A conserved coccidian gene is involved in *Toxoplasma* sensitivity to the anti-apicomplexan compound, tartrolon E

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**ABSTRACT**

New treatments for the diseases caused by apicomplexans are needed. Recently, we determined that tartrolon E (trtE), a secondary metabolite derived from a shipworm symbiotic bacterium, has broad-spectrum anti-apicomplexan parasite activity. TrtE inhibits apicomplexans at nM concentrations in vitro, including *Cryptosporidium parvum*, *Toxoplasma gondii*, *Sarcocystis neurona*, *Plasmodium falciparum*, *Babesia* spp. and *Theileria equi*. To investigate the mechanism of action of trtE against apicomplexan parasites, we examined changes in the transcriptome of trtE-treated *T. gondii* parasites. RNA-Seq data revealed that the gene, *TGGT1_272370*, which is broadly conserved in the coccidia, is significantly upregulated within 4 h of treatment. Using bioinformatics and proteome data available on ToxoDB, we determined that the protein product of this tartrolon E responsive gene (*trg*) has multiple transmembrane domains, a phosphorylation site, and localizes to the plasma membrane. Deletion of *trg* in a luciferase-expressing *T. gondii* strain by CRISPR/Cas9 resulted in a 68% increase in parasite resistance to trtE treatment, supporting a role for the *trg* protein product in the response of *T. gondii* to trtE treatment. Trg is conserved in the coccidia, but not in more distantly related apicomplexans, indicating that this response to trtE may be unique to the coccidians, and other mechanisms may be operating in other trtE-sensitive apicomplexans. Uncovering the mechanisms by which trtE inhibits apicomplexans may identify shared pathways critical to apicomplexan parasite survival and advance the search for new treatments.

**1. Introduction**

Apicomplexan parasites are the cause of significant diseases of both humans and domesticated animals. Despite the current therapies available to treat these parasitic infections, the challenge of overcoming the rapid development of parasite drug resistance underscores the need for new treatments.

Recently we described the broad-spectrum anti-apicomplexan activity of tartrolon E (trtE), a secondary metabolite of symbiotic bacteria of shipworms (O’Connor et al., 2020). Tartrolons are a class of macrolide dimers, or pseudodimers consisting of polyketide chains joined as diesters that commonly bind to boron. These compounds are produced by both marine and terrestrial bacteria and exhibit antibacterial (Elshahawi et al., 2013; Isrchik et al., 1995; Schummer et al., 1994) and insecticidal activity (Lewer et al., 2003). We demonstrated that trtE inhibits many different apicomplexans at low nM concentrations in vitro, including *Cryptosporidium parvum*, *Toxoplasma gondii*, *Sarcocystis neurona*, *Plasmodium falciparum*, *Babesia* spp. and *Theileria equi*. In vivo efficacy of trtE...
has also been demonstrated, as the treatment of *C. parvum*-infected neonatal mice with trtE significantly reduced intestinal infection (O’Connor et al., 2020).

No other compound with this broad activity against apicomplexans has been reported, which makes the identification of the mechanism of action of trtE particularly important. To gain insights into the compound's anti-parasitic mechanism, we conducted an RNA-Seq analysis of trtE treated *T. gondii*. This analysis identified a conserved gene of unknown function that was upregulated during trtE treatment. When this gene was disrupted, *T. gondii* parasites became significantly more resistant to trtE, supporting a role for this gene product in the parasite’s response to trtE treatment. We name this gene trg for trtE responsive gene, and present bioinformatic analyses characterizing the gene and protein in *T. gondii* and related species.

2. Methods

2.1. Parasites

*T. gondii* strains ME49 and RH were maintained by serial passage in human foreskin fibroblasts (HFF, ATCC SCRC-1041), cultured in DME supplemented with 15% fetal bovine serum as previously described (Roos et al., 1995).

