Selective and Specific Inhibition of the Plasmodium falciparum Lysyl-tRNA Synthetase by the Fungal Secondary Metabolite Cladosporin

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SUMMARY

With renewed calls for malaria eradication, next-generation antimalarials need to be active against drug-resistant parasites and efficacious against both liver- and blood-stage infections. We screened a natural product library to identify inhibitors of Plasmodium falciparum blood- and liver-stage proliferation. Cladosporin, a fungal secondary metabolite whose target and mechanism of action are not known for any species, was identified as having potent, nanomolar, antiparasitic activity against both blood and liver stages. Using postgenomic methods, including a yeast deletion strains collection, we show cladosporin specifically inhibits protein synthesis by directly targeting P. falciparum cytosolic lysyl-tRNA synthetase. Further, cladosporin is >100-fold more potent against parasite lysyl-tRNA synthetase relative to the human enzyme, which is conferred by the identity of two amino acids within the enzyme active site. Our data indicate that lysyl-tRNA synthetase is an attractive, druggable, antimalarial target that can be selectively inhibited.

INTRODUCTION

Malaria is a significant health problem, with 225 million annual cases and nearly 3.2 billion people at risk (WHO, 2010). Control and treatment of this disease is compounded by a lack of an effective vaccine. In addition, the emergence of multidrug-resistant parasites has compromised efficacy of many of the frontline chemotherapy treatments. Although there are many effective drugs (Burrows et al., 2011), endoperoxides are the only drug class for which clinically significant resistance has not been reported (Eastman and Fidock, 2009). However, endoperoxides, like many antimalarials, are inactive against the asymptomatic malaria liver stages. To ensure continued malaria control, with an aim for eradication, next-generation antimalarials are required to be active against multidrug-resistant parasites and efficacious against liver- and blood-stage infections.

Traditional drug discovery efforts have revolved around high-value targets identified for their essentiality in the parasite. However, finding targets that are essential for both blood and liver malarial stages has been technologically challenging, and few inhibitors with these desirable properties are known because of the difficulties associated with simultaneously demonstrating that a target is essential to support viability in both blood and exoerythrocytic stages. An alternative approach to target discovery is to first find compounds with promising activity in phenotypic cell-based screens (Gamo et al., 2010; Guiguemde et al., 2010; Plouffe et al., 2008) and then determine their mechanism of action through the identification of their specific targets (reviewed in McNamara and Winzeler, 2011). Successful validation of a target from this latter approach provides a proof of concept for small-molecule inhibition and supports continued drug discovery based on rational design of the hit compound.

In order to discover targets for both blood and liver stages, we performed a screen to identify inhibitors of P. falciparum blood- and liver-stage proliferation with a natural product library. Cladosporin, a fungal secondary metabolite whose primary target and mechanism of action are not known for any species, was identified as having potent, nanomolar, antiparasitic activity in both blood and liver stages. A member of the isocoumarin class, cladosporin is produced by various fungal genera such as Cladosporium, Aspergillus, Eurotium, and Chaetomium (Scott et al., 1971). It has been previously reported to have antifungal (Scott et al., 1971), insecticidal (Grove and Pople, 1981), and antibacterial properties (Anke, 1979) as well as plant growth regulatory effects (Springer et al., 1981) and anti-inflammatory
results in mouse lung tissue (Miller et al., 2010). Using both traditional and systems biology approaches, we show here that cladosporin potently and specifically inhibits cytosolic lysyl-tRNA synthetase in Plasmodium spp. In addition, we show that cladosporin is highly selective for the parasite enzyme and that selectivity is in part conferred by the amino acid identity at two key residues in the ATP binding pocket.

RESULTS

A Natural Product Screen Identifies Cladosporin with Potent Antiplasmodial Activity

Small molecules with activity against P. falciparum blood-stage parasites were previously identified in a phenotypic screen against a natural product library (Plouffe et al., 2008). Out of the 12,000 natural products, 275 compounds inhibited parasite growth with 50% inhibitory concentration (IC50) values in the submicromolar range. These hits were further screened by a high-content image-based assay to determine their ability to block in vitro P. yoelii liver-stage development (Meister et al., 2011). Cladosporin (Figure 1A) demonstrated exceptional growth-inhibitory activities against both blood- and liver-stage parasite forms (IC50 ~40–90 nM) while having little effect on the growth or viability of HepG2-CD81 cells (>10 μM) or other human cell lines (Table 1). The high selectivity index of cladosporin against Plasmodium parasites compared to mammalian cells (IC50/CC50 ≥ 111), as well as its equipotency against a diverse collection of multidrug-resistant Plasmodium lines (Table 1) and cidal action (see Table S1 available online), suggested that further investigation was warranted.

