Antimalarial proteasome inhibitor reveals collateral sensitivity from intersubunit interactions and fitness cost of resistance.

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Antimalarial proteasome inhibitor reveals collateral sensitivity from intersubunit interactions and fitness cost of resistance

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We describe noncovalent, reversible asparagine ethylenediamine (AsnEDA) inhibitors of the Plasmodium falciparum proteasome (P20S) β5 subunit that spare all active subunits of human constitutive and immuno-proteasomes. The compounds are active against erythrocytic, sexual, and liver-stage parasites, against parasites resistant to current antimalarials, and against P. falciparum strains from patients in Africa. The β5 inhibitors synergize with a β2 inhibitor in vitro and in mice and with artemisinin. P. falciparum selected for resistance to an AsnEDA β5 inhibitor surprisingly harbored a point mutation in the noncatalytic β6 subunit. The β6 mutant was resistant to the species-selective P20S β5 inhibitor but remained sensitive to the species-nonspecific β5 inhibitors bortezomib and carfilzomib. Moreover, resistance to the P20S β5 inhibitor was accompanied by increased sensitivity to a P20S β2 inhibitor. Finally, the β5 inhibitor-resistant mutant had a fitness cost that was exacerbated by irradiation. Thus, used in combination, multistage-active inhibitors of the P20S β5 and β2 subunits afford synergistic antimalarial activity with a potential to delay the emergence of resistance to artemisinins and each other.

Plasmodium | malaria | proteasome inhibitors | artemisinin | collateral sensitivity

Each year malaria causes an estimated 200 million cases and nearly 500,000 deaths in children under 5 y of age, with the large majority of serious illnesses and deaths due to Plasmodium falciparum (1). Resistance to older antimalarials, such as chloroquine, is common, and resistance to the most effective newer treatments, artemisinin-based combination therapies (ACTs), is established in Southeast Asia (2–4). Compounds that target the preerythrocytic hepatic stage and block the development of the transmissible gametocyte form that are taken up by the mosquito are ideal candidates for malaria prevention, but few antimalarials are effective against these stages. The growing threat of ACT failure and the need to target nonerythrocytic stages underscore the need for drugs with new targets in the parasite. Proteasome inhibitors have the potential to fulfill both requirements (5), and the proteasome has emerged as an important target for antimalarial drug development (6–10). Eukaryotic proteasomes have two copies of each of the three proteolytically active subunits—chymotryptic β5, trypsin β2, and caspase-like β1—in each 20S core particle. Several proteasome inhibitors are effective in vitro against Plasmodium spp. at multiple stages of the parasite lifecycle, including the erythrocytic, liver, and gametocyte stages, and for the treatment of Plasmodium berghei-infected mice (9, 11–15). The human host has two major isoforms of the 20S, constitutive proteasomes (c-20S) and immunoproteasomes (i-20S), with subtle but significant differences in proteolytic activity and functions. Proteasomes are important in all human cells. Inhibition of c-20S is cytotoxic, and inhibition of i-20S can dampen immune responses against infection (16, 17). Proteasome inhibitors designed to treat infections must achieve a high degree of species selectivity from intersubunit interactions and fitness cost of resistance to artemisinins and each other.

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Significance

Protozoal proteasome is a validated target for antimalarial drug development, but species selectivity of reported inhibitors is suboptimal. Here we identify inhibitors with improved selectivity for malaria proteasome β5 subunit over each active subunit of human proteasomes. These compounds kill the parasite in each stage of its life cycle. They interact synergistically with a β2 inhibitor and with artemisinin. Resistance to the β5 inhibitor arose through a point mutation in the nonproteolytic β6 subunit. The same mutation made the mutant strain more sensitive to a β2 inhibitor and less fit to withstand irradiation. These findings reveal complex interplay among proteasome subunits and introduce the prospect that combined inhibition of β2 and β5 subunits can afford synergism and thwart resistance.

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selectivity to avoid toxicity and impairment of the host’s immune response.

