targeting the tgfnt1 gene in Δtgfnt3 parasites then the tgfnt2 gene in Δtgfnt1/3 parasites (Supplementary Fig. S4).

For complementation studies, Δtgfnt1 parasites were transfected with vectors containing the tgfnt1, tgfnt2 or tgfnt3 gene fused at the 3’ end to a 3 × HA tag, under the control of the constitutive α-tubulin promoter. For TgFNT2 and TgFNT3, the vectors used were those described above for the localization of the ectopically expressed proteins. The complementation vector for TgFNT1 was made by amplifying the tgfnt1 open reading frame from genomic DNA using primers p14 and p5 (Supplementary Table S1). The PCR product was digested with AvrII and BglII and ligated into the AvrII and BglII sites of pUgCTH3.

**Immunofluorescence assays and western blotting.** Immunofluorescence assays (IFAs) and western blots were carried out as described previously. The primary antibodies used were rat anti-HA (Roche clone 3F10; 1:100 – 1:1000 dilution; used for IFAs and western blots) and mouse anti-P30 (Abcam clone TP3; 1:500 dilution; used for western blots). The secondary antibodies used for IFAs (all from Life Technologies) were goat anti-rat AlexaFluor 488 (1:100 – 1:250 dilution), goat anti-mouse AlexaFluor 546 (1:500 dilution), and goat anti-mouse AlexaFluor 647 (1:500 dilution). The microscopy was performed as described previously. For western blots the secondary antibody was horse radish peroxidase-conjugated goat anti-rat (Santa Cruz Biotechnology; 1:5000 dilution).

**Parasite growth assays.** Parasite growth was measured in the culture medium described above, which contains 25 mM glucose. Plaque assays were performed essentially as described previously, with 400 parasites added per 25 cm² flask. Fluorescence growth assays were performed in 96-well plates, with 4000 parasites added to each well and the fluorescence from tdTomato (excitation: 540 nm; emission: 590 nm) used to monitor parasite growth. Prior to estimating parasite growth rates, a value corresponding to the background fluorescence (averaged from wells to which parasites had not been added) was subtracted from all other data. The fluorescence intensity corresponding to maximum growth was then determined for each parasite line by fitting a sigmoidal curve to the data: \[ t = \frac{a}{1 + (t/t_{1/2})^b} \], where \( t \) is fluorescence intensity, \( a \) is the maximum fluorescence intensity, \( t \) is time and \( t_{1/2} \) is the time at which half-maximal growth is reached. The data for each parasite line were then expressed as a % of the maximum growth for that line. For each parasite line, the maximum growth rate was determined from the slope of the sigmoidal curve in a 4 h window surrounding the \( t_{1/2} \) (estimated by fitting the sigmoidal curve described above).

**pH measurements in extracellular tachyzoites.** Parasites were harvested by passage of cultures through a 26G needle. The parasites were then passed through a 3 μm filter to remove host cell debris and centrifuged for 20 min at 15000 × g. The supernatant media was removed and the parasites were suspended in ‘BCF’ (bicarbonate-free RPMI supplemented with 25 mM HEPES, 11 mM additional glucose, 0.2 mM hypoxanthine and pH-adjusted to 7.1) then centrifuged at 12,000 × g for 1 min. The supernatant media was removed and the parasites suspended again in BCF, to which the acetoxyethyl ester (AM) form of the pH-sensitive dye BCECF (Molecular Probes) was added at a final concentration of 5 μM. The parasites were incubated in the BCECF-containing medium for 14 min before being centrifuged (12,000 × g, 1 min), resuspended in BCF and maintained at 37 °C. Directly before their use in fluorescence measurements, parasites were centrifuged (12,000 × g, 1 min) and resuspended in pH 7.1 Saline Solution (125 mM NaCl, 5 mM KCl, 25 mM HEPES, 20 mM glucose and 1 mM MgCl₂). Fluorescence measurements were performed at 4 °C in a Cary Eclipse Fluorescence Spectrophotometer.
(Agilent). The excitation wavelengths were 440 nm and 495 nm and the emission wavelength was 520 nm. The ratio of fluorescence (495 nm/440 nm) was used as an indicator of cytosolic pH. The relationship between Fluorescence Ratio and pH was determined by suspending parasites in calibration salines of known pH containing 16 µM nigericin (to equilibrate H+ across the plasma membrane) and measuring the Fluorescence Ratio after it stabilized.

