Anti-malarial benzoxaboroles target *P. falciparum* leucyl-tRNA synthetase


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Running title: LeuRS is a target of antimalarial oxaboroles

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**ABSTRACT**
There is a need for new antimalarials, ideally with novel mechanisms of action. Benzoxaboroles have been shown to be active against bacteria, fungi, and trypanosomes. Therefore, we investigated the antimalarial activity and mechanism of action of 3-aminomethyl benzoxaboroles against *Plasmodium falciparum*. Two 3-aminomethyl compounds, AN6426 and AN8432, demonstrated good potency against cultured multidrug-resistant (W2 strain) *P. falciparum* (IC₅₀ 310 nM and 490 nM, respectively) and efficacy against murine *Plasmodium berghei* infection when administered orally once daily for 4 days (ED₉₀ 7.4 and 16.2 mg/kg, respectively). To characterize mechanisms of action, we selected parasites with decreased drug sensitivity by culturing with step-wise increases in concentration of AN6426. Resistant clones were characterized by whole genome sequencing. Three generations of resistant parasites had polymorphisms in the predicted editing domain of the gene encoding a *P. falciparum* leucyl-tRNA synthetase (LeuRS; PF3D7_0622800) and in another gene (PF3D7_1218100), which encodes a protein of unknown function. Solution of the structure of the *P. falciparum* LeuRS editing domain suggested key roles for mutated residues in LeuRS editing. Short incubations with AN6426 and AN8432, unlike artemisinin, caused dose-dependent inhibition of [¹⁴C]leucine incorporation by cultured wild type, but not resistant parasites. The growth of resistant, but not wild type parasites was impaired in the presence of the unnatural amino acid norvaline, consistent with a loss of LeuRS editing activity in resistant parasites. In summary, the benzoxaboroles AN6426 and AN8432 offer effective antimalarial activity, and act, at least in part, against a novel target, the editing domain of *P. falciparum* LeuRS.
There is an urgent need for new antimalarials. As older regimens are limited by resistance in *Plasmodium falciparum*, the most virulent human malaria parasite, artemisinin-based combination therapy regimens have been adopted as standard treatment for uncomplicated falciparum malaria (1). However, resistance to artemisinins is growing in Southeast Asia (2), and resistance has been seen to most artemisinin partner drugs (3). Drug resistance seriously jeopardizes efforts to control and eliminate malaria, and new agents, ideally with novel mechanisms of action, are needed.

Benzoxaboroles are boron-containing compounds that have demonstrated potent activity against a number of infectious pathogens, including bacteria (4, 5), fungi (6), and trypanosomes (7 an orally-active benzoxaborole for the treatment of stage 2 human African trypanosomiasis). The highly electrophilic nature of the boron component of these compounds leads to interaction with a variety of protein targets via reversible covalent bonds (8, 9), with identified targets including leucyl-tRNA synthetase (LeuRS) (5, 6) and β-lactamase (10). In particular, the benzoxaboroles tavaborole and AN3018 (6) inhibit fungal LeuRS, and AN3365 and AN3664/ZCL039 inhibit bacterial LeuRS (4, 5).

There is increasing interest in the exploration of aminoacyl-tRNA synthetases as antimicrobial targets in both prokaryotic and eukaryotic pathogens (11, 12). These enzymes catalyze the attachment (charging) of amino acids to their cognate tRNAs. For example, mupirocin inhibits bacterial IleRS and is used topically to control *Staphylococcus aureus* colonization (13 special features and applications of an antibiotic from a gram-negative bacterium). Class I aminoacyl-tRNA synthetases, including LeuRS, contain a cis editing domain that is distinct from the enzyme active site, and that recognizes non-cognate amino acids and hydrolyzes misacylated tRNAs (14) The LeuRS editing domain is the target of the benzoxaborole LeuRS inhibitors noted above. In *P. falciparum*, aminoacyl-tRNA synthetase inhibitors and their targets include borrelidin (ThrRS) (15, 16 borrelidin, against drug-resistant strains of Plasmodia, 17 synthesis and biological evaluation against *Plasmodium falciparum*
parasites), mupirocin (IleRS) (44), 4-thiaisoleucine (IleRS) (18), cladosporin (LysRS) (19), lysyl-
adenylate analogues (LysRS) (20), and halofuginone (ProRS) (21). The *P. falciparum* cytosolic
LeuRS, but not an apicoplast-directed LeuRS, contains a typical editing domain (22).

In a search for new antimalarial compounds we screened a benzoxaborole library rich in
LeuRS inhibitors for potency against cultured *P. falciparum*. The two most active compounds,
AN6426 and AN8432, were selected for further studies. The compounds demonstrated effective
activity against multiple *P. falciparum* strains and in a murine malaria model. Genetic and
biochemical evidence are consistent with activity of these compounds against *P. falciparum*
LeuRS.

**MATERIALS AND METHODS**

**Synthesis of benzoxaboroles.** Experimental procedures and analytical data for compounds 1,
2, 5, 6, 9 and 10 (Table 1) were described previously (23). Their analogs 3, 4, 7, and 8 were
prepared following similar synthetic routes and procedures. The compounds were analyzed by
1H NMR, LC-MS and HPLC for structural characterization and purity. 1H NMR spectra were
recorded on a Varian or Bruker spectrometer (300 or 400 MHz). LC-MS data were obtained
using an Agilent LC-MS 1200 with 6110 MS detector equipped with an electrospray ion source
operated in a positive or negative mode. HPLC analysis was performed on a Shimadzu HPLC
system or a Waters 600 Controller system. The columns used were a Venusil XBP-C18 (50 ×
4.6 mm I.D.), Shimpack VP-ODS (150 × 4.6 mm I.D.), or UPLC BEH C18 (50 × 2.1 mm I.D.).

