Double drugging of prolyl-tRNA synthetase provides a new paradigm for anti-infective drug development

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Abstract

Toxoplasmosis is caused by Toxoplasma gondii and in immunocompromised patients it may lead to seizures, encephalitis or death. The conserved enzyme prolyl-tRNA synthetase (PRS) is a validated druggable target in Toxoplasma gondii but the traditional ‘single target–single drug’ approach has its caveats. Here, we describe two potent inhibitors namely halo-fuginone (HFG) and a novel ATP mimetic (L95) that bind to Toxoplasma gondii PRS simultaneously at different neighbouring sites to cover all three of the enzyme substrate subsites. HFG and L95 act as one triple-site inhibitor in tandem and form an unusual ternary complex wherein HFG occupies the 3’-end of tRNA and the L-proline (L-pro) binding sites while L95 occupies the ATP pocket. These inhibitors exhibit nanomolar IC\textsubscript{50} and EC\textsubscript{50} values independently, and when given together reveal an additive mode of action in parasite inhibition assays. This work validates a novel approach and lays a structural framework for further drug development based on simultaneous targeting of multiple pockets to inhibit druggable proteins.

Author summary

Among infectious diseases, parasitic diseases are a major cause of death and morbidity. Toxoplasmosis is caused by an infection of the apicomplexan parasite Toxoplasma gondii. In immunocompromised patients Toxoplasmosis may lead to seizures, encephalitis or death. Novel therapeutics for human parasites are constantly needed. In recent years, the aminoacyl-tRNA synthetase (aaRS) enzyme family has been validated as a drug target for several parasitic infections. The Toxoplasma gondii prolyl-tRNA synthetase inhibitor...
halofuginone (HFG) has been validated earlier but here we show that an ATP-mimic called L95 is a potent inhibitor and can bind to the target enzyme in the presence of HFG. Thus, the two inhibitors described in this study simultaneously occupy all three natural substrate (ATP, L-amino acid and 3’-end of tRNA) binding pockets and thereby inhibit the enzyme leading to parasite death. This unprecedented double drugging of a pathogen enzyme may delay resistance mutation generation and this approach opens the path to multi-drugging of validated parasite proteins.

Introduction

Management of infectious diseases continues to be a huge priority for the global medical community as they comprise nearly half of the major global health concerns [1]. Amongst them, apicomplexan parasites like Plasmodia and Toxoplasma that cause malaria and toxoplasmosis are causes of morbidity and mortality worldwide [2,3]. Toxoplasma gondii (Tg) is an obligate intracellular parasitic protozoan which causes toxoplasmosis. It is one of the most common parasites in developed countries with a serological infection rate of up to 30–50% [4]. In humans, active toxoplasmosis can be treated with a combination of drugs such as pyrimethamine and sulfadiazine along with folinic acid [5]. Considering the likely development of drug resistance against available therapeutics, there is a continued need for identifying novel targeting strategies against this parasitic infection.

Aminoacyl-tRNA synthetases (aaRSs) are highly conserved and essential enzymes responsible for feeding charged tRNA molecules into the protein synthesis machinery [6,7]. The active site of aaRSs has three closely located sub-sites for natural substrates L-amino acid (aa), ATP and 3’-end tRNA [6]. Inhibiting these enzymes stalls protein translation. By virtue of this, several aaRSs have been recognized as druggable targets for infectious and human diseases [8–11]. In recent years, inhibitors targeting their active and allosteric sites have been reported [12–15]. The active site inhibitors were further grouped based on their binding modes such as single (any one of the substrate sub-sites inhibited) [16–19], dual (overlapping of two substrate subsites [20–22] or substrate sub-sites with an auxiliary pocket [23,24]), triple or multi-site (covering all substrate subsites) [25], while the non-aminoacylation site inhibitors were grouped into editing, allosteric and non-translational function-site inhibitors [26–29]. Prolyl-tRNA synthetases (PRS) charge tRNA molecules with L-pro for protein translation. Over the last decade, parasite- and host-encoded PRSs have been extensively studied using febrifugine (FF) and its derivatives including halofuginone (HFG) that block both the L-pro and the 3’-end of the tRNA binding pockets [20–21,30–32]. Hewitt et al. in 2017 showed that glyburide and the ligand TCMDC124506 can target the PRS enzyme allosterically [27]. Recent advancements towards ATP mimetics targeting the ATP pocket of PRSs have opened a new window towards drug development [33–36]. A partner group recently published a series of inhibitors that targets the ATP pockets of both malaria and host PRSs [37].

The simultaneous targeting of distinct catalytic sites of a single target using a cooperative ligand binding approach has also been explored recently. Although structurally incomplete and poorly understood, these discoveries highlight the importance of multivalent targeting strategies in modulating the efficacy of druggable hits [38–42].