Developing pathogen-selective proteasome inhibitors that spare both i-20S and c-20S has been challenging, as proteasome subunits are highly conserved. Species-selective proteasome inhibitors were first reported for Mycobacterium tuberculosis (18, 19) and then for Plasmodium sp (6, 10), trypanosomes and Leishmania (20). However, most studies confined assessment of selectivity to testing the impact of these compounds on host c-20S, whereas i-20S inhibition was not examined. Moreover, most studies tested activity against only one of the human proteasome subunits.

Here we present a class of proteasome inhibitors that is highly species-selective for the P20S β5 subunit over all active subunits of both human c-20S and i-20S. Use of these inhibitors revealed three previously unreported findings with P20S inhibition: late-stage gametocytocidal activity and inhibition of gamete activation; marked synergy between a P20S β5 inhibitor and a β2 inhibitor; and association of resistance to a P20S β5 inhibitor with markedly increased sensitivity to inhibition of β2. The findings of synergy and collateral sensitivity suggest the potential value of capitalizing on interactions among different subunits of the parasite proteasome.

Results
Identification of Antimalarial Asparagine Ethylenediynes. A focused proteasome inhibitor library of around 180 compounds including three unique classes was synthesized in-house (18, 19, 21–24). We randomly selected 95 of these compounds at 10 μM with bortezomib, a pan-proteasome inhibitor, serving as a positive control (Fig. 1). Initial screens measured the ability of the compound to inhibit hydrolysis of a fluorogenic chymotryptic substrate, suc-Leu-Leu-Val-Tyr (LLVY)-AMC, by a P. falciparum lysate. We focused further on compounds that afforded >85% inhibition of suc-LLVY-AMC hydrolysis, comparable to the impact of bortezomib (Fig. 1C).

We next tested compounds against the erythrocytic stage of the parasite using a standard in vitro growth inhibition assay over 72 h (Fig. 1D) (25). Compounds in classes of N,C-capped dipeptides (22) and β-amino acid dipeptidomimetics (23) were potent against P. falciparum but were also potent inhibitors of i-20S and hence were not sufficiently species selective. We next focused on compounds belonging to a recently reported class of proteasome inhibitors, the asparagine ethylenediynes (AsnEDAs) (17). Among these, PKS3080 and PKS21004 showed the most potent antimalarial activity (Fig. 1A and SI Appendix, Table S1). PKS21003, a regioisomer of PKS21004, was inactive against the parasite and served as a negative control.

We assessed subunit selectivity with the activity-based proteasome probe MV151, a fluorescent vinyl sulfone that irreversibly labels the three catalytic subunits of the parasite proteasome (7, 26). PKS21004 and bortezomib specifically inhibited labeling of P20S β5, whereas PKS21003 did not (Fig. 1B). In contrast, PKS21004 did not block MV151 from labeling P20S β2 or β1. Importantly, none of the AsnEDAs showed appreciable inhibitory activity against the human c-20S subunits β1c or β2c or the i-20S subunits β1i or β2i (17). In contrast, in our assay, Trp-Leu-Trp-vinyl sulfone (WLW-VS) (6), reported to be highly selective for P20S β2 over human c-20S β2c and human i-20S β2i, was highly potent against human c-20S β5c and human i-20S β5i, with IC50s of 100 nM and 23 nM, respectively (SI Appendix, Fig. S1). It is thus important to confirm species selectivity by testing all proteasome subunits, as inhibitors intended for one subunit of one species can inhibit different subunits of a different species.

The ability of PKS21004 to block the covalent reaction with P20S β5 and MV151 was dose-dependent (Fig. 1B, Lower). Poly-ubiquitinated proteins accumulated in P. falciparum (Pf3D7 strain) inhibited within 4 h of exposure to 1 μM PKS21004, similar to results with 10 μM bortezomib (Fig. 1C).

We tested the cytotoxicity of PKS21004 and PKS21003 against the human hematoma cell line HepG2, which predominantly expresses c-20S, the human lymphoma cell line Karpas 1106P, which predominantly expresses i-20S, and primary mouse bone marrow-derived macrophages (MBM), which also express i-20S. PKS21004 was selectively toxic for P. falciparum over the mammalian cells, while PKS21003 was nontoxic for both P. falciparum and mammalian cells (Fig. 1D and SI Appendix, Table S1).