2.2. RNA-seq of trtE-treated *T. gondii* RH parasites

Total RNA from *T. gondii* RH parasites treated with 24.2 μM trtE or 0.1% DMSO control was collected 4, 8, and 12 h post-treatment for RNA-Seq analysis. The details of the experimental analysis and results of the transcriptome are available in the GEO database (GSE140197). Briefly, the integrity and quantity of the total RNA samples were determined using a Fragment Analyzer (ThermoFisher Scientific, Waltham, MA, USA). The RNA samples that satisfied the input requirement were used as input for the Illumina TruSeq Stranded Total RNA with Ribo-Zero supplemented with 15% fetal bovine serum as previously described.

The integrity and quantity of the total RNA samples were determined using the Fragment Analyzer. The libraries were pooled and sequenced using HiSeq 2500 High Output V4 chemistry (Illumina). The 75 million to 95 million reads per sample obtained were analyzed using the TopHat-Cufflinks pipeline. The expression level was estimated with Cuffdiff and represented as fragments per kilobase of exons per million mapped fragments (FPKM) to the *T. gondii* GT1 genome available on ToxoDB (Gajria et al., 2008) (http://toxodb.org). Genes were considered differentially expressed when both the p-value and q-value, estimated using the Benjamini-Hochberg procedure, were below a significance value of 0.05. The RNA-Seq experiment was conducted once.

2.3. Reverse transcriptase quantitative PCR

To confirm RNA-Seq results, HFF cells were infected with 2 x 10⁶ *T. gondii* RH parasites per well in six-well plates. After an additional 24 h of incubation, infected cells were treated with 24.2 nM trtE, 10 μM pyrimethamine, or 0.1% DMSO carrier control in triplicate. Total RNA was extracted from treated, infected cells using an RNeasy kit (Qiagen, Germantown, MD, USA) at 1, 2, and 4 h post treatment. Synthesis of cDNA for each RNA sample was accomplished using iScript™ cDNA Synthesis Kit (Bio-Rad, Hercules, CA, USA) using 1 μg of RNA template. Quantitative PCR was performed on the CFX96 Touch™ Real-Time PCR Detection System (Bio-Rad) using SsoAdvanced™ Universal SYBR® Green Supermix (Bio-Rad). The amplification conditions were: 98 °C for 30 s, 40 cycles of denaturation at 98 °C for 30 s, and annealing/extension at 60 °C for 30 s. A melting curve, representing a 1 s hold at every 0.5 °C between 65° and 95 °C, was generated to confirm a single peak for each primer pair. Primer sequences and product sizes for each pair are provided in Table S1. Relative expression of TGGT1_272370 (hereafter referred to as trg, GenBank accession no: XM_018781656) and bradyzoite antigen 1 (bag1, GenBank accession no: XM_002365075) was determined using the 2-ΔΔCt method (Livak and Schmittgen, 2001) using expression of the endogenous parasite actin gene (act J; GenBank accession no: XM_002369622) for normalization.

To investigate the dose-dependent expression of trg during trtE treatment, HFF cells were infected with 2 x 10⁶ *T. gondii* RH parasites in six-well plates. Twenty-four hours post-infection, infected cells were treated with 60.5, 24.2, 12.1, 6.1, 1.2 nM trtE or 0.1% DMSO control for 4 h before RNA extraction. Three technical replicates were performed on each sample. Data analysis was accomplished using GraphPad Prism 8.0 Software (GraphPad Software, San Diego, CA, USA).