Resulting analytical data were consistent with corresponding structures, and HPLC purity was
≥95% at 214, 220, or 254 nm (Supporting Information).
Culture of malaria parasites. Erythrocytic *P. falciparum* were cultured using standard methods at 2% hematocrit in RPMI-1640 (Invitrogen) medium supplemented with 0.5% Albumax II (GIBCO Life Technologies), 2 mM L-glutamine, 100 mM hypoxanthine, 5 μg/ml gentamicin, 28 mM NaHCO₃ and 25 mM HEPES at 37°C in an atmosphere of 5% O₂, 5% CO₂, and 90% N₂. *P. falciparum* strains were from the Malaria Research and Reference Reagent Resource Center (www.mr4.org).

Activity of benzoxaboroles against cultured *P. falciparum*. Parasites were synchronized by treatment with 5% D-sorbitol and cultured in duplicate 96 well culture plates (200 μL per well) with serially diluted benzoxaboroles or the antimalarials chloroquine, lumefantrine, mefloquine, piperaquine, artemisinin, or dihydroartemisinin (from Sigma-Aldrich, except piperaquine was from Jinan Jiaquan International Trade Co.), at concentrations of 0.056-1000 nM, with ≤0.2% DMSO. After 48 h, cultures were fixed with 2% formaldehyde for 24 h at 37°C or 48 h at room temperature, cells were stained with 4 nM YOYO-1 dye (Molecular Probes), and counts of treated and control cultures were determined using FACS. IC₅₀ values were calculated by nonlinear regression using GraphPad Prism software. Activity against isolates from Ugandan children with uncomplicated falciparum malaria was determined using an ELISA directed against *P. falciparum* histidine-rich protein-2, as previously described (24).

Cytotoxicity Assays. Human Jurkat (T cell lymphoma) and HepG2 (hepatocarcinoma) cells were from the American Type Culture Collection. Jurkat cells were seeded in 96-well plates at 2 x 10⁴ cells per well in 100 μL RPMI-1640 medium with 10% fetal bovine serum and 2 mM L-glutamine, with 10-fold serial dilutions (0.1 nM-100 μM) of oxaboroles, and a final concentration of 0.25% DMSO. Plates were incubated at 37°C in 5% CO₂ for 72 h, 20 μL [3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H-tetrazolium was added, and after 4 additional h, absorbance was determined at 490 and 690 nm. IC₅₀ values were
calculated by non-linear regression analysis with the four parameter logistic equation using GraphPad Prism software. HepG2 cells were seeded in 96-well plates at 3000 cells per well in Dulbecco's Modified Eagle's medium with 10% fetal bovine serum, 2 mM L-Glutamine, and 50 units/mL penicillin-streptomycin. Serial half-log dilutions of oxaboroles in DMSO, at a final concentration of 0.5% DMSO, were added and cells grown for 7 days, at 37°C in 5% CO2 with replacement of medium and compounds on the fourth day. After 7 days, media was removed, cells were blotted dry, 50 uL of 1X Janus Green Stain (Mitosciences, Eugene, OR) was added per well, and the assay was developed according to the manufacturer's protocol (In-cell ELISA kit, cat # MS643). Absorbance at 595 nm was measured and IC50 values were calculated as described for Jurkat cells.

Activity of benzoxaboroles against murine malaria. Swiss Webster mice were infected intraperitoneally with 6x10^6 P. berghei-infected erythrocytes collected from a previously-infected mouse and then treated, beginning 1 h after inoculation, with benzoxaboroles (in 55% polyethylene glycol 300, 25% propylene glycol, 20% water) or chloroquine (in water) by daily oral gavage for 4 days. Negative controls were treated with vehicle only. Infections were monitored by daily microscopic evaluation of Giemsa-stained blood smears. ED90 values, based on comparisons of parasitemias between treated and control animals on the fourth day after initiation of treatment, were calculated using Graphpad Prism software. Mice were euthanized when parasitemias exceeded 50%.

Stage specificity assay. Using a previously-described protocol (25), synchronous W2 strain P. falciparum were cultured in triplicate wells in 96-well culture plates with 2 μM AN6426 or 1.3 μM chloroquine for 8 h intervals, beginning at the ring stage. At the end of each interval, the cultures were washed 3 times and resuspended in culture media without drug. After 48 h, when control
parasites were at the ring stage, the cultures were fixed with 2% formaldehyde, stained, and
counted with FACS as described above.

Selection of parasites with decreased sensitivity to AN6426. Triplicate 10 mL cultures of
Dd2-strain *P. falciparum*, each containing a clonal population of $6 \times 10^7$ asynchronous parasites,
were incubated with stepwise increasing concentrations of AN6426, beginning with 0.4 μM.
Media was changed and fresh AN6426 added daily. Once treated parasites grew at rates
comparable to those of untreated controls, parasite aliquots were cloned by limiting dilution,
drug sensitivity (IC₅₀ values) was assessed as described above, aliquots were stored, the
concentration of AN6426 was increased, and the selection process was repeated. To assess
the ease of resistance selection, 3 cultures each of $10^6$, $10^7$, and $10^8$ Dd2 strain parasites were
incubated with 1.5 μM AN6426, and cultures were followed for 67 days or until parasites regrew.