Chemogenomic Profiling Reveals that Cladosporin Targets Yeast Lysyl-tRNA Synthetase

We next sought to discover the target of cladosporin. Cladosporin demonstrated high micromolar inhibition against the model yeast, S. cerevisiae, although activity varied slightly due to compound batch and the growth media conditions (summarized in Table S2). In rich growth media, cladosporin displays a 30% inhibitory concentration (IC50) of 110 μM, making it a suitable compound for haploinsufficiency profiling (HIP) assay (Giaever et al., 1999). The HIP target discovery assay is based on a genome-wide collection of heterozygous knockout yeast strains, each of which contains a marked gene deletion (Giaever et al., 2002; Winzeler et al., 1999). It has been shown that heterozygous diploid strains that bear a deletion in one copy of a small molecule’s target gene show increased sensitivity to that small molecule’s target gene show increased sensitivity to that small molecule relative to those strains that have two copies of the gene (Giaever et al., 1999). This is a likely consequence of decreased expression of the target in the heterozygous diploid strain. Each of these strains also carries a distinct 20 base pair DNA barcode that can be PCR amplified and quantified in competitive growth assays using microarrays containing the barcode complements. As each barcode is linked to a specific strain, barcodes that disappear over time in the presence of a small molecule reveal the identity of the molecular target of the small molecule.

Three independent HIP assays reproducibly identified the heterozygous lysyl-tRNA synthetase (krs1/KRS1) knockout strain as being the only strain in the collection of 5,803 total strains (covering ~95% of the genome) that was hypersensitive to cladosporin (Figures 1B and 1C). Because strains that bear deletions in drug pumps or other proteins that attenuate the action of small molecules often also show haploinsufficiency, we also assessed whether the krs1/KRS1 strain is ubiquitously sensitive. Historical profiling of 1,800 compounds (our unpublished data) revealed that the krs1/KRS1 strain was exclusively sensitive to cladosporin (Figure 1D), indicating a high degree of selectivity for cladosporin. KRS1 is a nonredundant and essential gene in yeast and functions to load lysine onto the corresponding tRNA molecule for protein translation (Mirande and Waller, 1988).

Overexpression of KRS1 Decreases Cladosporin Antifungal Activity

To further test if cladosporin specifically inhibits lysyl-tRNA synthetase, KRS1 was placed downstream of a Gal1 promoter and transformed into wild-type yeast. The same was done for genes encoding isoleucyl-tRNA synthetase (ILS), glutamyl-tRNA synthetase (GLN4), and threonyl-tRNA synthetase.
We took advantage of this conservation and used the human crystal structure as a template to generate a ScKrs1 homology model. The ScKra1 model predicted that the identified mutations, Ile567Val (one strain), Gly551Ser (two strains), and Thr340Ile (seven strains), map to or near the binding pocket for ATP in ScKrs1 (Figures 2B and 2C and Figure S2A). Backcrossing of the ten selected mutants to isogenic wild-type strains of the opposite mating type and subsequent resistance assays revealed that all mutations were dominant. Subcloning of each of the mutated KRS1 genes into a low copy plasmid and reintroduction into the wild-type strain was sufficient to confer increased resistance to cladosporin (~5-fold increase), indicating that second-site mutations did not contribute to resistance. We conclude that the mutated KRS1 gene is responsible for resistance in S. cerevisiae and that the resistance phenotype is dominant to the wild-type allele (Figure 2D).

Amplification of the Lysyl-tRNA Synthetase Locus Confers Resistance in Parasites

To establish target conservation in P. falciparum, we evolved cladosporin-resistant parasites (Table 1 and Tables S3 and S4) with hopes that the parasites would acquire resistance mutations in the target gene. The genomes of the cladosporin-sensitive parental clone and cladosporin-resistant clones were compared using a microarray that has the capacity to reveal most genomic changes that are acquired by the resistant strains, including copy number variants, deletions, small insertion/deletion events, and single-nucleotide polymorphisms (Dharia et al., 2009). Since background mutations may appear at a low frequency, the selection was run in triplicate in order to distinguish resistance-conferring mutations from random mutations. Genome scanning revealed that each of the independent clones had acquired copy number variants that shared six genes in a syntenic region of chromosome 13: MAL13P1.254, and MAL13P1.255 and PF13_0262, encoding threonyl-tRNA synthetase (THS1). These strains were subsequently tested for growth defects in the presence of serially diluted cladosporin concentrations using galactose as a sole carbon source to drive promoter activity. Only the transgenic strain harboring the plasmid with the KRS1 gene conferred increased cladosporin resistance (2.2-fold shift), whereas the transgenic strains expressing other aminoacyl-tRNA synthetases (AARSs) exhibited no change in their sensitivity to cladosporin (Figure 2A). In the absence of cladosporin, all strains grew at an equal rate and concurrently reached the same final OD600, indicating the overexpression constructs did not have a discernible effect on growth.

Cladosporin Resistance in Yeast Is Mediated by Mutations in the Active Site of Krs1

To further strengthen the observed genetic link between the S. cerevisiae lysyl-tRNA synthetase (ScKrs1) and cladosporin, we examined mutations that confer cladosporin resistance in yeast. Yeast cells were mutagenized with ethylmethanesulfonate and then selected against 50 μM cladosporin, a concentration known to completely inhibit growth of wild-type strains in synthetic complete media (Table S2). Direct sequencing of the KRS1 gene in 37 clones revealed nonsynonymous mutations in ten of these clones (Figure 2B and Figure S2A).