In this study, we examined a pyrrolidine-based ATP mimic–PubChem ID SCHEMBL22505739 (hereafter labelled as L95, Figs 1A and S1) and validated it as an ATP mimic inhibitor against Toxoplasma gondii PRS (TgPRS) enzyme. Moreover, we hypothesized that L95 in concert with HFG can bind to and inhibit TgPRS as the two ligands reside in
close proximity but different substrate pockets. We validated our hypothesis via biochemical, structural and cellular evidence. Here, we present high-resolution crystal structures of two ternary complexes; a single liganded \( \text{TgPRS-L-pro-L95} \) and concurrently a bound double liganded \( \text{TgPRS-HFG-L95} \). Our work serves as proof for simultaneous targeting of PRS enzyme with two ligands and this study lays a framework for new drug development that focuses on a novel approach to counter drug resistance in major infectious diseases like malaria and toxoplasmosis.

**Results**

**Two inhibitors HFG and L95 binding simultaneously to the target TgPRS enzyme**

The binding potencies of the inhibitors (L95 and HFG) in the presence or absence of substrates (L-pro and ATP) were evaluated against recombinant TgPRS enzyme using a fluorescence-based thermal shift assay (TSA). The ligand (inhibitors/substrates) binding usually helps to stabilize the enzyme due to enzyme-ligand interactions, resulting in an increase of the melting temperature (T\( _m \)) of the target protein during the thermal denaturation process [43]. The \( \Delta T_m \) was calculated based on the difference between the \( T_m \) values of TgPRS with and without ligands. The \( T_m \) of TgPRS-apo is 52.3°C. In the presence of L95 alone, the \( T_m \) shifts to a higher temperature by \( \Delta T_m \) of 17.8°C, whereas in the additional presence of L-pro, the thermal stability of TgPRS enzyme improves significantly resulting in \( \Delta T_m \) value of 20.7°C suggesting that the inhibitor L95 is binding to TgPRS in an L-pro-dependent manner (Figs 1B and S2). The higher \( \Delta T_m \) of L95 in the presence of L-pro implies that induced conformational changes may stabilize the ternary complex of TgPRS-L-pro-L95 and suggests a cooperative binding mode of L95 with L-pro. Similarly, the measured \( \Delta T_m \) for HFG alone and HFG with ATP were 8.1°C and 37.6°C respectively, which is similar to previously reported values with the ATP facilitating HFG integration within the active site of the enzyme [20,21,44,45].

Next, we validated that the inhibitor L95 occupies the ATP binding pocket of TgPRS more effectively in the presence of L-pro (Fig 1B, compare L95 with L95+L-pro). Further, we
hypothesized the two drugs HFG and L95 occupying the three pockets together: L-pro, 3'-end of tRNA and ATP within the active site. For this, both HFG and L95 were added to TgPRS simultaneously and the ternary complex of TgPRS-HFG-L95 showed a $\Delta T_m$ of 23.6˚C. (Fig 1B). The similar $\Delta T_m$ values of TgPRS-HFG-L95 and TgPRS-L-pro-L95 suggest that the association of TgPRS with L95 in the presence of L-pro is quite similar to its binding with L95 in presence of HFG. However, the slightly higher $\Delta T_m$ value for TgPRS-HFG-ATP indicates that ATP (natural substrate) provides more stability than L95.

### PRS enzyme inhibition by L95

The enzymatic activity of recombinant proteins TgPRS and HsPRS were evaluated in the presence of either HFG or L95 by performing aminoacylation assays. The first part of the aminoacylation reaction where the aminoacyl-adenylate complex is formed was studied in the presence of these inhibitors. Here, the aminoacyl-adenylate complex is formed with the release of pyrophosphate (PPi) which can be indirectly measured by a malachite green assay that uses the pyrophosphatase-induced conversion of PPi into inorganic phosphate (Pi). For the enzyme inhibition assay, concentration ranges of 50 to 0.005 $\mu$M were used for HFG and L95. The measured half-maximal inhibitory concentration ($IC_{50}$) values computed for TgPRS in the presence of individual inhibitors HFG and L95 were 5.41 nM and 47.7 nM respectively (Fig 1C). The measured $IC_{50}$ value for HFG against TgPRS is very similar to our previous reports [20]. For HsPRS, the $IC_{50}$ values computed for HFG and L95 were 10.65 nM and 47.7 nM respectively. The high potency of these inhibitors is evident from their nanomolar $IC_{50}$ values.