Whole Genome Sequencing. To prepare genomic DNA, synchronized *P. falciparum*-infected
erthrocytes (100 mL, 2% hematocrit, 10% parasitemia) were treated with 0.15% saponin for 5
min on ice to lyse erythrocytes followed by 3 washes in PBS. Parasite pellets were lysed in 150
mM NaCl, 10 mM EDTA, 50 mM Tris-HCl pH 7.5, 0.1% sarkosyl (Sigma Aldrich), and 200
mg/mL proteinase K (Qiagen) overnight at 37°C. The samples were then subjected to extraction
with phenol/chloroform/isoamyl alcohol (25:24:1), pH 7.9 (Ambion), treatment with 0.05mg/mL
RNase A (1 h 37°C), two additional phenol/chloroform extractions, one chloroform extraction,
and then ethanol precipitation. All extractions used light phase lock tubes (5 Prime).

Genomic DNA libraries were prepared from 100 ng DNA using the Nextera DNA Sample
Prep Kit (Illumina) according to the manufacturer’s instructions, except with 6 cycles and the
bridge amplification step at 65°C for 6 min (26 Plasmodium falciparum, use a two-step genomic
strategy to acquire accurate, beneficial DNA amplifications). Each library was barcoded with
unique sets of two indices from the Nextera Index Kit (Illumina) to allow multiple samples to be
run on one flow cell. Next, fragments of 360-560 bp were extracted and collected using Lab
Chip XT (Caliper Life Sciences) according to the manufacturer’s instructions. The fragments
were amplified by limited-cycle PCR using Kapa HiFi DNA polymerase (Kapa Biosystems) with
dNTPs with an 80% AT coding bias (6 cycles of 95°C for 10 sec, 58°C for 30 sec, 65°C for 6
min). The primers for both PCR steps were AATGATACGGCGACCACCGA and
CAAGCAGAAGACGGCATACG (26 Plasmodium falciparum, use a two-step genomic strategy
to acquire accurate, beneficial DNA amplifications). Libraries were pooled at concentrations of
2 nM per library, determined with a DNA Bioanalyzer (Agilent). Final library quantification and
qualification, as detailed in (26 Plasmodium falciparum, use a two-step genomic strategy to
acquire accurate, beneficial DNA amplifications), were completed, followed by sequencing at
the UCSF Center for Advanced Technology on a HiSeq 2000 system (Illumina). Sequence data
for each library were aligned to the 3D7 reference genome using Bowtie (27), discarding reads
with >1 nucleotide mismatch and multiple alignments across the genome. For the identification
of SNPs, reads were matched to those from the parental strain, and the top 200 SNPs per
chromosome, ranked according to frequency of conflicting nucleotides per position in the
genome, were chosen and filtered based on standard parameters (26 Plasmodium falciparum,
use a two-step genomic strategy to acquire accurate, beneficial DNA amplifications). SNPs
were considered legitimate if the number of reads covering the position was >10 and the
frequency was at least 80%. Searches for novel SNPs included only non-synonymous SNPs in
exons, excluding hypervariable genes (pfemp1, rifin, and stevor). Copy number variation was
analyzed using the UCSC Genome Browser (28).

**Dideoxy Sequencing.** Genomic DNA was extracted using the QIAamp DNA mini kit (Qiagen)
according to the manufacturer’s instructions. PF3D7_0622800 was amplified in 4 fragments
using the Phusion Hot Start II High-Fidelity DNA Polymerase kit (Thermo Scientific) with 80%
AT dNTPs and primers as in Table S3 (95°C for 3 min; 30 cycles of 95°C for 10 sec, 52°C for 30
sec, 65°C for 1 min; 68°C for 10 min). The amplified fragments were cleaned using ExoSAP-IT (Affymetrix), mixed with sequencing primers (Supplementary Table 3), and sequenced at the UCSF Genome Core Facility. For PF3D7_1218100, the same approach was followed, but only one 800 bp fragment, including the M416T SNP, was amplified and sequenced.

Production of *P. falciparum* and *P. knowlesi* LeuRS editing domains. DNA fragments encoding LeuRS editing domains of *P. falciparum* (PfLeuRS; residues 272-687) and *P. knowlesi* (PkLeuRS; residues 266-600) were cloned into the NcoI-XhoI sites of pETM-11 (EMBL). Flexible loops of PfLeuRS (residues 328-361 and 475-519, corresponding to insertions 1 and 3 respectively) and PkLeuRS (residues 400-432) were deleted for crystallization. A glycine was introduced at position 475 of PfLeuRS. Cysteine 273 of PfLeuRS was mutated to serine to prevent formation of intermolecular disulfides bridges. Proteins were expressed in BL21-Codon+RIL *E. coli* at 18°C. The cells were lysed by sonication in 40-60 mL of lysis buffer (20 mM TrisHCl pH 8.0, 100 mM NaCl, 5 mM β-mercaptoethanol, 10 mM imidazole, and one tablet of cOmplete protease inhibitors (Roche). The soluble fraction was loaded onto a nickel column (Ni-NTA, Qiagen) and washed with 50 mL lysis buffer; 50 mL lysis buffer plus 1 M NaCl, and then 50mL lysis buffer. Proteins were eluted with 15 mL lysis buffer plus 400 mM imidazole. The His-tag of recombinant proteins was removed with TEV protease. Pure protein fractions were dialyzed in lysis buffer without imidazole and concentrated to 10-15 mg mL\(^{-1}\).