To better understand the significance of these mutations, we undertook homology modeling. The crystal structures of lysyl-tRNA synthetase from human (Guo et al., 2008) and two bacteria (Sakurama et al., 2009) have been published, and comparison reveals a high degree of structural conservation (Figure S2B).
Cladosporin Inhibits Cytoplasmic Protein Biosynthesis in *P. falciparum*

Malaria parasites have two lysyl-tRNA synthetases, one of which is targeted to the apicoplast (PF14_0166), a specialized subcellular organelle involved in fatty acid and isoprenoid biosynthesis, while the other one is targeted to the cytoplasm (PF13_0262). The apicoplast is essential to parasite viability, and inhibition of protein synthesis in this organelle yields a characteristic "delayed death" phenotype in which parasite death is observed one generation after drug treatment. Known apicoplast inhibitors include mupirocin, which targets the apicoplast-targeted isoleucyl-tRNA synthetase (Istvan et al., 2011); tetracycline to its activity in cellular screens (IC50 40–90 nM; Table 1). As expected, no activity was observed in the absence of exogenous product, AMP. Cladosporin possessed a half inhibitory concentration (IC50) of 61 nM against *Plasmodium falciparum* lysyl-tRNA synthetase (ScKrs1) protein domain organization. Resistance-conferring mutations (asterisks) are labeled and frequency of mutation provided in parentheses.

Bacterial antibiotics are known to inhibit protein synthesis in the cytoplasm by targeting ribosomal subunits. Eukaryotic protein synthesis also follows this paradigm, but ribosomes have an additional key step in the protein synthesis process, the recruitment of aminoacyl-tRNAs to the ribosomal A site, requiring a functional lysyl-tRNA synthetase. Thus, we decided to confirm if cladosporin inhibited *P. falciparum* lysyl-tRNA synthetase (PfKrs1).

The phenotype of cladosporin-treated parasites is indistinguishable from that of anisomycin, another eukaryotic protein synthesis inhibitor, and supports the notion that both have the same mechanism of action. In contrast, clindamycin, which specifically inhibits protein biosynthesis in the apicoplast, did not diminish growth until the end of the second blood-stage life cycle (Figure S3). Although these results do not preclude cladosporin as also targeting the apicoplast lysyl-tRNA synthetase, these data show that cladosporin is a fast-acting agent that shows the same characteristics as other known cytoplasmic protein synthesis inhibitors.

Cladosporin Is a Potent and Selective Inhibitor of *P. falciparum* Lysyl-tRNA Synthetase

To confirm that cladosporin is an inhibitor of *PfKrs1*, direct biochemical assays were performed on recombinant protein. *PfKrs1* activity was assayed in the presence or absence of the corresponding lysine-tRNA (tRNALys) substrate, and synthetase activity was measured using a Transcreener adenosine monophosphate (AMP) assay. This assay is a far-red, competitive fluorescence polarization immunoassay that detects the reaction product, AMP. Cladosporin possessed a half inhibitory concentration of 61 nM against *PfKrs1* (Figure 4A), which is comparable to its activity in cellular screens (IC50 40–90 nM; Table 1). As expected, no activity was observed in the absence of exogenous tRNA Lys. The agreement between the biochemical and biological inhibition constants further supports the notion that lysyl-tRNA synthetase is the primary target within the cell.

Our data show that cladosporin has relatively little activity on human cells, including the HepG2-CD81 cells used in the hepatic-stage screen (Table 1). To confirm that cladosporin is selective for *PfKrs1*, the human counterpart was assayed. Recombinant human lysyl-tRNA synthetase was assayed as described above. As predicted, cladosporin activity was only weakly detected at the high micromolar ranges for human Krs1 (Figure 4B). These data reinforce the notion that cladosporin specifically and selectively targets *P. falciparum* lysyl-tRNA synthetase.

Structural Basis for Selectivity and Specificity in *Plasmodium Krs1*

The *S. cerevisiae* and *P. falciparum* amino acid sequences share a high degree of identity to the human protein (yeast 58.3%, human 43.8%). To confirm that cladosporin is an inhibitor of *PfKrs1*, direct biochemical assays were performed on recombinant protein. *PfKrs1* activity was assayed in the presence or absence of the corresponding lysine-tRNA (tRNALys) substrate, and synthetase activity was measured using a Transcreener adenosine monophosphate (AMP) assay. This assay is a far-red, competitive fluorescence polarization immunoassay that detects the reaction product, AMP. Cladosporin possessed a half inhibitory concentration of 61 nM against *PfKrs1* (Figure 4A), which is comparable to its activity in cellular screens (IC50 40–90 nM; Table 1). As expected, no activity was observed in the absence of exogenous tRNA Lys. The agreement between the biochemical and biological inhibition constants further supports the notion that lysyl-tRNA synthetase is the primary target within the cell.

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plasmodium 53.4%), and the homology model could accommodate both primary sequences into the human template with high confidence. According to the predicted models, 30 of 36 amino acids within a 5 Å radius of the ATP/lysine-binding site are conserved between S. cerevisiae and P. falciparum. Using an in silico docking approach it was possible to model cladosporin into the ATP-binding pocket in an energetically favorable mode, mimicking that of ATP (Figure 4C). Residues Asn335 and Glu328, which hydrogen bond to ATP, are also predicted to hydrogen bond with the hydroxyl groups of cladosporin. The accuracy of this prediction was validated biochemically. Characterization of the recombinant PfKrs1 revealed that lysine had no effect on cladosporin inhibition (Figure S4A), whereas increasing concentrations of ATP in the reaction buffer significantly reduced cladosporin inhibition (Figure S4B). This was corroborated by the inability of exogenous lysine in culturing media to affect cladosporin potency against yeast cells (Table S5). Taken together, these data support the notion that cladosporin interacts with...
Cladosporin Selectively Targets P. falciparum Krs1

To further understand the basis of cladosporin selectivity, the amino acid residues of the ATP-binding pocket were compared to those in Krs1 of other species (Figure S4C). While the majority of amino acid residues in the binding pocket were highly conserved, a clear divergence existed at two amino acid positions corresponding to S. cerevisiae residues Gln324 and Thr340 (Table 2). In Plasmodium spp., these positions are occupied by a valine and serine residue, respectively. Based on the identity of the Plasmodium amino acid residues, smaller side chains are favored, presumably due to decreased steric hindrance. Additionally, a hydrophobic residue at position 324 also appears favorable for cladosporin binding.