### Structural features of single (L95) and concurrently double liganded (HFG-L95) TgPRS enzyme

To understand the binding mechanism of the single and double liganded TgPRSs, we determined two co-crystal structures of TgPRS in complex with L95 and L-pro (TgPRS-L-pro-L95) (PDB id 7EVV) and another in complex with HFG and L95 (TgPRS-HFG-L95) (PDB id 7EVU) at a resolution of 1.7 Å (Figs 2 and 3 and Table 1). Difference Fourier electron density

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**Fig 2. Crystal Structure summary of HFG and L95 bound to TgPRS.** (a and b) Difference Fourier map (Fo-Fc) of the bound ligands in the aminoacylation site of TgPRS. The Fo-Fc maps are contoured at 3 $\sigma$ and shown as green mesh representations. Ligands L-pro (green), L95 (yellow), HFG (pink) are shown as sticks. (c) Superposition of TgPRS-L-pro-L95 and TgPRS-HFG-L95 complexes showing the ligand induced movement of loops. (d) The interactions between TgPRS and bound L-pro-L95. The salt bridge interaction with L-pro and interacting residues with L95 are shown as sticks. The hydrogen bonds are shown as black dashed lines.

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maps confirmed the presence of L95 in the ATP pocket (Fig 2A). There is one monomer per asymmetric unit in the C2 space group of the TgPRS-L-pro-L95 ternary complex, while the biologically active dimer has crystallised in the P1 space group for the concurrently double liganded TgPRS enzyme (Figs 2B and S3). Superimposition of two chains in the TgPRS-HFG-L95 complex shows a high degree of similarity as evidenced by the overall mean RMSD values of 0.28 Å for 480 Cα-atoms. Therefore, for all structural descriptions, here on we have only considered chain A.

Superposition of the ATP pocket of both L95 bound structures showed that L95 adopts a similar but not identical conformation with minute deviations in the phenylacetamide moiety (Fig 2C). The class II signature motifs 2 (residues 460–494) and 3 (residues 585–603) of TgPRS form multiple interactions with L95. The inhibitor L95 is stabilized mainly by three direct and three water-mediated hydrogen bonds. The three direct hydrogen bonds are: (1) 6-methylpyridine N1 atom with main-chain N atom of Thr482 (2) phenylacetamide N-atom with main-chain O of Thr482 and (3) oxo-pyrrolidine keto O atom with side-chain OG1 atom of Thr592 (Figs 2D and S4). The hydrogen-bonding interactions between the hinge region of L95 and the protein main chain is very similar to that observed in Tyk2 inhibition [46]. The 6-methylpyridine ring is sandwiched by Phe485 via π-π stacking interaction (~3.9 Å) and Arg594 via cation-π interaction (S5 Fig). Superposition of the ligand complexes HFG-L95 (PDB id 7EVU) and HFG-ANP (non-hydrolysable ATP analog) (PDB id 5XIQ) showed that the inhibitor HFG adopts a similar conformation with identical enzyme-HFG interactions except for the stacking interaction between the quinazoline-4-one moiety of HFG and Phe415 (Fig 3A and 3B).

Structural analysis of TgPRS-L-pro-L95 shows that the binding of L-pro induces the movement of the side chain of Arg470 which forms a salt bridge between the carboxylic group of L-pro and the guanidium group of Arg470 (S6 Fig). This consequentially provides ample space to accommodate the binding of L95 in the ATP pocket—providing a good rationale as to why L95 shows tight binding in the presence of L-pro as indicated by Tm measurements. Superimposition of TgPRS-HFG-L95 and TgPRS-HFG-ANP shows that the ATP binding loop (residues 470–483) is largely displaced and significant side-chain deviations were found for Glu472 and Arg481 (Fig 3B). This displacement is indicative of an accommodative effort for the
phenylacetamide moiety of L95 by the active site of the enzyme. Differential loop movements
induced by ligand binding (528–538, particularly Phe534–Ala535–Gly536 where in Phe534
adopts unique rotameric conformations) were observed within HFG-ANP bound, HFG-L95

Table 1. Summary of data collection and refinement statistics. Values in parentheses are for the highest resolution shell.