Docking and Structure–Activity Relationship Studies to Improve Species Selectivity. To guide structure–activity (SAR) studies, we modeled the binding mode of PKS21004 in P20S, based on the recently solved cryo-electron microscopy structure of PKS21004 bound to i-20S (17). Given the high sequence and structural similarity between human i-20S and P20S for the relevant β5 and β6 subunits (~60% sequence similarity and ~1.1-Å rmsd for the dimers’ Cos), we predicted that the binding mode of PKS21004 in P20S would be similar to that in i-20S. In i-20S β5i (Fig. 2A), the biphenyl moiety of PKS21004 binds to the S1 pocket and S1 side pocket, while the Asn(β6-butyl) binds to the S3 pocket, forming hydrogen bonds with Ser27 and Ser129 of β5i. N-capped phenylpropionate of the inhibitor binds to S4, a solvent-exposed pocket formed by amino acid residues of β6 (17). In P20S β5i, PKS21004 is predicted to form very similar interactions with the S1, S3, and S4 pockets (Fig. 2B and SI Appendix, Fig. S2).

Using this model, we conducted SAR studies and developed the species-selective P20S inhibitors PKS21224 and PKS21287 (Fig. 2C) by replacing the biphenyl of PKS21004 with heteroaromatic rings (PKS21208 and PKS21229) and by replacing the tosyl with a hydrophilic isoaxazol carboxamide. Both compounds were highly potent and modestly selective for P20S over c-20S and i-20S (Fig. 2F). All compounds were highly active against P. falciparum and selective against parasite versus HepG2 cells. PKS21224 and PKS21287, which both have lower logD (the log of the partition of a chemical compound between the lipid and aqueous phases) than PKS21004, were both the least active against i-20S and c-20S but were highly potent against P. falciparum, with EC50 of 9 nM and 34 nM, respectively. However, PKS21287 had poor cell permeability in a parallel artificial membrane permeability assay (SI Appendix, Table S2). We therefore modified the pyrazole moiety by changing the C–C linker to a C–N linker.
resulting in TDI4258. PKS21287 and TDI4258 showed comparable IC50s against c-20S and i-20S and EC50s against *P. falciparum* 3D7 (Fig. 2F). The logD of TDI4258 at pH 7.4 (1.85) was greater than that of PKS21287 (1.49). The change was associated with an improvement in permeability from an undetectable level for PKS21287 to 64 nm/s at pH 7.4 and 56 nm/s at pH 5.0 for TDI4258. However, only 2.3% of an orally administered dose of TDI4258 was bioavailable. Consistent with their reduced inhibitory activity against human proteasomes, PKS21224, PKS21287, and TDI4258 were much less cytotoxic to mammalian cells than PKS21004 and PKS21221 (Fig. 2F). As expected, PKS21221 and TDI4258 dose-dependently blocked the labeling of Pf20S β5 by MV151 (Fig. 2D).

### Determination of IC50s Against Pf20S β5

Using purified Pf20S, we observed that hydrolysis of substrate suc-LLVY-AMC was reduced by 20–30% in the presence of the β5-specific inhibitors PKS21004, PKS21221, or TDI4258 at 10 μM and by 50% by the β2-specific inhibitor WLW-VS at 0.5 μM (Fig. 2E) (6). This suggested that both β2 and β5 of the Pf20S can hydrolyze