Despite being separated by 15 amino acids in the primary sequence, the homology model predicted that these residues were juxtaposed in the active site (Figure S4D). The docking model also predicts that the pyrene moiety of cladosporin points toward both of these residues and that Thr340 resides at an intimate distance of 3.7 Å (Figure 4C and Figure S4D). Extending the comparison to other eukaryotic and prokaryotic pathogens with Krs1 orthologs, a clear correlation exists between cladosporin activity and the identity of the amino acids at these two key positions in the ATP-binding pocket (Table 2). The trend predicts a significant loss in cladosporin potency whenever a bulkier residue (replacement of serine with threonine) is present at position 324. If this mutation is accompanied by a modestly more bulky and polar residue at position 324 (valine to glutamine or asparagine), then an even more significant decrease in activity is observed. The presence of a methionine at position 340 in bacteria species, which would likely impose an even greater steric hindrance than threonine, leads to the most significant loss in cladosporin potency.

To validate whether ScKrs1 positions 324 and 340 are significant factors in cladosporin selectivity and specificity, these positions were mutated in yeast to mimic those residues found in Plasmodium spp. Substantial increases in cladosporin potency were observed for both mutations. The single mutation Gln324Val produced a 5.7-fold increase in cladosporin potency compared to wild-type S. cerevisiae strain (IC50 = 163 μM; triangles). The double mutant, ScKrs1-Gln324Val/Thr340Ser, shows a 1,000-fold increase in sensitivity to cladosporin (IC50 = 4 μM; circles), which more closely resembles PfKrs1, is 41-fold more sensitive to cladosporin. See also Figure S4. Enzymatic data were from at least two independent assays performed in triplicate and expressed as means ± SD. A nonlinear regression curve fit is shown for each.

The ATP-binding pocket of lysyl-tRNA synthetase reveals that cladosporin has low nanomolar inhibition (IC50 = 61 nM) against PfKrs1 (A, squares) and high micromolar activity (IC50 > 20 μM) against human Krs1 (B, squares). As a control, recombinant enzyme was assayed under the same conditions but in the absence of the tRNA^lys^ substrate (circles). Enzymatic data were from at least two independent assays performed in triplicate and expressed as means ± SD. A nonlinear regression curve fit is shown for each.

(C) Superimposed structures of cladosporin (green) and ATP (cyan) docked to the yeast homology model (gray). Oxygen and nitrogen atoms are colored red and blue, respectively, in all molecules. Cladosporin is predicted to bind in the ATP-binding pocket of the yeast homology model with the isocoumarin moiety located in the same region as the adenine of ATP. Also, cladosporin’s pyrane moiety superimposes between the sugar and the phosphate of ATP and projects toward Gln324 and Thr340. The two hydroxy groups of cladosporin are predicted to form hydrogen bonds with Asn335 and Glu328 (dotted lines; bond length label given in Å).

(D) ScKrs1-Gln324Val mutant (IC50 = 28 μM; squares) and ScKrs1-Thr340Ser mutant (IC50 = 16 μM; diamonds) mimic the differences in the Plasmodium ATP pocket and significantly increase the potency of cladosporin compared to wild-type S. cerevisiae strain (IC50 = 163 μM; triangles). The double mutant, ScKrs1-Gln324Val/Thr340Ser (IC50 = 4 μM; circles), which more closely resembles PfKrs1, is 41-fold more sensitive to cladosporin. See also Figure S4. Enzymatic data were from at least two independent assays performed in triplicate and expressed as means ± SD. A nonlinear regression curve fit is shown for each.

The model also predicts that the pyrene moiety of cladosporin points toward both of these residues and that Thr340 resides at an intimate distance of 3.7 Å (Figure 4C and Figure S4D). Extending the comparison to other eukaryotic and prokaryotic pathogens with Krs1 orthologs, a clear correlation exists between cladosporin activity and the identity of the amino acids at these two key positions in the ATP-binding pocket (Table 2). The trend predicts a significant loss in cladosporin potency whenever a bulkier residue (replacement of serine with threonine) is present at position 324. If this mutation is accompanied by a modestly more bulky and polar residue at position 324 (valine to glutamine or asparagine), then an even more significant decrease in activity is observed. The presence of a methionine at position 340 in bacteria species, which would likely impose an even greater steric hindrance than threonine, leads to the most significant loss in cladosporin potency.