|                          | TgPRS-L-pro-L95 | TgPRS-HFG-L95 |
|--------------------------|-----------------|---------------|
| **PDB code**             | 7EVV            | 7EVU          |
| **Data collection**       |                 |               |
| Beamline                 | PROXIMA-1 (PX1) | I03, DLS      |
| Wavelength (Å)           | 0.979           | 0.976         |
| Detector type            | EIGER X 16M     | Eiger2 XE 16M |
| Crystal-to-detector distance (mm) | 270.6        |
| Oscillation (°)          | 0.1             | 0.1           |
| Exposure (s)             | 0.5             | 0.02          |
| No. of images            | 3600            | 3600          |
| Software used for data processing | XDS           | XIA2/DIALS   |
| **Space group**          | C2              | P1            |
| **Cell dimensions**       |                 |               |
| a, b, c (Å)              | 111.35, 75.20, 74.74; | 70.75, 70.83, 76.04; |
| α, β, γ (°)              | 90.0, 110.7, 90.0 | 92.16, 101.44, 115.61 |
| Resolution (Å)           | 41.85–1.70 (1.71–1.70) | 61.97–1.70 (1.73–1.70) |
| Rmeas (%)                | 8.9 (60.2)      | 11.3 (252.2)  |
| I/σI                     | 13.3 (2.9)      | 6.0 (1.3)     |
| Completeness (%)         | 99.1 (95.7)     | 96.9 (89.3)   |
| Redundancy (%)           | 2.2 (2.3)       | 3.5 (3.7)     |
| CC1/2 (%)                | 99.2 (86.8)     | 98.9 (31.4)   |
| No. of unique reflections| 62494 (9672)    | 138636 (6454) |
| **Refinement**           |                 |               |
| Resolution (Å)           | 31.52–1.7 (1.71–1.7) | 59.62–1.7 (1.71–1.7) |
| No. of reflections /test set | 62474/3122    | 138595 / 6805 |
| Rwork/ Rfree (%)         | 15.3/18.1       | 19.8/23.0     |
| No. of protein residues  | 490             | 484 + 481     |
| No. atoms                |                 |               |
| Protein /Water           | 4040/287        | 7895/556      |
| Ligand (non-H atoms) L95+ L-pro/HFG+L95 | 36 (28 + 8) | 104 (48 + 56) |
| Ions/Solvent molecules×  | 1/16            | 2/42          |
| Average B-factors (Å²)   |                 |               |
| Protein /Water           | 31.6/37.9       | 35/45.4       |
| Ligand (non-H atoms) L95- L-pro/HFG + L95 | 21.7 (23.5/19.8) | 27 (27.1/28.8) |
| CI ion / MPD/MES         | 77.4/42.6       | 33.5          |
| R.m.s deviations         |                 |               |
| Bond lengths (Å)         | 0.018           | 0.016         |
| Bond angles (°)          | 1.508           | 1.400         |
| Ramachandran plot        |                 |               |
| Favoured/Allowed (%)     | 97.9 / 2.1      | 98 / 2        |
| Clash score              | 3.1             | 3             |
| Disordered Regions       | 408–415, 787–788 | 408–416, 784–790 |

* Ions: CI/Br and Solvent molecules: MPD, MES, EDO, IMD

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bound and apo TgPRS enzyme when compared to the L-pro-L95 bound structures (S7 Fig). Similar loop movements and side-chain conformation flips were also found between HsPRS-HFG-ANP (PDB ID: 4HVC) and HsPRS-ATP pocket ligand (PDB Id: 7OT3) [21,36]. The HFG binding loop (residues 405–420) is found to be disordered in almost all HFG bound structures (Fig 3B).

The above ligand-induced structural movements of loops are coherent with the differential ΔT_m values reported above. The similarity in ΔT_m values of TgPRS-HFG-L95 and TgPRS-L-pro-L95 suggest that the binding mode of L95 in the presence of L-pro mirrors that of L95 with HFG. However, the higher ΔT_m value for TgPRS-HFG-ANP reflects a more stable association of HFG with ATP than HFG with L95 or L95 with L-pro. Within the ternary complexes, the inhibitors/substrates contact each other peripherally and the interface area for HFG-L95, L-pro-L95 and, HFG-ANP are 27, 35, and 77 Å² respectively (S8 Fig and S1 Table). The significantly higher surface interface overlap between HFG and ANP in the complex is likely to be due to the phosphate groups between HFG and the adenine ring of ANP.