2.4. Bioinformatics analysis of trg predicted protein product

The complete amino acid sequence of the ME49 strain Trg protein (GenBank accession no: XP_0186366692) was retrieved from the National Center for Biotechnology Information (NCBI) database. The presence of multiple post-translational modification sites were predicted including phosphorylation sites using NetPhos 3.1 server (Blom et al., 1999) (https://services.healthtech.dtu.dk/service.php?NetPhos-3.1) with kinase-specific predictions (Blom et al., 2004), and palmitylation sites using CSS-Palm 4.0 (Ben et al., 2008) (http://cspalm.biocuckoo.org/online.php). The TMHMM server v2.0 (Krogh et al., 2001) (https://services.healthtech.dtu.dk/service.php?TMHMM-2.0) was used to estimate transmembrane domains. Both WoLF PSORT (Horton et al., 2007) (https://wolfpsort.hgc.jp) using “Animal” as the organism type) and data from hyperplexed Localization of Organellar Proteins by Isotopic Tagging (hyperLOPIT) experiments using extracelluar *T. gondii* tachyzoites (https://proteome.shinyapps.io/toxolopittzex/) accessed through ToxoDB (Gajria et al., 2008) (http://toxodb.org) were used to predict Trg localization. Identification of short linear motifs in the Trg sequence was accomplished using the eukaryotic linear motif (ELM) resource (Gouw et al., 2018) (http://elm.eu.org) filtering for predicted cell compartment and *T. gondii* taxonomic context. Protein sequences of Trg homologs in other coccidia were identified by searching NCBI Database resources (NCBI Resource Coordinators, 2016) using the basic local alignment search tool (Altschul et al., 1990), ToxoDB (Gajria et al., 2008) (http://toxodb.org), and Ensembl Genomes (Hove et al., 2019) (http://www.ensemblgenomes.org) aligned using the Clustal W (Thompson et al., 1994) algorithm of the AlignX module of Vector NTI (https://wwwVECTORNTI.com) and filtered against the Eukaryotic Pathogen CRISPR guide RNA/DNA Design Tool (Tarleton and Peng, 2015) (http://grna.ctegd.uga.edu) using the identified guide sequence targeting *T. gondii* act 1 (GenBank accession no: XM_002369622) for normalization.

2.5. CRISPR/Cas9-mediated deletion of trg

A clustered regularly interspaced short palindromic repeats (CRISPR)/Cas9-associated protein 9 method (CRISPR/Cas9) was designed to knockout trg in the *T. gondii* ME49 ΔHPT:Luc strain (Tobin and Knoll, 2012). A guide sequence (5’-AGTTGATGGCGAGGACCTGCTG-3’) targeting trg was identified using the Eukaryotic Pathogen CRISPR guide RNA/DNA Design Tool (Tarleton and Peng, 2015) (http://gRNA.cTED.UGA.EDU) based on the efficiency score against the *T. gondii* ME49 genome available on ToxoDB (Gajria et al., 2008) (http://toxodb.org). The pSAG1: CAS9-U6:sg272370 (Fig. 1A) was prepared by replacing the single guide (sgRNA) sequence of the pSAG1: CAS9-U6 plasmid (Shen et al., 2014) with the identified guide sequence targeting trg using primers listed in Table S1 and the Q5® Site-Directed Mutagenesis Kit (New England Biolabs, Ipswich, MA, USA). A second plasmid provided the regions of sequence identity for homologous recombination and the mCherry gene substitution for trg. This plasmid, pBC-GFP-272370-mCherry (Fig. 1B, designated the “flanking plasmid”), was generated by Gibson assembly® (New England Biolabs) of PCR
The plasmid containing a mCherry expression cassette (red, pSAG1-mCherry-DHFR 3’ UTR) amplified from the pBC-GFP-mCherry plasmid and the linearized pBC-GFP-mCherry plasmid (containing a GFP expression cassette, pTUB1-eGFP-SAG1 3’ UTR). Plasmids were produced in transfected E. coli D5α (New England Biolabs) and isolated using the QIAGEN Plasmid Maxi Kit (QIAGEN). T. gondii ME49 ΔHPT:tachyzoites were transfected with the pSAG1:Cas9-U6:sg272370 plasmid (pink) targeted to trg and induced a double-stranded break. Linearized pBC-mCherry-272370-GFP plasmid provided a template for double homologous recombination, which is designed to replace the entire gene with mCherry and lose GFP for sorting. (For interpretation of the references to colour in this figure legend, the reader is referred to the Web version of this article.)