**Fig. 2.** AsnEDAs with improved potency and selectivity. (A and B) Binding mode of PKS21004 in the i-20S structure (17) (A) and in Pf20S (8) predicted by docking. Only β5 and β6 are shown. Key amino acid residues involved in ligand binding are labeled in the respective enzymes. (C) Structures of AsnEDAs. (D) Dose-dependent inhibition of Pf20S by PKS21221 and by TDI4258 assessed by the activity-based probe MV151. (E) LLVY-AMC hydrolysis by Pf20S was assessed in the presence of AsnEDAs and WLW-VS, both separately and in combination. (F) IC50s of all AsnEDAs against human i-20S, c-20S, and Pf20S and EC50s against *P. falciparum* 3D7 and HepG2. All data are averages of at least three independent experiments.
Activity Against the Asexual Erythrocytic Stage. We next determined the EC50 of PKS21004 against established laboratory strains of *P. falciparum* with different points of origin and with different drug-resistance profiles (Fig. 3A). PKS21004 was potent against all lines tested. EC50 ranged from 1.5 nM against *P. falciparum* V1S, a multidrug-resistant South East Asian strain, to 10 nM against *P. falciparum* HB3, a Central American strain resistant to pyrimethamine. Five more recently isolated parasite strains from the Greater Mekong subregion, two of which are artemisinin sensitive and three of which are artemisinin resistant, showed a narrow range of EC50 (1.5 nM–6 nM) to PKS21004, PKS21221, PKS21287, and PKS21224 also showed activity (SI Appendix, Fig. S4).

TDI4258 was further tested for ex vivo activity against *P. falciparum* samples from 38 malaria patients in Uganda. The EC50 ranged from 11–81 nM, with a mean of 30 nM (Fig. 3B and SI Appendix, Table S3), which is consistent with EC50 against the *P. falciparum* laboratory strains. There was no detectable association between sensitivities to TDI4258 and chloroquine among the tested Ugandan isolates (SI Appendix, Table S3) (27).

To understand the dynamics of antimalarial action, we assessed the in vitro killing profile of PKS21224. The in vitro parasite reduction ratio (PRR) assay directly measures the number of parasites that remain viable at different time points post drug exposure (28). PKS21224 showed a *P. falciparum* killing profile similar to that of pyrimethamine (log PRRs of 2.7 and 2.8, respectively) (Fig. 3C) with a lag phase (the time needed to observe the maximal

Fig. 3. Erythrocytic stage and ex vivo activities and PRR of AsnEDAs. (A) IC50s of PKS21004 against various strains of *P. falciparum*. (B) Ex vivo activity of TDI4258 against 38 clinical samples of isolates from Uganda. The geometric mean EC50 was calculated as 30 nM. (C) PRR of PKS21224. An in vitro parasite reduction rate assay was used to determine onset of action and rate of killing as described (28). *P. falciparum* was exposed to PKS21224 at a concentration equivalent to 10× EC50. The number of viable parasites at each time point was determined as described. Four independent serial dilutions were done with each sample to correct for experimental variation; error bars indicate SD. Previous results with current antimalarial drugs tested using the same protocol are shown for comparison (28).

Activity Against Hepatic-Stage Parasites. The activity of PKS21004 against preerythrocytic-stage parasites was tested using a *P. berghei*-HepG2 model. Luciferase-expressing *P. berghei* sporozoites were added to HepG2 cells in the presence or absence of PKS21004 (30). Infectivity was determined 48 h postinfection by quantification of luciferase activity. PKS21004 demonstrated potent inhibition of liver-stage development with an EC50 of 16.3 nM (Fig. 4D). Due to the selectivity of our compounds, we were able to determine the effect of proteasome inhibition directly on sporozoite-to-merosone development before and after hepatic cell invasion. Our findings killing effects of the drug) of 24 h and a parasite clearance time (the time needed to kill 99.9% of the initial population) of 61 h.

Activity Against *P. falciparum* Transmission and Mosquito Stages. We tested activities of PKS21004, PKS21224, and PKS21287 against stage III (Fig. 4A) and stage V *P. falciparum* gametocytes with dihydroartemisinin (DHA) as a positive control (Fig. 4B). All three compounds showed in vitro antigametocyte activity after 24, 48, and 72 h, with EC50s of 28, 140, and 212 nM, respectively. Antigametocyte activity increased by 2.3- to 3.6-fold from 24 to 48 h (SI Appendix, Fig. S5 and Table S4) with little additional enhancement of killing at 72 h. In comparison, the activity of DHA did not improve with exposure longer than 24 h, consistent with its rapid killing mechanism. Our compounds were also effective against mature stage V gametocytes at 72 h, with EC50s of 57 nM, 227 nM, and 229 nM for PKS21004, PKS21224, and PKS21287, respectively (Fig. 4B).