To validate whether ScKrs1 positions 324 and 340 are significant factors in cladosporin selectivity and specificity, these positions were mutated in yeast to mimic those residues found in Plasmodium spp. Substantial increases in cladosporin potency were observed for both mutations. The single mutation Gln324Val produced a 5.7-fold increase in cladosporin sensitivity, whereas an even more pronounced increase by 10.4-fold was observed for the Thr340Ser mutant (Figure 4D). As expected, the double mutant Gln324Val/Thr340Ser led to an even greater sensitization of yeast to cladosporin (38.7-fold increase), as the active site of this mutant more closely resembled that in Plasmodium spp. (Figure 4D). Finally, because mutating two residues might alter the efficiency of the active site, we replaced the entire yeast aminoacylation domain with those from either human or P. falciparum (Figure 5A). These strains were engineered to be dependent on the hybrid synthetases, and all showed normal growth (Figure 5B). However, the S. cerevisiae strain bearing the P. falciparum hybrid synthetase showed a 1,000-fold increase in sensitivity to cladosporin (IC50 from >200 μM to 0.17 μM, Figure 5C). In contrast, the strains bearing the human or yeast synthetases showed little change in cladosporin sensitivity (Figure 5C). These data demonstrate that the conformation of the aminoacylation domain renders
the malaria parasite protein uniquely susceptible to cladosporin and that lysyl-tRNA synthetase is the primary target of cladosporin.

**DISCUSSION**

Cell-based screening has recently been shown to be an attractive way to find new leads for malaria drug development (Gamo et al., 2010; Guiguemde et al., 2010; Plouffe et al., 2008). In this respect, cladosporin could be an attractive lead, given the species selectivity and potent nanomolar activity against liver-stage parasites. Unfortunately, cladosporin possesses poor oral bioavailability, a key requirement for an anti-malarial. Therein lies a key disadvantage of the cell-based lead-finding approach—if an initial hit is chemically intractable and the target is not known, there may be no path forward. However, if a chemically validated target can be discovered, then a more focused and extensive lead-finding biochemical screen can be readily initiated.

Methodological choices for finding targets in malaria parasites have been traditionally limited. Cheminformatic predictions based on historical screening data against different biochemical targets (Gamo et al., 2010) and/or in silico docking typically only creates hypotheses, and extensive, time-consuming experimentation is required for confirmation. Affinity chromatography, a method by which an inhibitor is immobilized to resin and incubated with cell lysate to recover the molecular target, has been used with limited success (Kato et al., 2008; Knockaert et al., 2000). However, nonspecific binding, variable parasite protein target abundance (1,000-fold differences in protein abundance have been observed), and limitations in generating enough parasite lysate for analysis make this a challenging approach. Thus, evolution of drug-resistant parasites combined with genome scanning (Dharia et al., 2009) is our preferred method for discovering the target of antimalarials (McNamara and Winzeler, 2011; Meister et al., 2011; Nam et al., 2011; Rottmann et al., 2010). Nevertheless, some ambiguity often remains. Specifically, more than one gene may be mutated, or in some cases, resistance is conferred by a copy number variant encompassing several genes. The use of yeast deletion strains to complement the parasite evolution studies provides a powerful method of confirmation. However, an inherent disadvantage of genetic methods in both yeast and parasites is that only a resistance mechanism may be revealed, and thus additional experiments will always be necessary to validate the mechanism of action.

While it is possible that the amplification in lysyl-tRNA synthetase only confers resistance to cladosporin, or that cladosporin affects its killing through a different mechanism than by inhibiting lysyl-tRNA synthetase in *P. falciparum*, our biochemical evidence suggests that lysyl-tRNA synthetase is the primary target of cladosporin. Recent evidence suggests that human Krs1 has diverse biological roles beyond protein synthesis. Among these functions, it is secreted as a proinflammatory cytokine (Park et al., 2005) and is an integral component of a multimeric protein complex that affects regulatory signal pathways (reviewed in Park et al., 2010). The large therapeutic window between species makes inhibition of human Krs1 a minor concern. However, it is not known whether these same diverse functions are conserved in *Plasmodium* spp. Therefore, it is presently unclear if specifically targeting PfKrs1 and/or potential components in these putative pathways makes this enzyme a more desirable parasite target than other AARSs.

Aside from the noncanonical functions, selective inhibition of the protein translational function of PfKrs1 has obvious value because of the significant parasite proliferation during both liver- and blood-stage infections. Indeed, AARSs have recently gained recognition as desirable targets in antibacterial drug discovery programs (Hurdle et al., 2005). For instance, mupirocin has been shown to be a selective inhibitor of bacterial isoleucyl-tRNA synthetase (Hughes and Mellows, 1980; Sheng and Zhang, 2011), and a series of diaryl diamines specifically target *Trypanosoma brucei* methionyl-tRNA synthetase (Shibata et al., 2011). Also, a broad-spectrum inhibitor of fungal leucyl-tRNA synthetases, AN2690, is in development for the treatment of onychomycosis (Rock et al., 2007). However, targeting the AARS of an invasive micro-organism without inhibiting the human counterpart is a major challenge in drug discovery. Unlike the highly divergent prokaryotes, eukaryotic pathogens such as fungus (*Candida* spp.) and apicomplexan parasites (*Plasmodium*, *Toxoplasma*, and *Trypanosoma* spp.) generally have higher conservation with the homologous human AARS. Moreover, the lack of three-dimensional structures from some eukaryotic pathogens limits rational structure-based drug design to help identify and exploit differences. Despite this, we show here that selective eukaryotic AARS inhibitors can be achieved, opening the door for target-based drug discovery efforts focused on lysyl-tRNA synthetase. This in turn may lead to small molecules with the