Crystallization of *P. falciparum* and *P. knowlesi* LeuRS editing domains. Crystallization was carried out at 20°C by the hanging drop vapor diffusion method. PfLeuRS D3 crystallized in two different space groups. Monoclinic crystals of PfLeuRS D3 were obtained by mixing 1 μL 10 mg/ml protein with 1 μL reservoir solution containing 0.2 M sodium nitrate, pH 6.8, 20% (w/v) PEG 3350, and 12% glycerol. Triclinic crystals of PfLeuRS D3 were obtained by mixing 1 μL 10
mg/ml protein with 1 μL reservoir solution containing 0.1 M HEPES, pH 7.5, 10% isopropanol, 20% (w/v) PEG 4000, and 5% glycerol. Crystals of PfLeuRS D13 were obtained by mixing 1 μL of 10 mg/mL protein with 1 μL reservoir solution containing 0.1 M HEPES, pH 7.5, 20% (w/v) PEG 10000, and 8% ethylene glycol. Crystals of PkLeuRS were obtained by mixing 1 μL of protein at 12 mg mL⁻¹ with 1 μL reservoir solution containing 0.05 M sodium cacodylate, pH 6.5, 0.2 M potassium chloride, 0.1 M magnesium acetate, 10% (w/v) PEG 8000, and 25% ethylene glycol.

**Structure determination and refinement.** All data collection and refinement statistics are shown in Table S2. Diffraction data sets were collected at the European Synchrotron Radiation Facility and integrated and scaled with the XDS suite (29) or with the HKL2000 program package (30). Further data analysis was performed with the CCP4 suite (31 The CCP4 suite: programs for protein crystallography). The structure of PkLeuRS D3 was initially solved by molecular replacement with PHASER (32) using the structure of the C. muris LeuRS editing domain (PDB 5FOM; Palencia et al., unpublished) as a model. All models were improved by manual adjustments with COOT (33) and refined using REFMAC5 (34). Structure quality was analyzed with the PDBe validation server (http://wwpdb-validation.wwpdb.org/validservice/), which showed all residues for the different models in allowed regions of the Ramachandran plot. Figures were drawn with PYMOL (http://www.pymol.org/). Docking of AN6426-Ade76 into the editing site of Plasmodial LeuRS was done by structural alignment of the PkLeuRS structure to the complex of CmLeuRS-AN6426-AMP (PDB: 5FOM). The root-mean square deviation calculated over backbone atoms of CmLeuRS and PkLeuRS was 1.27 Å.

**Protein Synthesis Assay.** Synchronous Dd2-strain trophozoites at 10-15% parasitemia were washed 3 times with leucine-free culture media and resuspended at 5% hematocrit in the same
medium. Triplicate 100 μL cultures were incubated with 0X, 1X, 10X, or 100X IC₅₀ concentration of test compounds for 15 min at 37°C, 0.5 μCi of [¹⁴C]leucine (324 mCi/mmol, Perkin Elmer) was added, cultures were incubated for two h at 37°C, erythrocytes were lysed with 0.15% saponin as described above, pellets were resuspended in 100 μL 0.02% sodium deoxycholate (Sigma Aldrich) to lyse the parasites, and 100 μL ice cold 10% (w/v) TCA was added to precipitate proteins. Precipitates were transferred to 0.45 μm nitrocellulose membrane filter plates (Millipore Multiscreen HTS, MSHAN4B50), washed 4 times with 400 μL ice cold 5% TCA, and air-dried. [¹⁴C]leucine incorporation was determined by adding 60 μL OptiPhase Supermix scintillation cocktail (PerkinElmer) and counting in a Wallac MicroBeta Trilux 1450 liquid scintillation counter.

**Norvaline Sensitivity Assay.** Synchronized Dd2-strain parasites were cultured at 1% parasitemia, beginning at the ring stage, in either leucine-free or complete culture media with varying concentrations of L-norvaline (Sigma-Aldrich) for 48 h, and the parasites were then fixed, stained, and counted as described above.

Atomic coordinates and structure factors have been deposited in the Protein Data Bank under the following accession codes: 5FO4 for the PfLeuRS D3 structure in P1 space group, 5FOC for PfLeuRS D3 in P2₁ space group, 5FOD for the PfLeuRS D13, and 5FOF for the PkLeuRS structure.

**RESULTS**

Structure-activity relationships for 3-aminomethyl benzoxaboroles active against *P. falciparum*. A benzoxaborole library was screened against cultured multidrug-resistant W2 strain *P. falciparum*. Multiple 3-aminomethyl benzoxaboroles were active, and IC₅₀ values were
determined for the closely related compounds 1-10 (Fig. 1). Compound 1, with a phenyl group at R4, had moderate activity; replacement with a chloro atom in compound 2 led to sub-micromolar activity. Larger and more lipophilic groups at R7, such as a 7-benzyloxy group in compound 3, had minimal effect on antimalarial activity. A free basic amino group at R3 was required for activity, as a Boc-blocked amino group abrogated activity in compound 5.

Replacement of the R7 ethoxy group (6, AN6426) with other alkoxy groups, such as methoxy in 7 and cyclopropoxy in 8, led to decreased activity. Replacement of the R4 Cl with F in 9 reduced activity; replacement of this Cl with Br (10, AN8432) had a minimal impact on activity.

From this analysis compounds AN6426 and AN8432 emerged as the most active antimalarials, and they were selected for further study.

In vitro and in vivo antimalarial activity of AN6426 and AN8432. We characterized activities of AN6426 and AN8432 against multiple strains of *P. falciparum*, including field isolates. Activities were similar against laboratory strains known to be sensitive (3D7) or resistant (W2, Dd2) to chloroquine and other antimalarials, and against fresh isolates collected from malaria patients in Uganda in 2012 (Table 1). Assessment of IC50s showed minimal differences after 48 or 96 hour incubations, and thus AN6426 and AN8432 lack the delayed death phenotype (with much greater activity after 96 hours) seen with tetracyclines and some other antimicrobials. The compounds were tested for cytotoxicity against human Jurkat and HepG2 cells; toxicity was seen only at concentrations at least 50-fold greater than concentrations toxic to cultured parasites.