**Cell-based drug gradient assays indicate synergism between HFG and L95**

To explore the anti-parasitic effects of L95 and HFG drug combination, an ex vivo cell-based assay was done to monitor the intracellular growth of Toxoplasma gondii (T. gondii) tachyzoites through expression of NanoLuc luciferase (RH NLuc) in a human foreskin fibroblast (HFF) host-cell monolayer [47]. Firstly, the efficiency of L95 and HFG were determined in single drug assays using pyrimethamine as reference for parasite growth inhibition (Fig 4A). In these assays, both L95 and HFG inhibited parasite proliferation at nM concentrations with measured half maximum effective concentration (EC_{50}) of 160 and 1.6 nM respectively, which is more efficient than the pyrimethamine positive control (EC_{50} 257 nM) (Fig 4A). In parallel, we evaluated the cytotoxicity of these compounds against HFFs and determined a selectivity index (SI). The HFFs half-maximal cytotoxicity concentration (CC_{50}) for L95 and HFG was calculated to be 3.2 µM and 0.4 µM with a selectivity index (SI) of 20 and 250 respectively. The SI was calculated by comparing the T. gondii EC_{50} to the CC_{50} of HFF cells used for Toxoplasma cultures (Fig 4B and 4C). These data indicate that L95 has a parasite selectivity, which is intriguing considering its inhibitory effectiveness against HsPRS enzyme (See Figs 1C, 1D and 4). To investigate the combinatorial effects of L95 and HFG against T. gondii, we performed a multi-drug combination profiling analysis of parasite growth inhibition. The observed drug combination responses (the input dose-response matrix can be seen in Fig 5A) were compared with expected responses calculated by means of the Zero Interaction Potency (ZIP) synergy reference model using the SynergyFinder web-application to identify any synergistic/additive effects [48,49]. The ZIP score measures synergy by comparing the change in potency of dose-response curves between individual drugs and their combination [48,49]. Fig 5B shows an overall additive score of 10.12 with a synergistic area score of 31.96 for L95 concentrations ranging from 12 to 75 nM in combination with HFG ranges of 1.5 to 2.5 nM. Together these results indicate the potency of L95 and HFG inhibitor combination to inhibit T. gondii growth and reveal an overall additive effect when used in combination.

**Discussion**

Aminoacyl-tRNA synthetases (aaRSs) have over the last decade evolved as effective, novel and specific targets for many disease conditions [16,20,21,24,31,32,45]. The aaRS inhibitors are now also being used as scaffolds for anti-oncogenic drug development [34,35]. Inhibition of these enzymes stalls protein translation thus providing a roadblock in the further development of the parasite or diseased cell [16,20,31,32,45,50,51].
Here, we tested two inhibitors with distinct mechanisms of action. L95 and HFG act wherein the former inhibits PRS activity by binding to the ATP site while the latter binds to the L-pro and tRNA binding sites. Therefore, the prospective therapeutic combination of HFG and L95 occupies all three substrate sites, rendering the target completely blocked and inactive.

The study highlights an ‘ortho-steric cocktail’—a strategy of simultaneous targeting of multiple pockets within the active site by two different ligand scaffolds. The thermal shift profile of L95 in presence of L-pro implies a strong complex. Further, both TgPRS-L-pro-L95 and TgPRS-HFG-L95 complexes have similar thermal profiles (Fig 1B). Interestingly, the TgPRS-HFG-ATP complex has a higher ΔT_m value (37.6˚C) than those of TgPRS-L-pro-L95 (20.7˚C) and TgPRS-HFG-L95 (23.6˚C) (Fig 1B). This is most likely because of the ligand induced movement of loops and consequent differences in the side chain conformation (particularly Arg470 and Phe534).

The potencies of L95 and HFG against Toxoplasma parasites and TgPRS enzyme were validated via enzymatic and cell-based assays. Both IC_{50} and EC_{50} data revealed that these drugs are effective at nM concentrations when used individually (Figs 1C, 1D and 4). In context of selectivity, HFG was more selective against T.gondii parasite and the parasitic enzyme than...
L95 (cellular SI of HFG = 250 vs cellular SI of L95 = 20; enzymatic SI of HFG = ~2 vs enzymatic SI of L95 = ~1). In thermal stability, the \( T_g \) of the PRS-HFG-L95 combination had a higher \( \Delta T_m \) value than each ligand alone (23.6˚C compared to 8.1˚C for PRS-HFG and 17.8˚C for PRS-L95), suggesting that when used in tandem, they could improve \( T_g \) inhibition in an additive fashion (Figs 1B and S2). When the cell-based drug-combination assay was performed using a 7x7 matrix, we discovered that these compounds act synergistically when used in a specific range at low doses (see the most synergistic area with a high synergy score of 31.96; Fig 5). However, the overall synergy score of ~10 proves that the inhibitor combination is more likely to have a general additive effect. Indeed, HFG is more active than L95, and we cannot rule out the possibility that if HFG is used at higher concentrations, its effect would mask L95 activity.

The similarities observed in the structures of PRS-L-pro-L95 and PRS-HFG-L95 indicate that the potency of L95 remains unaffected in presence of either L-pro or HFG, which would be consistent with an additive activity at concentrations close to the IC\(_{50}\). This is in contrast to the cooperative binding of ligands thus indicating a gap in our understanding of the structural machinations of this approach.