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**Table 2. Summary of Amino Acid Conservation at Key Residues in the ATP Pocket of Krs1 and Corresponding IC₅₀ Value of Cladosporin**

| Organism           | Cladosporin Inhibition | Key Active Site Residues Position | Position |
|--------------------|------------------------|----------------------------------|----------|
| *Plasmodium falciparum* | 0.04–0.08 IC₅₀ (µM)  | Val 324 Ser 340                  |          |
| *Plasmodium yoelii*  | 0.04 IC₅₀ (µM)         | Val 324 Ser 340                  |          |
| *Trypanosoma brucei* | 2.05                  | Val 324 Thr 340                  |          |
| *Leishmania donovani* | 2.56                  | Val 324 Thr 340                  |          |
| *Toxoplasma gondii*   | 2.63                  | Asn 324 Ala 340                  |          |
| *Homo sapiens*       | >10                    | Gln 324 Thr 340                  |          |
| *Saccharomyces cerevisiae* | 30–110                  | Gln 324 Thr 340                  |          |
| *Escherichia coli*    | >100                   | Asn 324 Met 340                  |          |
| *Bacillus stearothermophilus* | >100                  | Val 324 Met 340                  |          |

*IC₅₀ (µM) and MICb (µg/ml) were determined on primary sequence alignments (see also Figure S4). The SYBR Green proliferation assay of blood-stage parasites and high-content imaging of liver-stage parasites are methods used to confirm the targets.*

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Figure 5. Yeast Cells Dependent on Chimeric Lysyl-tRNA Synthetases Exhibit Wild-Type Growth Kinetics but Show Differential Sensitivity to Cladosporin
(A) The homology model of ScKrs1 with cladosporin docked into the ATP binding site (asterisk). Amino acids 1–220 corresponding to the tRNA-binding domain are colored black, whereas residues beyond 220, which correspond to the aminoacylation domain, are shown in color.

(B) Growth curves of yeast cells solely dependent on heterologous, chimeric lysyl-tRNA synthetases with aminoacylation domains of the indicated species. Data for (B) and (C) were collected using the optical density assay and are a representation of triplicate experiments. The independent data points for each experiment are shown. In (C), a nonlinear regression curve fit was performed on the means ± SD of these data.

EXPERIMENTAL PROCEDURES

HIP Profiling
Cladosporin was tested in three independent HIP experiments at 110 μM (two times) and 150 μM (one time). Each experiment identified the krs1/KRS1 strain as the most sensitive strain. The experimental execution and processing of results were based on previous protocols (Pierce et al., 2007). The HIP assay was performed in 24-well plates with 1,600 μl of YPD per well. Per experiment, cladosporin was tested in two wells and compared to eight wells with DMSO. At the beginning of each experiment, 100 μl per well of a 1.5 OD600/ml log phase pre-culture comprising all strains of the heterozygous deletion collection strains were added (OpenBiosystems, part numbers YSC1056). Plates were incubated for 16 hr in a robotic shaking incubator at 30 °C/55 RPM, allowing for approximately five doublings and then a robotic system back-diluted 50 μl/well into fresh medium with compound. This procedure was repeated two more times until the yeast had doubled 20 times. The HIP pool construction, gDNA extraction, hybridization procedure, and calculation of sensitivity scores were based on the methods published by Pierce et al. (Pierce et al., 2007). In addition to the sensitivity score, for each gene, a z score was calculated from the TAG4 microarray intensities by taking the logarithm of the ratio of mean experiment versus mean control tag-averaged intensities. The z scores of all genes were then robustly normalized experiment- and gene-wise.

S. cerevisiae Aminoacyl-tRNA Synthetase Overexpression
Lysyl-tRNA synthetase (KRS1), isoleucyl-tRNA synthetase (IIS1), glutamyl-tRNA synthetase (GLN4), and threonyl-tRNA synthetase (THS1) were obtained from the previously characterized yeast GAL overexpression collection (Gelperin et al., 2005) (OpenBiosystems, part numbers YSC870). Cells were grown in synthetic medium supplemented with 2% galactose and serial dilutions of cladosporin (200 μM max concentration, 3:1 dilution). DMSO was normalized to 2%. Dose-response curves were calculated by taking the 20 hr OD600 measurements and applying a logistic regression curve fit.