When administered orally to *P. berghei*-infected mice once daily for 4 days, the compounds demonstrated activity, with Day 4 ED90 values of 7.4 mg/kg/day for AN6426 and 16.2 mg/kg/day for AN8432. At a dosage of 200 mg/kg/day the compounds afforded long-term cures in 40-60% of infected mice (Fig. S1).
Stage-specificity and morphological effects of AN6426. Synchronized W2-strain parasites were incubated with the compound or chloroquine for 8 h intervals across the erythrocytic life cycle. AN6426 was removed at 8 h time points, cultures were maintained until untreated control parasites had reached the ring stage, and the parasitemias of control and treated cultures were compared. For both AN6426 and chloroquine, inhibition of parasite development was seen across the life cycle, but activity was greatest against trophozoites (Fig. 2). AN6426-treated parasites had no obvious morphological abnormalities during the ring and early trophozoite stages, but the parasites treated during these stages were unable to develop into multinucleated schizonts or new ring stage parasites (Fig. 2).

Selection and analysis of parasites with decreased sensitivity to AN6426. To gain insight into the antimalarial mechanism of action of AN6426, we cultured Dd2 strain *P. falciparum* in step-wise increasing concentrations of the compound, assessed parasite sensitivity after each step, and characterized cloned parasites with decreased sensitivity by whole genome sequencing. Parasites were incubated with AN6426 at 0.4 μM for 4 weeks (R1), 1 μM for 10 weeks (R2), and then 10 μM for 14 weeks (R3) (Fig. 3). At each step of selection, parasites were initially undetectable on Giemsa-stained smears (for ~3 weeks for R1 and 7-8 weeks for R2 and R3) followed by regrowth, suggesting selection of mutations allowing growth under drug pressure. Parasites at each step of AN6426 resistance were similarly resistant to AN8432, indicative of a shared mechanism of resistance. In contrast, selected parasites did not demonstrate decreased sensitivity to the antimalarials chloroquine, lumefantrine, mefloquine, piperazine, or dihydroartemisinin (Fig. 3).

Parasites from each step of selection were cloned by limiting dilution, and DNA from multiple clones was analyzed by whole genome sequencing. Sequencing identified a number of genetic changes between parent and AN6426-resistant parasites, including SNPs and copy number variations. SNPs in only two genes were seen in multiple clones from all selection.
steps: PF3D7_0622800, which encodes cytoplasmic LeuRS (35), and PF3D7_1218100, which encodes a protein of unknown function classified as a membrane protein based on homology models (36). For the LeuRS gene, 4 different SNPs were identified in the predicted editing domain (14, 37, 38) (Fig. 3). SNPs identified in R2 and R3 parasites (T400I, V568L), but not those identified in R1 parasites (E628G, V630L), map to conserved editing active-site regions of LeuRS from other eukaryotic pathogens (Fig. 4). For PF3D7_1218100, a single SNP (M416T) was present in R1, R2, and R3 parasites (Fig. 3). In addition, 2 to 4 fold amplification of two clusters of genes on chromosome 12 was observed at all levels of resistance; one of these clusters includes PF3D7_1218100 (Table S1). R3 parasites were subsequently cultured without drug pressure for 5 months. These parasites showed partial reversion to drug sensitivity, with ~5-fold decreased sensitivity to AN6426 compared to wild type, but they retained mutations in both PF3D7_0622800 and PF3D7_1218100.

To assess ease of selection of resistance to AN6426, we cultured different numbers of Dd2 strain P. falciparum with 1.5 μM AN6426. In 3 cultures with inocula of 10^8 parasites, parasites regrew after 16, 23, and 23 days. With 3 cultures each of 10^7 or 10^6 parasites, no growth was seen over 67 days of observation. For parasites incubated with 5 nM atovaquone, all of 6 cultures with inocula of 10^6 parasites regrew over 24-50 days, 2 of 6 cultures with inocula of 10^7 parasites regrew after 45 days, and 1 of 6 cultures with inocula of 10^6 parasites regrew after 45 days.

**Structure of the LeuRS editing domain of P. falciparum and P. knowlesi.** To better understand AN6426-LeuRS interactions we obtained crystal structures of the editing domain from two plasmodial species. Based on the crystal structure of the LeuRS editing domain of the related apicomplexan parasite Cryptosporidium muris (Palencia et al., unpublished), we
designed several constructs of the LeuRS editing domains of *P. falciparum* (residues 272–687) and of *P. knowlesi* (residues 266–600), a monkey species that causes human infections in southeast Asia. One or more of three apicomplexan specific and presumed flexible insertions were deleted with the aim of promoting crystallization. Crystals were obtained for two *P. falciparum* LeuRS constructs, one with deletion of insertion 3 (residues 475-520; *Pf*LeuRS D3), and another with deletion of insertion 1 as well (residues 328-360; *Pf*LeuRS D13). These crystals diffracted to 1.5 and 1.7 Å, respectively (Table S2, Fig. S2 and S3). In addition, we obtained crystals of *Pf*LeuRS D3 in a second space group (*P*1), which diffracted to 1.85 Å. A construct with deletion of insertion 3 in the *P. knowlesi* LeuRS editing domain (*Pk*LeuRS D3) gave crystals diffracting to 2.4 Å (Table S2). All structures showed the canonical editing domain fold of eukaryotic LeuRS (39), but with unique partially ordered insertions similar to those observed in the LeuRS editing domain of *C. muris* (Palencia et al., unpublished). The role of these insertions, which are distant from the editing active site, is unknown.