PKS21224, PKS21287, and TDI4258 were further tested in a gamete activation assay. The activation of male but not female gametes was susceptible to inhibition of P20S β5, with EC50s of 140 nM–240 nM (Fig. 4C and SI Appendix, Table S5) (29). The inability of male gametes to undergo activation would preclude the generation of infectious sporozoites and block malaria transmission.

Fig. 4. Antimalarial activities of AsnEDAs against gametocyte stage, gamete activation, and liver stage. (A and B) AsnEDAs and DHA as a positive control were dose-dependently evaluated against gametocytes at stages III (A) and V (B). (C) Inhibition of *P. falciparum* gamete activation. PKS21224, PKS21287, and TDI4258 were dose-dependently evaluated in a dual gamete activation assay, FG, female gametes; MG, male gametes. (D) Dose–response curve of PKS21004’s activity against *P. berghei* preerythrocytic stages in the HepG2 model. Data shown are the average of three independent experiments, each performed in triplicate.
suggest that functional proteasomes are required for parasite development within the hepatocyte.

**Synergy Between DHA and Proteasome Inhibitors.** Parasite proteasome inhibition reportedly can reverse artemisinin resistance (5), where artemisinin resistance is defined as slow clearance post drug exposure as measured by a ring-stage survival assay (RSA) (31). We tested TD4258 in combination with DHA using a modified ring-stage survival assay (RSA) with artemisinin-resistant parasites that contain a mutant Kelch (R539T) protein in a multidrug-resistant Dd2 strain background (32). We exposed early 1- to 3-h ring-stage parasites to a 3-h pulse of DHA and continuous TD4258 for 72 h, as described (5, 31). Parasitemia was measured at 72 h post initial exposure. Isobologram analysis demonstrated synergistic killing of artemisinin-resistant parasites by the combination of TD4258 and DHA (Fig. 5A). Wild-type Dd2 parasites remained sensitive to DHA and thus could not be assessed for synergy in this assay.

Synergistic in Vitro Activity Against *P. falciparum* of Proteasome Inhibitors That Target β2 and β5 Pf20S Subunits. To test if the synergy we noted above between β2 and β5 inhibitors against P20S pertained to intact parasites, we tested β5 inhibitors together with various concentrations of WLW-VS against the erythrocytic stages of *P. falciparum* as described (33) and determined the viability of intraerythrocytic parasites at 72 h. PKS21004, TD4258, and bortezomib all gave fractional inhibition concentration (FIC) values of ≤0.5, demonstrating a synergistic effect of simultaneous inhibition of P20S β5 and β2 (Fig. 5C). This suggests that the binding of a ligand to one subunit can affect the binding of another ligand to a distal subunit of the P20S.

Next we tested combined inhibition of P20S β2 and β5 (in a fixed ratio of TD4258:WLW-VS of 4:1) in the presence of DHA in a ring-stage survival assay with an artemisinin-resistant strain, as described above. Potent synergy after 3 h exposure to the compounds was evident when viability was assessed 72 h after initial exposure (Fig. 5B).

**In Vivo Antimalarial Activity.** The in vivo efficacy of TD4258 was tested in a mouse model of *Plasmodium yoelii* infection. Female CF1 mice were infected i.v. with *3 × 10^5* *P. yoelii*-parasitized erythrocytes and were treated daily for 4 d beginning 1 d after inoculation. TD4258 (15 mg/kg), WLW-VS (60 mg/kg), and WLW-VS (60 mg/kg) plus TD4258 (15 mg/kg) were administered i.p. in DMSO. Parasitemia was determined on day 5. The experiment was performed twice, and the data were pooled (Fig. 5D). TD4258 did not yield a statistically significant reduction in parasitemia compared with DMSO (21.5% reduction, *P* = 0.16), likely reflecting its short (30-min) half-life in the mouse (SI Appendix, Table S2). WLW-VS reduced parasitemia by 42% (*P* = 0.0014). The combination of TD4258 and WLW-VS reduced parasitemia by 95% (*P* < 0.0001). Thus, the synergy observed in vitro between inhibition of P20S β2 and β5 appeared to manifest in vivo during murine *P. yoelii* infection.