2.7. PCR test for trg

A total of 25 ng of gDNA from T. gondii ME49 ΔHPT:LUC parental strain and each of the eight ME49 ΔHPT:LUC Δtrg clones, was mixed with primers designed to amplify a 172 bp fragment of trg (Table S1) and Q5® Hot Start High-Fidelity 2X PCR Master Mix (New England Biolabs). The amplification conditions were: 98°C for 30 s, 30 cycles of denaturation at 98°C for 5 s, annealing at 68°C, extension at 72°C for 20 s, and a final extension at 72°C for 2 min. Reactions were resolved on a 2% agarose gel using a 100 bp DNA ladder (New England Biolabs) for size estimation.

2.8. Southern blot analysis

Approximately 5 µg of gDNA from each strain was digested overnight with the restriction endonuclease Bsu36I (New England Biolabs) and resolved on a 0.8% agarose gel using a DIG-labeled DNA Molecular Weight Marker II (MilliporeSigma, St. Louis, MO, USA) for size estimation. After blotting and u. v. crosslinking to a nylon membrane (BrightStar™ Plus Positively Charged Nylon Membrane, Thermo Fisher Scientific), the membrane was hybridized to digoxigenin (DIG)-labeled mCherry gene probes generated by PCR using the PCR DIG Probe Synthesis Kit (MilliporeSigma), the primers shown in Table S1, and pBG-GFP-mCherry plasmid template. Hybridizations were performed overnight at 47°C using DIG Easy Hyb™ hybridization buffer (Milli- poreSigma), and final washes were made in 0.1X saline sodium citrate (SSC)/0.1% sodium dodecyl sulfate (SDS) solution for 15 min at 68°C. Chemiluminescence of probe-targeted hybrids was detected by anti-DIG-alkaline phosphatase antibody (MilliporeSigma) and visualized using the ChemiDoc Touch™ Imaging System (Bio-Rad).

2.9. In vitro inhibitor response assays

HFF cells in 96-well plates were infected with 5 × 10⁴ parental or trg knockout clones for 24 h before treatment with eight 2-fold dilutions of trtE (60.5–0.5 nM) or DMSO control with three technical replicates for each concentration. Treated parasites were incubated at 37°C and 5% CO₂ for 24 h before measuring luciferase (LUC) activity using the Bright-


Glo™ Luciferase Assay System (Promega, Madison, WI, USA). Inhibition was calculated by comparing the LUC activity of trtE-treated and untreated parasites. Estimation of the EC$_{50}$ value for trtE treatment using data from three independent experiments was accomplished using a four-parameter logistic regression, and comparisons between estimated EC$_{50}$ values of the strains were made using an extra sum-of-squares F test using GraphPad Prism 8.00 Software (GraphPad Software).

3. Results

3.1. T. gondii responds to trtE treatment by upregulating expression of trg

To identify proteins that may be targeted by trtE, we conducted a preliminary RNA-Seq experiment to examine the transcriptome of trtE-treated T. gondii parasites. Of the 90 significantly and differentially expressed T. gondii genes over the 12-hr. treatment period (Fig. S1, Table S2), only trg (Fig. 2A) and TGGT1_311100 (Fig. S1) were upregulated within 4 h of treatment and remained upregulated throughout the entire treatment period.

To confirm RNA-Seq results, we treated T. gondii ME49 strain with trtE or DMSO for 1–4 h and measured the expression of trg and TGGT1_311100 homologs by RT-qPCR. Expression of trg but not TgME49.311100 (Fig. S2), was upregulated within 4 h of trtE-treatment (Fig. 2B). Treatment with the known T. gondii inhibitor pyrimethamine did not affect trg transcript abundance. The expression of bag1 was decreased after 4 h of trtE treatment (Fig. S2). Additionally, we observed a linear dose-dependent expression of trg 4 h after the treatment of T. gondii parasites with trtE (Fig. 2C).