Measurements of the uptake of radiolabelled compounds by parasites. The uptake of radiolabelled compounds was measured in extracellular tachyzoites suspended in pH 6.1 Saline Solution (for l-[14C] lactate and l-[14C]arginine experiments) or Glucose-free Saline Solution (for [14C]2-DOG experiments; 125 mM NaCl, 5 mM KCl, 25 mM HEPES, 1 mM MgCl2; pH 6.1). The reactions for l-lactate experiments contained 200 µM unlabelled l-lactate and 0.2 µCi/ml l-[14C]lactate, those for the 2-DOG experiments contained 25 µM unlabelled 2-DOG and 0.2 µCi/ml [14C]2-DOG, and those for the L-arginine experiments contained 50 µM unlabelled L-arginine and 0.1 µCi/ml l-[14C]arginine.

The l-[14C]lactate and [14C]2-DOG experiments were conducted on ice, with centrifugation steps performed at 4 °C, to slow the rate of uptake such that initial rates could be determined. For the L-[14C]arginine experiments, parasites were incubated on ice for 30 min prior to commencing the experiments to mimic exposure conditions for the l-[14C]lactate experiments, but the experiments themselves were performed at 37 °C.

Uptake assays were performed as described previously. To commence an experiment, a volume of parasite suspension was added to an equal volume of a saline solution containing the radiolabelled compound. The uptake of radiolabelled compounds was stopped at pre-determined time points (in duplicate 200 µL samples) by centrifuging the samples in tubes containing 250 µL oil mix (84% Clerco PM-125 silicone fluid, 16% light mineral oil), thereby sedimenting the parasites below the oil layer. Aliquots (10 µL) of the supernatant solution were taken from above the oil layer to enable the extracellular concentrations of the radiolabelled compounds to be determined. For each tube, the remaining supernatant solution was removed, the tubes washed twice in running water, the oil aspirated, and the parasite pellet lysed with 500 µL 0.1% v/v Triton X-100. The lysate was transferred into a scintillation vial containing 1.5 mL scintillation fluid (Ultima Gold; Perkin Elmer), and the radioactivity measured using a scintillation counter.

To account for the radioactivity present in the extracellular fluid trapped around the cell pellet under the oil layer, we subtracted the amount of radioactivity present at time zero from the remaining data. Where there was significant uptake of radioactivity early in the time course (l-[14C]lactate and l-[14C]arginine experiments), we fit a curve to the time course data for each experiment \( y = y_0 + a(1 - e^{-bx}) \) to determine the amount of radioactivity present at time zero \( y_0 \). For [14C]2-DOG experiments the radioactivity measured at the first time point (taken within 30 s of exposure of parasites to [14C]2-DOG) was subtracted directly. A curve \( y = a(1 - e^{-bx}) \) was fitted to the data for intracellular radioactivity versus time and the initial rates were determined from the first derivative.

Experiments with Xenopus laevis oocytes. The frogs were purchased from Nasco (Cat# LM00535M) and were housed in the Xenopus Frog Facility of the ANU Research School of Biology in compliance with the relevant institutional and Australian Government regulations.

The oocyte expression vector containing pffnt was made previously. The oocyte expression vector containing tgfnt1 was made by amplifying the tgfnt1 open reading frame from gDNA using primers p14 and p5 (Supplementary Table S1). The PCR product was digested with AvrII and XmnI and ligated into the AvrII and XmnI sites of pGHJ-HA.

Oocytes were harvested from X. laevis frogs and prepared as outlined previously. cRNA encoding PfFNT and TgFNT1-3 HA was made using the mMessage mMachine T7 transcription kit and the MEGAclean kit (Ambion) and 30 ng was microinjected into oocytes, as described elsewhere. The uptake of l-[14C]lactic acid (Na‘ salt; 150.6 mCi/mmol; Perkin Elmer) was measured 3–4 days post-injection at 25.7 °C in ND96 buffer (96 mM NaCl, 2 mM KCl, 1 mM MgCl2, 1.8 mM CaCl2, 10 mM MES and 10 mM Tris-base; pH 6.4). Within each experiment measurements were made from 7 to 10 oocytes per condition. The influx of l-[14C]lactate was terminated by removing the reaction buffer and washing the oocytes twice in 3.5 mL of ice-cold ND96 buffer. The oocytes were then lysed and the radioactivity measured as described elsewhere.

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Author contributions

J.M.Z., K.K., G.G.v.D. and A.M.L. designed the study. J.M.Z., S.V.H. and S.H.S. performed experiments. J.M.Z., S.V.H., S.H.S., R.E.M., G.G.v.D. and A.M.L. analysed data. R.E.M., K.K., G.G.v.D. and A.M.L. supervised the study and provided resources. J.M.Z., G.G.v.D. and A.M.L. wrote the manuscript. All authors reviewed the manuscript.

Competing interests

The authors declare no competing interests.