The two mutations identified in R2 and R3 resistant parasites, T400I and V568L, map to the threonine-rich and AMP-binding loops of the *P. falciparum* LeuRS editing domain, respectively. Both loops are involved in binding and proof-reading of non-cognate amino acids in bacterial and fungal LeuRS homologs (40) and in binding of benzoxaboroles (6, 7 an orally-active benzoxaborole for the treatment of stage 2 human African trypanosomiasis, 23, 39). T400 in the *P. falciparum* LeuRS structure aligns with T252 of *E. coli* LeuRS; this residue is responsible for steric exclusion of cognate leucine from the editing site, thereby preventing hydrolysis of correctly charged Leu-\text{tRNA}^{\text{Leu}}$, but allowing hydrolysis of incorrectly incorporated amino acids (41). The substitution of threonine by the more bulky isoleucine would likely impact negatively on the predicted binding of AN6426, which extends into the non-cognate amino acid binding site. V568 structurally aligns to V335 of *E. coli* LeuRS, which establishes hydrophobic contacts with the terminal adenosine-ribose of the tRNA and is important for the stabilization of the substrate in the LeuRS editing site (40). The V568L substitution is predicted to destabilize or
impede the formation of the inhibition adduct (AN6426-AMP) in the editing site of *P. falciparum* LeuRS due to the larger size of leucine. The LeuRS mutations found in R1 resistant parasites, E628G and V630L, which are associated with low-level resistance, map outside the editing active site (Fig. 4).

**AN6426 and AN8432 inhibit protein synthesis.** To further consider *P. falciparum* LeuRS as the target for the 3-aminomethyl benzoxaboroles, we characterized the effects of AN6426 and AN8432 on uptake of leucine, an indicator of parasite protein synthesis. Treatment of wild type Dd2 strain parasites with 2 h pulses of AN6426 or AN8432 inhibited incorporation of $[^{14}C]$leucine, indicative of a block in protein synthesis, as also seen with the protein synthesis inhibitor cycloheximide, but not with artemisinin, a rapid-acting antimalarial that does not act against protein synthesis (Fig. 5A). In parasites with high level resistance to AN6426, the compound did not inhibit $[^{14}C]$leucine incorporation (Fig. 5B). To specifically assess LeuRS editing, we cultured parasites in leucine-free media in the presence of norvaline, a non-cognate analogue of leucine that is readily charged to tRNA by LeuRS enzymes (40) and then hydrolyzed from tRNA$^{Leu}$ by the editing active site (37). Misincorporation of norvaline into proteins is toxic (6). Norvaline was not toxic to wild type parasites, but markedly inhibited the growth of R2 and R3 parasites (Fig. 4C), an effect that was not seen when cultures contained leucine (Fig. 4D). These results support the hypothesis that AN6426 acts against *P. falciparum* through inhibition of LeuRS editing.

**DISCUSSION**

We identified two benzoxaboroles, AN6426 and AN8432, with nanomolar in vitro activity against *P. falciparum* and efficacy in a rodent malaria model. To gain insight into mechanisms of action, we selected and characterized parasites with markedly decreased sensitivities to AN6426. Resistant clones consistently showed SNPs predicted to be within the editing domain.
Biochemical studies showed that AN6426 and AN8432 inhibited protein synthesis in wild type, but not AN6426-resistant parasites, and that resistant parasites were inhibited by exogenous norvaline, consistent with a loss of LeuRS editing. Taken together, our data suggest that a target of the antimalarial benzoxaboroles AN6426 and AN8432 is \textit{P. falciparum} LeuRS.

The mutations seen in parasites selected for resistance to AN6426 were predicted, based on comparison with homologs from other organisms, to be located within the LeuRS editing domain (4-6, 42), and this conclusion was supported by determination of the structure of this domain in \textit{P. falciparum} LeuRS and by our biochemical studies. Brief incubations with AN6426 and AN8432, but not the rapid-acting antimalarial artemisinin, led to dose-dependent inhibition of parasite incorporation of leucine, consistent with a block in protein synthesis. Cycloheximide blocked leucine incorporation at concentrations closer to its antiparasitic IC\textsubscript{50}, presumably because its action against polypeptide elongation (43, 44) is more rapidly toxic to parasites than inhibition of aminoacylation. In leucine-free media, growth of AN6426-resistant, but not wild type parasites was inhibited by norvaline, consistent with a block in LeuRS editing in resistant parasites. Taken together, our genetic and biochemical results suggest that AN6426 and AN8432 act against \textit{P. falciparum} via inhibition of LeuRS.

In contrast to the case with \textit{C. muris}, we did not detect binding of the AN6426-AMP adduct into the isolated \textit{P. falciparum} LeuRS editing domain \textit{in vitro}. Although the \textit{P. falciparum} and \textit{C. muris} editing domain structures are similar (sequence similarity 38.4%, root mean squared deviation calculated over backbone atoms 1.3 Å), the \textit{P. falciparum} editing active site contains three idiosyncratic amino acid substitutions (also seen in plasmodial homologs) at usually conserved residues. These substitutions might explain differences in the ease of binding of AN6426 to the editing domain (Fig. 4, Fig. S3). In \textit{P. falciparum} LeuRS, K397 makes a unique hydrogen bond to Q649, but does not interact with nearby E399. In LeuRS of \textit{C. muris} and many other organisms these three residues are R397, E649, and E399, and the multivalent arginine forms a salt bridge to both glutamates. Another \textit{P. falciparum} specific residue, I563,
which in other species is a lysine, is predicted to lead to the loss of interaction with the
phosphate of the 3' terminal adenosine (Ade76) of tRNA^{Leu}; this interaction contributes to
trapping the tRNA^{Leu} in the editing site of LeuRS. These idiosyncratic differences make it
plausible that for \textit{P. falciparum}, unlike other LeuRS enzymes, high affinity adduct formation only
occurs in the context of the full length LeuRS and tRNA.