3.2. Bioinformatic analyses of trg predicted protein sequence

The ME49 strain trg gene is 12,805 bp in length comprised of 9 exons, of which 7107 bp encode a protein of 2368 aa. To gain insight into the function of the protein, the presence of predicted post-translational modifications and functional motifs was investigated. NetPhos 3.1 (Blom et al., 1999) was used to predict 137 potential sites with a probability score higher than 75%. Of these, three positions (T393, S502, S530) were identified as protein kinase A (PKA) family-specific sites, and six positions (S134, S174, T443, S1076, T2172, T2300) were identified as protein kinase C (PKC) family-specific sites and six positions (S134, S174, T443, S1076, T2172, T2300) were identified as protein kinase C (PKC) family-specific sites; the rest were uncharacterized.

Predicted palmitoylation sites identified by CSS-Palm 4.0 (Ren et al., 2010) for Serine, 76.9% (105/137) threonine, 22.6% (31/137) tyrosine, and 2.2% (3/137) threonine. Predicted palmitoylation sites identified by CSS-Palm 4.0 (Ren et al., 2008) clustered around two areas (C635, C636, C641, C646; C1637, C1647, C1684) of Trg. While bioinformatics analyses predicted multiple phosphorylation and palmitoylation sites, only one residue (T2262) has experimental evidence of phosphorylation (Treek et al., 2011). Trg was not identified in the palmitoylated proteins discovered by DIA-MS (Foe et al., 2015) or ABE-MS (Caballero et al., 2016).

To examine the predicted localization of Trg, we used both computational prediction tools and reports of experimental proteome data of targeted proteins. TMHMM v2.0 (Krogh et al., 2001) predicted five transmembrane domains in the protein sequence (positions 610–632, 748–770, 1618–1640, 1653–1675, and 2192–2214). WolF PSORT (Horton et al., 2007) analysis of the protein sequence predicted the uncharacterized protein to be localized to the plasma membrane with 81.25% (26/32) nearest neighbors associated with the plasma membrane, 12.5% (4/32) neighbors associated with the nucleus, and 6.25% (2/32) neighbors associated with the cytoplasm. The hyperLOPIT proteomic data for T. gondii tachyzoites accessed through ToxoDB (Gajria et al., 2008) also localized Trg to the plasma membrane.

Because trtE is broadly effective against multiple apicomplexan parasites, we looked for proteins that shared sequence identity with Trg among the apicomplexa. However, after searching various databases (NCBI, ToxoDB.org, and Ensembl Genomes), we identified homologs with similar predicted transmembrane structures in the coccidia only (Table 1, Fig. 3, and Fig. S3). No homologs of Trg were found in other more distantly related apicomplexans.

3.3. Deletion of trg results in an increase in parasite resistance to trtE-treatment

To determine if Trg played a direct role in the response of T. gondii...
parasites to treatment with trtE, we generated trg deletion mutants using CRISPR/CAS9. After the selection of putative knockout parasites by FACS, and two rounds of cloning by limiting dilution, the presence of trg was assessed by PCR in eight ME49 ΔHPT:LUC Δtrg isolates (Fig. 4A); and screened for erroneous insertion events by Southern blot analysis (Fig. 4B). We were unable to detect trg in ME49ΔHPT:LUC Δtrg clones by PCR (Fig. 4A), indicating disruption of the gene. No erroneous insertion events of the mCherry expression cassette into the chromosome of ME49 ΔHPT:LUC Δtrg clones were detected by Southern blot (Fig. 4B).

There were no apparent differences in growth rates between wt and the ME49 ΔHPT:LUC Δtrg mutants, as was previously observed (Sidik et al., 2016).