The authors understand the limitations of dual targeting using these exact scaffolds and the caveats that come along with it for bradyzoite targeting, which remains critical for Toxoplasma targeting. This exact therapeutic combination might not be of ultimate utility considering the similarity in PK properties and potential toxicity of both these compounds. Bioavailability and cost feasibility would remain a critical bottleneck for the actual medical translation of this combination therapy. Furthermore, as far as the possibility of off-site targets by ATP-mimetics is concerned–previously characterized inhibitors have given ample evidence for the highly specific nature of ATP recognition by enzymes and why such an occurrence is rare [16]. Here we provide proof that two ligand moieties can simultaneously occupy all catalytic pockets of an
essential aaRS enzyme (here, TgPRS). Indeed, our approach can be extended to other enzyme-drug systems where novel assays screen for potency as well as synergy. Drug targeting of proteins has so far relied on finding single inhibitors of high potency but our novel data show that it is feasible to design two (or more) inhibitors that may simultaneously attack any available druggable pockets in target proteins. Borrelidin is a natural product and is an example of a simultaneous multi-site aaRS inhibitor. Designing and producing such molecules synthetically has many caveats including but not limited to selectivity and cost feasibility. The future prospects of such and other multi-site inhibitors remains to be explored in a systematic manner. Our approach detailed here will allow rescreening of even known enzyme-drug combinations with new compound libraries in order to discover synergistic inhibitors that can co-occupy and co-inhibit the same target enzyme.

**Materials and methods**

**Protein purification**

TgPRS was produced in an *E. coli* BL-21 strain in accordance with methods published previously [20,45]. Briefly, protein expression was induced by adding 0.5 mM isopropyl β-D-thiogalactopyranoside (IPTG) to cells grown at 37˚C for 4 h, and for 20 h post-induction at 18˚C. The cells were harvested by centrifugation at 5,000 g for 15 min. The bacterial pellet was suspended in a buffer containing 50 mM Tris–HCl pH 8.0, 200 mM NaCl, 3 mM ßME, 15% v/v glycerol, 0.1 mg ml⁻¹ lysozyme and EDTA free protease inhibitor cocktail (Roche). The cells were lysed by sonication and cleared by centrifugation at 20,000 g for 45 min. The cleared supernatant with maltose-binding protein (MBP) and 6xHis tagged proteins were applied to NiNTA beads (GE Healthcare) and protein was eluted with buffer 50 mM Tris–HCl pH 8.0, 200 mM NaCl, 10 mM ßME and 250 mM Imidazole. The eluted protein fractions were dialyzed against 30 mM HEPES pH 7.5, 20 mM NaCl, 1 mM DTT and 0.5 mM EDTA (buffer A). The protein was further purified by heparin chromatography (GE Healthcare) using NaCl gradients with buffer B containing 30 mM HEPES pH 7.5, 500 mM NaCl, 1 mM DTT and 0.5 mM EDTA. The protein peak was found at 40% buffer B. The MBP and 6xHis tag was removed by incubating with TEV protease at 20˚C for 24 h. The cleaved TgPRS protein was concentrated using 10 kDa cut-off Centricon centrifugal device (Millipore) and purified by gel filtration chromatography on a Superdex 200 column 16/60 GL (GE Healthcare) equilibrated with 20 mM HEPES pH 7.5, 200 mM NaCl and 2 mM DTT. Bovine serum albumin (66 kDa, Sigma) was used as a standard for the molecular mass estimation. The eluted fractions were checked by SDS-PAGE and the pure fractions were pooled, concentrated in 50 mM Tris–HCl pH 8.0, 200 mM NaCl, 10 mM ßMe and stored at -80˚C.

**Thermal shift assays**

Thermal shift assays were performed for TgPRS and HsPRS in the presence of substrates (L-pro and ATP) and inhibitors (L95 and HFG) as per previous protocol [20,45]. Briefly, the purified enzymes (1 μM) alone and/or with substrates/inhibitors (2 mM/50 μM) were heated from 25 to 99˚C at a rate of 1.2˚C min⁻¹ and fluorescence signals of the SYPRO orange dye was monitored by a quantitative real-time PCR system (Life Technologies). Note that titration experiments determined that saturation of active pocket enzymes was obtained at 1:50 protein:compound ratio (S2B Fig). The melting curve is an average of three measurements and data were analysed using Protein Thermal shift software (v1.3, Thermofisher). The inhibitors and substrate alone in assay buffers, along with no enzyme controls were used and flat lines were observed for these fluorescence readings across the temperatures. The derivative Tₘ was used for analysis.
Enzyme inhibition assays