**Mechanism of Resistance.** For additional target validation and to explore potential mechanisms of resistance, we cultured the Dd2 strain of *P. falciparum* parasites in gradually increasing concentrations of PKS21004 from the initial EC_{50} of 4 nM to 192 nM. After 4 mo, we obtained a stably resistant parasite line. We were not able to generate any resistant parasites using the same protocol with 3D7-strain parasites. We cloned parasites from the resistant Dd2 line and maintained two clonal lines, *P. falciparum* Dd2-R1 and Dd2-R2. Both demonstrated ∼130-fold resistance to PKS21004 (Fig. 6A). Resistance extended to other members of the AsnEDA class; for example, both lines were 40-fold more resistant to PKS21224 and 13-fold more resistant to TD4258 than unselected Dd2 parasites. However, resistance was specific, in that there was little or no change in EC_{50} against DHA, atovaquone, or piperaquine (Fig. 6A).

Intriguingly, the AsnEDA-resistant parasites did not demonstrate resistance to another β5 inhibitor, bortezomib; on the contrary, they demonstrated a twofold increase in susceptibility to bortezomib. Bortezomib is a dipeptide boronate that forms a reversible covalent boron–oxygen bond with the hydroxyl group of the active site Thr^{205} and primarily targets the β5 subunit (34). Its P1 Leu and P2 Phe side chains bind to S1 and S2 pockets, respectively, and its pyrazinoyl moiety binds along the substrate-binding groove (35). The mutation conferring resistance to the AsnEDA β5 inhibitors slightly enhanced the action of a dipeptidyl boronate with a different binding pose. Increased sensitivity was also seen with carfilzomib, a peptide epoxyketone proteasome inhibitor drug. Moreover, the AsnEDA-resistant parasites had 12- to 14-fold increased sensitivity to the β2 inhibitor WLW-VS.

To gain insight into the mechanism of resistance, we sequenced the genomes of the parental Dd2 strain and the two resistant clones, *P. falciparum* Dd2-R1 and Dd2-R2. Surprisingly,
as summarized in SI Appendix, Table S6, both resistant clones bear a G-to-A mutation in the gene encoding the β6 proteasome subunit (PF3D7_0518300), leading to an A117D amino acid change; no mutation was found in the target β5 subunit. Both clonal lines also harbored mutations in PF3D7_0822400, a gene of unknown function (SI Appendix, Table S6). One clone also carried a mutation in PF3D7_0724700, which encodes another protein of unknown function. There was no evidence of copy number variation of any genes.

The β6 proteasome subunit has no proteolytic activity. A117 is predicted to reside in an α-helix adjacent to a loop (Tyr152–Cyc159), part of which forms a wall between the β5 S1 side pocket (S1SP) and the β5 S3 pocket. A117 is closely opposed to β6 Tyr150 and β6 Tyr158, whose side chains are only 3.4 and 3.7 Å distant from A117, respectively (Fig. 6B). The A117D mutation introduces a negatively charged side chain that is expected to clash with these two tyrosines (Fig. 6B), forcing the loop to undergo conformational changes that would alter the shape and polarity of the S1SP and the S3 pocket in β5, directly affecting the binding of PKS21004. In contrast, the A117D and accompanying conformational changes are distal to the binding sites of both bortezomib and carfilzomib (Fig. 6C), potentially explaining why the mutant was not resistant to both bortezomib and carfilzomib.

To confirm that the mutation affected the binding affinity of PKS21004, we enriched mutant Pf20S(β6A117D) and conducted subunit labeling experiments (Fig. 6D, Upper). In agreement with the shift in EC_{50}, PKS21004’s potency in blocking the labeling of Pf20S by MV151 was reduced compared with its ability to block the labeling of the wild-type proteasome. Conversely, the potency of WLW-VS in blocking the labeling of Pf20S by MV151 was greater in the mutant than in the wild type (Fig. 6D, Right).