We compared the susceptibility of the parental strain and three of the eight ME49 ΔHPT:LUC Δtrg clones (1, 3, and 4) to trtE-treatment by quantifying the EC50 of the compound for each clone. ME49 ΔHPT:LUC Δtrg parasites were, on average 68% more resistant to trtE treatment than the ME49 ΔHPT:LUC parental line (Fig. 4C) with an estimated EC50 of 5.088 ng/ml (4.144–6.247 ng/ml; 95% CI) for the 3 deletion mutants clones as compared to an EC50 of 3.027 ng/ml (2.309–3.968 ng/ml; 95% CI) for the parental line.

### Table 1

| Organism          | GeneID  | Accession No. | Identity (%) | Similarity (%) | Coverage (%) |
|-------------------|---------|---------------|--------------|----------------|--------------|
| H. hammondi       | HHA_272370 | KEP66945.1   | 73.31 [1736] | 77.53 [100]    | 84           |
| B. besnoiti Ger 1 | BESB_032870 | XP_029215099.1 | 35.43 [958]  | 47.08 [315]    | 30           |
| N. caninum        | NCLIV1034940 | CEL67703.1  | 32.07 [813]  | 42.96 [276]    | 48           |
| C. suis Wien I    | CU1_002351 | PHI23796.1  | 20.33 [615]  | 31.93 [351]    | 23           |
| C. cayetanensis NFI_C8 | LOC34621828 | XP_026189647 | 17.09 [440]  | 25.27 [212]    | 11           |
| E. tenella Houghan | ETH_00032795 | CDJ37990.1 | 16.08 [411]  | 23.79 [197]    | 12           |
| S. neurona SN3    | SN3_00202415 | NA            | 10.19 [269]  | 18.55 [221]    | 14           |

* Not available in the NCBI database.
4. Discussion

This study has identified a component of the response of T. gondii parasites to the pan-anti-apicomplexan compound, trtE. Because trtE-treated T. gondii parasites die within 1 h of treatment (O’Connor et al., 2020), genes that were differentially expressed after 4 h of treatment were targeted in this study. RNA-Seq of trtE-treated T. gondii parasites identified five genes in addition to trg (TGGT1_311100, TGGT1_233460, TGGT1_214080, TGGT1_311425, and TGGT1_310740) that were differentially expressed after 4 h of treatment (Fig. S1 and Table S2). Two of these genes have characterized protein products (GenElD:protein; TgME49_214080:toxofilin, and TgME49_233460:RSRS98:SAg1). Toxofilin is an actin-binding protein (Poupel et al., 2000) that has been demonstrated to disassemble host actin and facilitate rapid parasite invasion of host cells (Delorme-Walker et al., 2012). The parasite surface protein SAG1 has been shown to play an active role in attachment and invasion of T. gondii tachyzoites (Mineo and Kasper, 1994) and anti-SAG1 antibodies partially block invasion (Mineo et al., 1993). The expression of both genes was downregulated 4 h post trtE-treatment suggesting trtE may affect tachyzoite invasion into host cells. Pretreatment of tachyzoites with trtE prevents the parasite from establishing infection, but it was not determined if growth inhibition was pre or post-invasion (O’Connor et al., 2020). Of the remaining genes differentially expressed within 4 h of treatment, only trg and TGGT1_311100 were significantly upregulated throughout the entire RNA-Seq experiment; these genes were selected for further study.

Using qRT-PCR, we confirmed that the trg transcript (but not TGGT1_311100) was significantly upregulated in a rapid, dose-dependent manner in response to trtE treatment. This observation appears to be specific to trtE-treatment and not related to a general inhibition of the parasites since treatment with the anti-Toxoplasma compound pyrviniummephenamine did not induce an increase in trg transcripts. Stress can cause T. gondii parasites to form bradyzoites in vitro (Mayoral et al., 2020). However, trtE treatment reduced expression of the bradyzoite-specific gene bag1 after 4 h of treatment, suggesting that the increase in expression of trg is not related to stress-associated bradyzoite formation. This observation is corroborated by studies showing that trg expression is not upregulated during bradyzoite formation (Behnke et al., 2008). In general, it appears that trg is not stage-specific as transcripts are detected during the merozoite (Behnke et al., 2014), oocyst, tachyzoite, and bradyzoite stages (Fritz et al., 2012). In a CRISPR genome-wide loss of function screen (Sidik et al., 2016), trg was found to be dispensable and, consistent with this report, we did not observe any obvious differences in growth rate between the parental and the trg knock out clones.