Selection of Drug-Resistant S. cerevisiae
Strain BY4743Δ8, derived from BY4741 but deleted for eight genes involved in drug resistance (efflux pumps, SNQ2, PDR5, YOR1; transcription factors, PDR1, PDR2, PDR3, YAP1, YRM1) was incubated in 2.5% ethanolamine sulfate until only 50% of the cells formed colonies. A total of 2 × 10^7 mutagenized CMB970 cells were plated on two 14 cm dishes with synthetic complete medium (0.7 g/Difco Yeast Nitrogen Base without amino acids, 0.79 g/Mpbio CSM amino acid mixture, 2% glucose) containing 50 μM cladosporin. After 4 days, 37 resistant colonies appeared. Resistance was confirmed by re-streaking on 50 μM cladosporin. Genomic DNA was extracted (YeaStar Genomic DNA Extraction Kit, Zymo Research), and the KRS1 gene including promoter and terminator was sequenced (P1, CCTTACCTGAGGAAAGG; P2, GGTCAAGGAGATCCTGG; P3, CATCCTATTCCCTCCT; P4, CTGCA CACCAACGATATC). Of these 37, ten colonies contained nonsynonymous mutations in the KRS1 open reading frame. To distinguish between recessive and dominant mutations, the ten strains were crossed with an isogenic wild-type strain of opposite mating type and resistance confirmed by restreaking on 50 μM cladosporin. All ten strains resulted in resistant diploids. To exclude secondary mutations, KRS1 genes bearing three different mutations were cloned into the Smal and Kpn1 sites of the single copy pRS416 plasmid by PCR amplification with primers, containing restriction sites at their 5’ ends (KRS1-Smal, ATAAATCCCGGACCCATGAGCACAATAG; and KRS1-Kpn1, ATAAATGCTAGCGGTAGCAATAG) and transformed into cladosporin-sensitive BY4741 yeast. Resistance was confirmed by recording dose-response curves in YPD medium with serial dilutions of cladosporin (200 μM max concentration, 3:1 dilution). DMSO was normalized to 2%. Curves were calculated by taking the 16 hr OD600 measurements and applying a logistic regression curve fit.

Generation and Assay of Mutant KRS Lines in Yeast
S. cerevisiae KRS1 wild-type gene was cloned into a pRS416 low copy number plasmid. Single mutations were introduced by site-directed mutagenesis, and the double mutant was generated by excising the 623 bp fragment containing the Gin324Val mutation and cloning it between the EcoRI sites of the Thr340Ser-containing plasmid. The resulting plasmids were verified by direct DNA sequencing, transformed into KRS1/krts heterozygous deletion mutants, sporulated, and the tetrads dissected. Selecting for uracil auxotrophy and G418 resistance identified colonies exclusively dependent on the plasmid encoding mutant KRS1. IC50 values were determined by testing growth of four independent colonies in SD-ura medium with 12-point serial dilutions at a dilution factor of 3. The top concentration of cladosporin was 200 μM and resulted in a final DMSO concentration of 2%. Growth was monitored by reading OD600 values over 24 hr, and these data were fit to a nonlinear regression by Prism to calculate the IC50 value.

Selection of Drug-Resistant Malaria Parasites
Cladosporin-resistant parasites were selected using the protocol described in Rottmann et al. (Rottmann et al., 2010). In brief, clonal Dd2 parasites were established in triplicate flasks and evolved independently. Drug challenge was initiated at the IC50 value for cladosporin (40 nM) and increased in 10–40 nM steps. The continuous exposure and challenge to sublethal concentrations of inhibitor were carried out for 2 months, with final drug concentration between 380 and 400 nM. Clonal cladosporin-resistant clones were selected by limiting dilution using complete media supplemented with 340 nM cladosporin. Each clone was tested against a panel of antimalarial compounds (Table S4).

pharmacokinetic and pharmacodynamic properties needed for the orally bioavailable antimalarial drugs needed to support global eradication efforts.
Cladosporin Selectively Targets *P. falciparum* Krs1

**Genome Scanning of *P. falciparum* gDNA**
Genomic DNA samples, extracted from cladosporin-sensitive and cladosporin-resistant clones, were digested, labeled, and analyzed as previously described by Dharia et al. (Dharia et al., 2009). Briefly, 15 μg of digested biotin end-labeled genomic DNA was hybridized to a custom high-density tiling array (Pttling). PttGenominator version 2.0 (freely available at http://www.scripps.edu/winzeler/software/) was used to perform a comparative genomic analysis to detect hybridization differences between the parental Dd2 clone and each of the independently derived cladosporin-resistant clones. Modular analyses were performed to identify copy number variation and single nucleotide polymorphism (SNP) events. A p value of cutoff <1 x 10^-8 was enforced on the SNP analysis to filter out false positives, and a probe-by-probe analysis was performed on each SNP to validate its likelihood.

**Metabolic Labeling with [35S]-Methionine/Cysteine for Protein Synthesis Inhibition Studies**
A metabolic labeling assay was performed according to the protocol described in Rottmann et al. (Rottmann et al., 2010). For more information, refer to the Supplemental Experimental Procedures.

**Molecular Modeling**
Structural alignment of the X-ray crystal structures of human (Protein Data Bank [PDB] ID 3BUJ) and bacterial lysyl-tRNA synthetases (PDB ID 1BBU and 3E9H) using ICM software (Molsoft, LLC) revealed a high degree of similarity (see Figure S4). The yeast homology model was built using the MOE software (Chemical Computing Group). The human lysyl-tRNA synthetase X-ray structure showed the greatest amino acid conservation to the yeast sequence and was therefore selected as the template for the yeast homology model. The homology model was built using MOE default parameters. Prior to docking studies with cladosporin, a model for the binding of ATP in the yeast protein was created. ATP was manually placed within the yeast homology model, and the protein side chains were minimized in order to best accommodate this compound. Cladosporin was docked in this protein using Glide SP (Schrodinger, LLC) and refined using the MM/GBSA method implemented in Prime (Schrodinger, LLC).