Drug resistance-mediating mutations may be accompanied by loss of fitness (36). To test if the β6 A117D mutation imposed a fitness cost, we performed a plaque assay comparing the Pf Dd2 parent with the Pf Dd2 R1 parasite line and identified subtle but significant differences in growth rates under normal culture conditions, with the Pf Dd2 parent appearing to have more robust growth than the resistant parasite line (SI Appendix, Fig. S6). We then exposed intraerythrocytic P. falciparum to X-ray irradiation at 50 and 100 Gy, which is likely to damage DNA and protein, and measured the parasite’s ability to recover from this damage. As summarized in SI Appendix, Table S6, both resistant clones bear a G-to-A mutation in the gene encoding the β6 proteasome subunit (PF3D7_0518300), leading to an A117D amino acid change; no mutation was found in the target β5 subunit. Both clonal lines also harbored mutations in PF3D7_0822400, a gene of unknown function (SI Appendix, Table S6). One clone also carried a mutation in PF3D7_0724700, which encodes another protein of unknown function. There was no evidence of copy number variation of any genes.

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stress by monitoring the recovery of parasitemia to 2% (Fig. 6E, Left) (38). PlurD2 recovered by days 1.3 (after 50 Gy) and 6.8 (after 100 Gy), whereas PFDd2 R1 recovered by days 4.3 (50 Gy) and 16.7 (100 Gy) (Fig. 6E, Right), demonstrating a defect in the ability of the resistant parasite line to withstand radiation stress.

Discussion

The spread of drug-resistant parasites is a major obstacle to the WHO’s goal of reducing morbidity and mortality attributable to malaria by 90% by 2030. Here we present a set of compounds that spare both human constitutive and immuno-proteasomes and target the malaria parasite proteasome. The proteasome is a key regulator of the parasite’s ability to proliferate and adapt to stress, likely including stresses imposed by artemisinin antimalarials (5). We show that simultaneously targeting two different catalytic proteasome subunits provides synergistic killing of P. falciparum as well as synergism with artemisinin against artemisinin-resistant parasites. Moreover, a mutation in the parasite proteasome that confers resistance to blockade of one subunit can lead to increased sensitivity to blockade of another. These results suggest that, despite the conservation of proteasomes among eukaryotes, targeting P. falciparum proteasome subunits offers a more promising approach for antimalarial chemotherapy than was previously apparent.

We determined the activity of the AsnEDAs at multiple stages of the malaria lifecycle. Activity against the gametocyte stage has previously been determined only with nonselective inhibitors. Here we found that the relatively selective AsnEDAs were effective against stage III and stage V gametocytes, the stage poised for transmission to the mosquito. The AsnEDAs also blocked male gamete activation, supporting the idea that proteasome inhibitors can act as transmission-blocking antimalarials at several stages. Our studies on the liver stage are unique in that the species-selective nature of the compounds allowed us to treat intrahepatic parasites for a prolonged time without toxicity to the host cell. The efficacy of AsnEDAs at the preerythrocytic stage indicates the potential use of proteasome inhibitors for malaria prevention. This observation opens the door to study the biology of the parasite proteasome in this important stage, which is associated with extensive parasite remodeling (30).

The availability of cryo-EM structures (17) to guide modeling studies helped us better understand and improve the selectivity of inhibitors for the P. falciparum proteasome against both the human constitutive proteasome and immunoproteasome. Expression of human c-20S is ubiquitous, while i-20S is expressed predominantly in immune cells and proportionately more so when they are stimulated with IFN-γ. The immunoproteasome plays important roles in the activation of immune cells and in immune surveillance. Selectivity for the pathogen’s proteasome is crucial for proteasome inhibitors to advance as antiinfective agents.