Of note, \textit{P. falciparum} encodes two LeuRS enzymes, the nuclear-encoded enzyme that
is expressed in the cytoplasm and that we have shown to be the target of AN6426, and a
second enzyme that is encoded by the apicoplast genome (PF3D7_0828200; previously
annotated as PF08_0011) (45). We did not see mutations in the apicoplast-encoded LeuRS
gene in parasites selected for resistance to AN6426. This result is consistent with the
observation that AN6426 targets the LeuRS editing domain and the understanding from
informatic analysis that, unlike the cytoplasmic LeuRS, the \textit{P. falciparum} apicoplast LeuRS does
not contain a canonical editing domain (46).

Parasites selected for resistance to AN6426 consistently contained a SNP in a second
\textit{P. falciparum} gene, PF3D7_1218100, and 2-4 fold amplification of a gene cluster that includes
this gene. The function of the PF3D7_1218100 product is unknown. The product is predicted to
be a 55.7 kDa protein that features two transmembrane domains and an apicoplast signal
peptide (36). The gene is transcribed in mature erythrocytic parasites and in sexual stages.
Disruption of the \textit{P. berghei} ortholog of PF3D7_1218100 (PBANKA_143370) led to a defect in
the development of sporozoites, but no apparent effect in erythrocytic parasites (35). Thus,
limited available data concerning this gene product do not explain its role in resistance to
AN6426 and AN8432. Further consideration of both LeuRS and the product of PF3D7_1218100
as targets of AN6426 should include reverse genetic experiments to explore the impact of gene
alterations on sensitivity to the compound.

Malaria drug discovery is especially challenging. In addition to obvious requirements for
a drug to be safe and efficacious, antimalarial drug candidates should meet additional criteria,
including rapid clinical response, requirement for no more than 3 days of treatment (and ideally single-dose treatment), oral bioavailability, low tendency to select for drug resistance, lack of cross resistance with existing antimalarials, safety in children and in pregnancy, and low cost of production (47). Our results with AN6426 and AN8432, suggest that the 3-aminomethyloxaborole class warrants further exploration. First, the compounds exerted good activity against \textit{P. falciparum}, although increased potency is a goal in optimization of this class. Second, the compounds were active against murine malaria with once daily dosing. Third, safety of other benzoxaboroles has been demonstrated, with extensive \textit{in vitro} and \textit{in vivo} toxicology studies and progression to trials in humans (22, 48, 49). Fourth, AN6426 and related benzoxaboroles are easy to synthesize, requiring a 4 to 6 step scheme starting from inexpensive reagents (50). Lastly, our results suggest that AN6426 has a novel antimalarial mechanism of action, the inhibition of \textit{P. falciparum} LeuRS. Taken together, these results support continued efforts to develop 3-aminomethyl benzoxaboroles as novel antimalarial agents.

\textbf{ACKNOWLEDGMENTS}

We thank Fernando Rock, Anacor Pharmaceuticals; Case McNamara, California Institute for Biomedical Research; Christian Nsanzabana, Foundation for Innovative New Diagnostics; and Melissa Conrad, University of California, San Francisco, for helpful discussions.

\textbf{FUNDING INFORMATION}

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REFERENCES


Figure legends

Fig. 1. *In vitro* activity of benzoxaboroles against cultured *P. falciparum*. IC_{50} values against the W2 strain of *P. falciparum* were determined from two independent experiments, each performed in duplicate. cLogD_{7.4} was calculated using ChemAxon software under the condition of pH = 7.4. Boc, tert-butoxycarbonyl.

Fig. 2. Stage specificity of action of AN6426. (A) Parasitemias were compared between W2 strain *P. falciparum* incubated with 2 μM AN6426, 1.3 μM chloroquine, or 0.1% DMSO for 8 h intervals across the life cycle and then continued in culture until the following ring stage, when parasitemias were counted and compared. Error bars represent SD for 3 independent experiments. (B) Photomicrographs of representative parasites treated with AN6426 and untreated controls are shown. ER, LR: early and late rings; ET, MT, LT: early, mid, and late trophozoites; S: schizonts.

Fig. 3. Selection and analysis of AN6426-resistant parasites. The selection schematic is shown at the top, including concentration of AN6426 and duration of incubation for each step. IC_{50}s for the indicated compounds are shown after each selection; values include SDs for at least 3 independent experiments, each with 2 replicates. SNPs identified in these parasites by whole genome and dideoxy sequencing (**), or by dideoxy sequencing alone (*), based on comparison with the 3D7 reference strain, are shown.

Fig. 4. Structure of *P. falciparum* LeuRS editing domain and mapping of resistant mutants. (A) *P. falciparum* LeuRS editing domain surface model (white) and cartoon representation (blue) with resistance mutant residues shown as red sticks. (B) Zoomed view of the *Pf*LeuRS editing site. (C) Docking of AN6426-AMP adduct into the editing site of *Pf*LeuRS.
The adduct was placed by overlapping PfLeuRS to the complex of C. muris LeuRS with AN6426–AMP (PDB: 2FOM). Root mean square deviation over backbone atoms was 1.3 Å. (D) Sequence alignment of LeuRSs of plasmodia, other eukaryotic pathogens, and E. coli. Residues that are changed in P. falciparum AN6426-resistant mutants are highlighted in red.