These were done using malachite green assay and the assay was performed according to the earlier published report [20]. Briefly, the reaction was observed for 25 μM ATP, 25 μM L-pro and 400 nM recombinant PRS enzyme in a buffer containing 30 mM HEPES (pH 7.5), 140 mM NaCl, 30 mM KCl, 50 mM, MgCl₂, 1mM DTT and 2 U/ml E. coli inorganic pyrophosphatase (NEB) at 37˚C. The enzymatic reactions (100 μl total volume) were performed in a clear, flat-bottomed, 96-well plates (Costar 96-well standard microplates). The reaction mixture was incubated for 40 min at 37˚C. The reaction was stopped by adding 25 μl of malachite green dye solution to the reaction mixture and levels of inorganic phosphate (Pi) were detected after incubation of 5 min at room temperature. Absorbance was measured at 630 nm using a Spectramax M2 (Molecular Devices). Reactions without PRS enzyme were performed as background controls, values of which were subtracted from the reactions with enzyme. HFG and L95 were added to the aminoacylation assay reaction buffer in varying concentrations ranging from 50 to 0.005 μM. The IC₅₀ values for the data is shown for three replicates as the mean ± SD.

Crystallization and structure determination

Purified TgPRS enzyme was used for crystallization via the hanging-drop vapour-diffusion method at 20˚C using commercially available crystallization screens (Index, JCSG⁺, Morpheus, PACT premier, PGA, Crystal Screen, PEG/Ion and ProPlex (Hampton Research and Molecular Dimensions). Initial screening was performed in 96-well plates using a nano drop dispensing Mosquito robot (TTP Labtech). Three different drop ratios were used for the crystallization trials by mixing 75, 100 or 50 nl purified protein solution with 75, 50 or 100 nl reservoir solution, respectively (i.e., 1:1, 2:1 and 1:2 drop ratios). Each of the drops was equilibrated against 100 μl of the corresponding reservoir solution. Before crystallization, 1 mM L95 and 2 mM L-pro were added to TgPRS in one set up and with HFG in another while the mixtures were incubated at 4˚C for 10 min. Diffraction quality crystals of ternary complexes were obtained at 20˚C by the hanging-drop vapour-diffusion method using 50 nl each of the protein mixture solutions and 50 nl crystallization well solution. The TgPRS-L-pro-L95 crystals were obtained in B5 of JCSG⁺ screen (0.1 M sodium cacodylate pH 6.5 and 40% v/v MPD 5% w/v PEG 8000). The TgPRS-HFG-L95 crystals were obtained in Morpheus G2 screen [0.1 M carboxylic acids (0.2 M sodium formate, 0.2 M ammonium acetate, 0.2 M sodium citrate tribasic dihydrate, 0.2 M potassium sodium tartrate tetrahydrate, 0.2 M sodium oxamate), 0.1 M buffer pH 6.5 (imidazole; MES monohydrate (acid) and 30% v/v precipitant mix (40% v/v Ethylene glycol; 20% w/v PEG 8000)]. Plate/rod-shaped crystals appeared within 10 days. The crystals were mounted in nylon loops (Hampton Research) or litho loops (Molecular Dimensions) after being soaked for 10–30 s in a cryoprotectant containing the corresponding crystallization mother liquor with 20% (v/v) glycerol. The crystals were subsequently flash-cooled in liquid nitrogen. The X-ray diffraction data sets were collected on beamline I03 at Diamond Light Source (DLS), United Kingdom at a wavelength of 0.976 Å for the TgPRS-HFG-L95 structure while for the TgPRS-L-pro-L95 structure at PROXIMA-1, SOLEIL, France at a wavelength of 0.979 Å. The data were processed by the Xia2 auto-processing pipeline using DIALS for integration [52]. The initial models were determined by the molecular-replacement (MR) method using Phaser [53] with A chain of TgPRS-HFG-ANP (PDB ID 5XIQ) without bound ligands (HFG and ANP) as the template. The structure was further refined by iterative cycles of refinement with Phenix [54] and model building with COOT [55]. The final models were refined to 1.7 Å resolution with Rwork/Rfree values of 15.3/18.1 and 19.8/23.0%, respectively for ternary complexes TgPRS-L-pro-L95 and TgPRS-HFG-L95. The stereo-chemical quality of the models
was assessed and corrected using MolProbity [56]–finally showing good geometry quality with 98% residues in most favoured regions of the Ramachandran plot for the TgPRS-L-pro-L95 and TgPRS-HFG-L95 complexes, respectively. Statistics of data collection and structure refinement are given in Table 1. The atomic coordinates and structure factors of TgPRS-L-pro-L95 and TgPRS-HFG-L95 complexes have been deposited in the Protein Data Bank (PDB) with the accession codes 7EVV and 7EVU, respectively. Surface areas for the protein and ligand molecules were computed using the PDBePISA server [57]. The figures were prepared using Chimera [58] and PyMOL [59].