Resistance to AsnEDA β5 inhibitors emerged in vitro only after prolonged in vitro selection in one of two P. falciparum lines tested. The resistance was associated with a mutation not in the target β5 subunit but in the adjacent β6 noncatalytic structural subunit. Modeling studies suggest that the amino acid change in β6 may alter the S1SP and S3 pockets formed by β5 and β6. This demonstrates that a mutation in the β6 subunit confers resistance to Pl20S β5 inhibitors by affecting distal region of the substrate binding cleft. Similarly, a single amino acid mutation in the trypansomal proteasome nonproteolytic β4 subunit conferred resistance to a newly identified kinetoplastid-specific proteasome inhibitor (20). The additional mutation in PF3D7_0822400, a conserved protein with no known function, may contribute to resistance to proteasome inhibitors; a supportive or adaptive role for this mutation has yet to be explored.

The efficacy of AsnEDAs and WLW-VS with cultured parasites and in murine malaria infection suggests that combination therapy with inhibitors of both the Plasmodium Pl20S β5 and β2 subunits could offer an effective therapy for malaria, provided that ongoing medicinal chemistry efforts lead to improved oral bioavailability. Targeting both these subunits was synergistic against purified proteasomes, erythrocytic-stage parasites in vitro, and erythrocytic-stage parasites in the mouse. Similarly, enhanced cytotoxicity from combined inhibition of the β2 and β5 subunits has been described in human tumor cells (39, 40). The findings reported here provide a biologic rationale for medicinal chemistry campaigns directed at both subunits and suggest that it will be fruitful to consider each set of inhibitors in the context of its interaction with others. In addition, given that resistance to a β5 inhibitor led to increased sensitivity to a β2 inhibitor both in enzymatic studies and in vitro culture, dual inhibition appears to lead to a highly desirable resistance-thwarting strategy, collateral sensitivity (41). Further structural studies with wild-type and A117D-mutant strains of Pl20S, with and without inhibitors, should shed light on the mechanism driving the increased sensitivity to WLW-VS observed in PKS21004-resistant parasites.

A fitness cost that accompanies the development of drug resistance has been described for various microorganisms, including P. falciparum, and can help limit the spread of drug-resistant parasites. Similarly, yeast bearing mutations associated with bortezomib resistance grew more slowly than wild type, especially under stress conditions (42). The majority of mutations reported to confer resistance to proteasome inhibitors in yeast or cancer cells are in β5, in contrast to the β6 mutation we identified in each of two resistant lines of P. falciparum.

The present findings strengthen the case for targeting the parasite proteasome in antimalarial drug development. The inhibitors described demonstrate the potential to inhibit the P. falciparum proteasome while sparing not only the human constitutive proteasome but also the human immunoproteasome. Further, they show that inhibition of the P. falciparum β5 subunit can kill or block parasite development in all stages of the parasite lifecycle—in the asexual blood stage, during hepatic invasion and development, and in the development and activation of gametocytes—offering the potential for chemoprevention (by killing liver stages) and transmission blocking (by inhibiting gametocytes and gamete activation), stages for which there are few active antimalarials. Finally, these compounds revealed the synergistic potential of exploiting long-range interactions between distinct proteolytic subunits of Pl20S as well as the opportunity for collateral sensitivity. Further development of parasite proteasome inhibitors will be aided by a deeper understanding of the complex relationships among the proteasome subunits. Such an understanding may be translated clinically to maximize potency and raise the barrier to the development of resistance.

Materials and Methods

The SI Appendix, SI Methods details materials and methods for P. falciparum cultivation; Pl20S enrichment; biochemical characterization of Pl20S; IC50 determinations against c-20S, i-20S, and Pl20S; dose–response studies against parasites in erythrocytic, preerythrocytic, and gametocyte asexual stages and for chemoprevention (by killing liver stages) and transmission blocking (by inhibiting gametocytes and gamete activation), stages for which there are few active antimalarials. Finally, these compounds revealed the synergistic potential of exploiting long-range interactions between distinct proteolytic subunits of Pl20S as well as the opportunity for collateral sensitivity. Further development of parasite proteasome inhibitors will be aided by a deeper understanding of the complex relationships among the proteasome subunits. Such an understanding may be translated clinically to maximize potency and raise the barrier to the development of resistance.

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