Fig. 5. Biochemical support for action of benzoxaboroles against LeuRS. (A) Effects of benzoxaboroles and positive (cycloheximide) and negative (artemisinin) controls on [14C]leucine incorporation by wild-type Dd2 strain P. falciparum. (B) Effects of AN6426 on [14C]leucine incorporation by parasites selected for resistance. Concentrations studied were multiples of IC50s, determined based on triplicate measures (IC50s for AN6426, 420 nM; cycloheximide, 130 nM; artemisinin, 8.7 nM). (C, D) Effects of norvaline on growth of cultured parasites selected for resistance in the absence (C) or presence (D) of exogenous leucine (0.025 g/L). Error bars represent standard deviations for at least three independent experiments.

Table 1: In vitro and In vivo antimalarial activities of AN6426 and AN8432

IC50 results (±SD) are means from at least 3 experiments, each with duplicate readings. CC50 results for Jurkat and HepG2 cells are means from 2 experiments, each with at least 2 replicates. The value for field isolates is the mean of readings for 7 independent isolates collected in Tororo, Uganda in 2012 and performed as described previously (24). In vivo ED90 values are based on comparisons of parasitemias between treated and control animals on the fourth day after inoculation of parasites and initiation of treatment. ND, not determined.
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<th>R₃</th>
<th>R₄</th>
<th>R₅</th>
<th>R₇</th>
<th>cLogD₇.₄</th>
<th>IC₅₀ (μM)</th>
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Table 1: *In vitro* and *In vivo* antimalarial activities of AN6426 and AN8432

<table>
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<th>Mammalian cells</th>
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<td>CC₅₀ (µM)</td>
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<td>AN6426</td>
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<td>7.4</td>
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<td>0.31 ± 0.18</td>
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<td>0.42 ± 0.15</td>
<td>0.35 ± 0.13</td>
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<td>AN8432</td>
<td>0.28 ± 0.01</td>
<td>106 ± 38</td>
<td>16.2</td>
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<td>0.49 ± 0.29</td>
<td>&gt;25</td>
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IC₅₀ results (±SD) are means from at least 3 experiments, each with duplicate readings. CC₅₀ results for Jurkat and HepG2 cells are means from 2 experiments, each with at least 2 replicates. The value for field isolates is the mean of readings for 7 independent isolates collected in Tororo, Uganda in 2013 and performed as described previously (40). In *vivo* ED₉₀ values are based on comparisons of parasitemias between treated and control animals on the fourth day after inoculation of parasites and initiation of treatment. ND, not determined.
Wild type

6 × 10⁷ Dd2 parasites/flask

0.4 µM
31 Days

R1-1 R1-2

IC₅₀  (µM)

AN6426  0.42  ± 0.15
0.70 ± 0.15
0.63 ± 0.21
6.7 ± 0.06
8.6 ± 1.2
22 ± 1.3
25 ± 1.2
4.7 ± 0.5

AN8432

0.49  ± 0.18
0.94 ± 0.13
6.4 ± 0.19
18 ± 0.63

Chloroquine 48 ± 8.1
34 ± 5.3
20 ± 6.9
9.5 ± 3.9

Lumefantrine

5.6 ± 0.1
2.6 ± 1.3
0.95 ± 1.1
0.97 ± 0.3

Mefloquine

2.2 ± 0.43
0.13 ± 0.1
0.97 ± 0.1
0.23 ± 0.1

Piperaquine

15 ± 0.03
7.5 ± 1.7
11 ± 2.5
5.7 ± 2.2

Dihydroartemisinin

7.9 ± 1.7
6.1 ± 0.7
7.0 ± 8.1
3.6 ± 1.0

IC₅₀ (nM)

PF3D7_0622800

None

V630L** E628G* T400I** V568L*

V630L** E628G* T400I** M416T**

V568L** V568L* M416T* M416T*

PF3D7_1218100

None

M416T**

M416T* M416T*

M416T* M416T*

Cloned by limiting dilution

R3Rev

IC₅₀  4.7 µM
0 µM
150 Days

Revertant R3

0.4 µM
1 µM
10 µM
IC₅₀  6.2 µM
32 µM
150 Days

IC₅₀  1.7 µM
IC₅₀  6.2 µM
IC₅₀  32 µM
150 Days

0.4 µM
10 µM
150 Days

Revertant R3
on June 14, 2016 by UCSF LIBRARY

A B

C

D

AMP binding loop
Thr rich region

AN6426-AMP Inhibition adduct

P. falciparum
LeuRS

D

T400

V568

E628-V630

P. falciparum 395TLKPSTAYGQN405 563ISTGIVPCVSS573 625YSEKRYVSVSY633
P. knowlesi TLKYSTMYGQN IATGIVPCVSS YESEKITSY
P. yoelii TLKYSTMYGQN IATGIVPCVSS YESEKITSY
P. chabaudi TLKYSTMYGQN IATATVPCVSS YESEKISSY
T. brucei TLCPSTMYGQT KGTGVTVCVPS VNDKLQSP
C. reinhardi TLCPSTMYGQT KGTGIVTSVPS CNQYIQOS
E. coli TTRPDTFMCGT YGTGAVMDVPG LAAAGSERD

Editing site

V630

E628