Measurement of EC\textsubscript{50} for Toxoplasma gondii parasites

A confluent Human Foreskin Fibroblast (HFF) monolayer was infected with 2,000 tachyzoites of RH parasites expressing the NLuc luciferase (RH NLuc) for 2 h. After parasites invasion, each drug was added to the culture medium. After 48 h of incubation at 37°C, the medium was replaced by 50 μL of PBS. The luminescence assay was performed using a Luciferase Assay System according to manufacturer’s instructions (Promega). After 3 min of incubation, luminescence was measured using a (BMG Labtech) plate reader. EC\textsubscript{50} were determined using non-linear regression analysis of normalized data and assuming a sigmoidal dose response. EC\textsubscript{50} values for each compound represent an average of three independent biological replicates. Statistical analyses were performed using one-way ANOVA test with the GraphPad software.

Measurement of CC\textsubscript{50} for human cells and determination of selectivity index

10,000 HFFs were plated in 96-well plates for 1 h and incubated with exponential concentrations of indicated compounds in a final volume of 100 μL. After 72 h of culture the CellTiter-Blue dye (Promega) (20μL/well) was added directly to each well. The plates were then incubated at 37°C for 2 h to allow cells to convert resazurin to resorufin before recording fluorescence (560(20) Ex/590(10) Em) with a (BMG Labtech) plate reader. Human cells cytotoxicity concentration (CC\textsubscript{50}) was determined using non-linear regression curve of normalized data. The CC\textsubscript{50} values represent the average of four biological experiments. Selectivity Index (SI) was obtained by the average of the human CC\textsubscript{50} divided by the average of T. gondii EC\textsubscript{50}.

Gradient combinational assay

Drug dilutions used to determine the dose-response curves were utilized to predict the drug combination response. We defined a 7x7 dose matrix (% inhibition range between 0 and 100) on 96-well plates. Confluent HFFs monolayer was infected with RH NLuc parasites for 2 h before adding drugs combination. After 48 h of culture the medium was replaced by PBS and the luminescence was read as described above. The overall synergy score for a drug pair is calculated based on Zero Interaction Potency (ZIP) reference model using SynergyFinder web tool. Deviations between predicted and expected responses with positive and negative score denote synergism and antagonism, respectively [48,49]. When synergy score is < -10, the interaction between two drugs is likely to be antagonistic; from -10 to 10, the interaction between two drugs is likely to be additive; and >10, the interaction between two drugs is likely to be synergistic. The most synergistic area represents the most synergistic 2-by-2 dose-window in a dose-response matrix.

Supporting information

S1 Fig. (a) Chemical structure of ligand L95 and its functional groups. (b) Structural overlay of L95 with ATP. (TIFF)
S2 Fig. TSA Melt Curves (a) TgPRS melt curves with different combinations of inhibitors and substrates. (b) TgPRS differential ratios for saturation of protein. (c) HsPRS melt curves with different combinations of inhibitors and substrates. (TIFF)

S3 Fig. The difference Fourier map (Fo-Fc) of the bound ligands in the aminoacylation site of chain B of TgPRS-HFG-L95 complex. The Fo-Fc maps are contoured at 3 σ and shown as green mesh representations. Ligands L95 (yellow), HFG (pink) are shown as sticks. (TIFF)

S4 Fig. Important interactions between ligand L95 with protein residues and water molecules. Ligand L95 (yellow) and protein residues (purple) are shown as sticks representations. The water molecules are shown as red spheres. Hydrogen bonds are shown as black dashed lines. (TIFF)

S5 Fig. The 6-methylpyridine ring of ligand L95 sandwiched between Arg594 and Phe485 via π-π stacking interaction (~3.9 Å). (TIFF)

S6 Fig. Overlay of L-pro-L95 (purple) bound structures with apo form of TgPRS (pink) showing the different conformations of active site residues Arg470, Arg594 and F534, and the ATP binding loop. (TIFF)

S7 Fig. Overlay of apo (pink), L-pro-L95 bound (purple), HFG-L95 bound (steel blue) and HFG-ANP bound (olive) TgPRS structures showing the displacement of the loop residues 528–538. (TIFF)

S8 Fig. Molecular surface representation showing the association of bound ligands in tertiary complexes of TgPRS-L-pro-L95, TgPRS-HFG-L95 and TgPRS-HFG-ANP. The TgPRS surface is displayed as 70% transparent surface and the bound ligands surface are shown as solid surfaces. (TIFF)

S1 Table. Buried surface areas of the inhibitors and substrates within TgPRS. (XLSX)

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