Bioinformatic analysis of the Trg sequence did not provide definitive clues as to the function of the protein. Trg has multiple transmembrane domains; it likely localizes to the plasma membrane and is a substrate for phosphorylation. The predictions for localization and endosomal targeting, common to membrane-bound proteins, are supported by both mass spectrometry data that identified the protein in the membrane fraction of T. gondii RH parasites (Dybas et al., 2008), and hyperLOPIT proteome data of extracellular tachyzoites (https://proteome.shinyapps.io/toxolopittzex/). There is experimental evidence that one of the predicted threonine phosphorylation sites (T2262) of Trg is phosphorylated (Treeck et al., 2011). Altogether these observations suggest the protein may play a role in a signaling cascade.

The trg gene is conserved amongst coccidia, and homologs were identified in Besnoitia besnoiti, Cyclospora cayetanensis, Cytospora suis, Eimeria tenella, Hammondia hammondi, N. caninum, and S. neurona (Table 1, Fig S3). Of the coccidia, we have confirmed the activity of trtE against S. neurona (O’Connor et al., 2020). Homologs to Trg in other trtE-sensitive apicomplexans may have sufficiently diverged such that we could not identify them by protein sequence identity, especially as there are no identified functional motifs in Trg. Alternatively, this response to trtE may be unique to the coccidians, and other mechanisms may be operating in Plasmodium, the pirolamops, and Cryptosporidium.

We anticipate that trtE has potassium ionophore activity since it is structurally similar to the known potassium ionophores trb (Surup et al., 2018) and boromycin (Moreira et al., 2016; Pache and Zahner, 1969). Ionophores are a promising source of treatments for drug-resistant parasites (Kevin Li et al., 2009). For example, monensin, a sodium ionophore isolated from the bacterium Streptomyces cinnamomi, has activity against Plasmodium spp., C. parvum, and T. gondii (Couzinet et al., 2000; Gulmia et al., 1997; McDonald et al., 1990). Using a forward genetic screen to identify mutant T. gondii parasites resistant monensin, Garrison and Arrizabalaga found that the disruption of a mitochondrial DNA repair enzyme (TgMSH-1) contributed to parasite resistance to both monensin and the potassium ionophore salinomycin (Garrison and Arrizabalaga, 2009). A study of the transcriptomic response of T. gondii parasites to monensin and salinomycin identified significant upregulation of histones and other genes associated with cell cycle arrest (Lavine and Arrizabalaga, 2011). However, the transcriptomic profile of trtE-treated T. gondii (Table S2 and Fig S1) was not in any way similar to that of monensin/salinomycin-treated parasites, suggesting that trtE has a different mechanism of action from these ionophores.

Using a combination of transcriptomic and genomic strategies, we identified a conserved coccidian gene that plays a role in the susceptibility of T. gondii parasites to the anti-apicomplexan compound, trtE. The function of the trg gene product remains entirely unclear, as does its relevance to the mechanism of action of trtE against non-coccidian apicomplexans. Further study into the function of Trg may clarify the specific mechanism of action of trtE against coccidans and identify a new drug target common to coccidian parasites of human and animal importance.

Declaration of competing interest

Please declare any financial or personal interests that might be potentially viewed to influence the work presented. Interests could include consultancies, honoraria, patent ownership or other. If there are none state ‘there are none’.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.ijpddr.2020.07.003.

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