**Cloning and Expression of *Plasmodium* Lysyl-tRNA Synthetase**
The sequence of the lysyl-tRNA synthetase (PF13_0262) was optimized for E. coli codon usage and synthesized in pUCminus vector (1st Base). The gene was subcloned into pET-28a(+) expression vector (Novagen) using BamHI and EcoRI sites, and transformed into *E. coli* Tuner (DE3) cells (Novagen). Of 2xYT media, 2 L containing 50 μg/ml kanamycin was inoculated with 20 μl of overnight culture and grown at 37°C. Expression was induced with 0.1 mM IPTG at A600 = 0.8 and incubated for 24 hr at 13°C. The cells were harvested by centrifugation at 6,000 rpm for 10 min and frozen at –80°C.

**Purification of *Plasmodium* Lysyl-tRNA Synthetase**
Frozen cell pellets were thawed and resuspended in 60 ml cold lysis buffer (20 mM Tris HCl [pH 8.0], 500 mM NaCl, 0.5 mM TCEP) supplemented with 0.01% SDS and a protease inhibitor tablet (Roche Diagnostics). Cells were lysed by sonication, and the crude lysate was centrifuged (22,000 g, 1 hr at 4°C). The supernatant was clarified through a 0.22 μm filter (Millipore) and loaded onto a Ni-affinity column. The column was washed with 10 column volumes wash buffer (20 mM Tris [pH 8.0], 500 mM NaCl, 50 mM imidazole, and 0.5 mM TCEP) and the protein eluted with a gradient of 50–500 mM imidazole over 20 column volumes. The protein-containing fractions were dialyzed against low-salt buffer (20 mM Tris [pH 8.0], 100 mM NaCl, and 0.5 mM TCEP) and load to a MonoQ 5/50GL column (GE Healthcare). The column was washed with 4 column volumes of wash buffer 2 (20 mM Tris [pH 8.0], 150 mM NaCl, 0.5 mM TCEP), a shallow gradient elution was carried out with 150–1,000 mM NaCl over 50 column volumes. A final purification step was repeated with the Ni-affinity column. The concentration of the eluent-containing target protein was measured by absorbance at 280 nm with ε = 41,780 M^-1 cm^-1.

**Lysyl-tRNA Synthetase Activity Assay**
Purified recombinant human enzyme was purchased from OriGene (catalog number TP300311). Enzymatic aminoclaylation by lysyl-tRNA synthetase was assayed with the Transcreener AMP assay system (BellBrook) in a Corning 384-well black plate. The P7RNA<sup>75</sup> substrate was chemically synthesized (Trilink). The assay was performed in a 10 μl volume containing either 250 nM of Plasmodium purified enzyme or 30 nM of human lysyl-tRNA synthetase (OriGene) in 40 mM HEPEs (pH 7.5), 1 mM DTT, 40 μM ATP, 10 μM P7RNA<sup>75</sup>, 8 mM MgCl₂, and 40 μM L-lysine. Reagent mixing was performed on orbital shaker for one minute at 900 rpm, followed by incubation at RT for 4 hr. Enzymatic activity was terminated by the addition of an equal volume (10 μl) of detection reagent and yielded a final concentration of 0.5x stop buffer, 1 mM AMP-Alexa633 tracer, and 2.5 μg/ml ADP antibody. Fluorescence polarization measurements were performed on the Infinite M1000 plate reader (Tecan) using a 635 nm excitation and 680 nm emission (20 nm bandwidth) filter settings. IC<sub>50</sub> values were determined by subtracting the background activity and performing nonlinear regression with Prism (GraphPad).

**Construction of Hybrid Synthetase Yeast Strains**
Protein sequences of *Homo sapiens* (NM_001130089) and *P. falciparum* lysyl-tRNA synthetase (PF13_0262) were aligned with *S. cerevisiae* Krs1 sequence. The first 220 ScKrs1 amino acids where taken (tRNA binding domain) and then fused to the aminoacylation domains for the two respective sequences. These hybrid protein sequences were back-translated into DNA using yeast codon usage, and synthetic DNA fragments were created with a Smal site at the 3' end. 332 bp of *S. cerevisiae* KRS1 promoter sequence, the hybrid KRS1 sequence, 206 bp of *S. cerevisiae* KRS1 terminator sequence, and a KpnI site. As a control, the same construct was generated using the full-length *S. cerevisiae* KRS1 sequence. The DNA fragments where then cloned into the cloning site of pBYIntURA plasmids using the Smal and KpnI sites, linearized with BstX1, and transformed into the heterozygous BY4743 KRS1/krsp1::kanMX4 (ura3J) strain where the constructs integrated into the TRP1 locus by selecting on medium lacking uracil. Transformants were verified by analytical PCR, sporulated, and dissected. Germinating spores were further analyzed for the integration of the plasmid in the TRP1 locus and presence of the KanMX deletion by analytical PCR. Integrity of the integrated lysyl-tRNA synthetase constructs was verified by sequencing. Growth kinetics of the strains were assessed by measuring OD<sub>600</sub> values in hourly intervals on a robotic system in 96-well plates. Sensitivity against cladosporin was tested by growing the stains in serially dilutedcompound as described previously.

Additional methods are provided in the Supplemental Information.

**SUPPLEMENTAL INFORMATION**
Supplemental Information includes four figures, five tables, Supplemental Experimental Procedures, and Supplemental References and can be found with this article online at doi:10.1016/j.chom.2012.04.015.

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