Targeting the *Plasmodium vivax* equilibrative nucleoside transporter 1 (PvENT1) for antimalarial drug development

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**Abstract**

Infection with *Plasmodium falciparum* and *vivax* cause most cases of malaria. Emerging resistance to current antimalarial medications makes new drug development imperative. Ideally a new antimalarial drug should treat both *falciparum* and *vivax* malaria. Because malaria parasites are purine auxotrophic, they rely on purines imported from the host erythrocyte via Equilibrative Nucleoside Transporters (ENTs). Thus, the purine import transporters represent a potential target for antimalarial drug development. For *falciparum* parasites the primary purine transporter is the *P. falciparum* Equilibrative Nucleoside Transporter Type 1 (PfENT1). Recently we identified potent PfENT1 inhibitors with nanomolar IC50 values using a robust, high throughputscreening assay. In the current work we characterized the *Plasmodium vivax* ENT1 (PvENT1) homologue and its sensitivity to the PfENT1 inhibitors. We expressed a yeast codon-optimized PfENT1 gene in Saccharomyces cerevisiae. PvENT1 expressing yeast imported both purines ([3H]adenosine) and pyrimidines ([3H]uridine), whereas wild type (fixIΔ) yeast did not. Based on radiolabel substrate uptake inhibition experiments, inosine had the lowest IC50 (3.8 μM), compared to guanosine (14.9 μM) and adenosine (142 μM). For pyrimidines, thymidine had an IC50 of 183 μM (vs. cytidine and uridine; mM range). IC50 values were higher for nucleobases compared to the corresponding nucleosides; hypoxanthine had a 25-fold higher IC50 than inosine. The archetypal human ENT1 inhibitor 4-nitrobenzylthioinosine (NBMPR) had no effect on PvENT1, whereas dipyrядimole inhibited PvENT1, albeit with a 40 μM IC50, a 1000-fold less sensitive than human ENT1 (hENT1). The PfENT1 inhibitors blocked transport activity of PvENT1 and the five known naturally occurring non-synonymous single nucleotide polymorphisms (SNPs) with similar IC50 values. Thus, the PfENT1 inhibitors also target PvENT1. This implies that development of novel antimalarial drugs that target both *falciparum* and *vivax* ENT1 may be feasible.

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1. Introduction

Malaria is a major global health problem and a socioeconomic burden in malaria endemic countries (Sachs and Malaney, 2002). According to the World Health Organization (WHO), in 2014 approximately 3.4 billion people were at risk for malaria infection (World Health Organization, 2014). Over 200 million clinical cases of malaria resulted in ~600,000 deaths. Most deaths occurred in sub-Saharan Africa in young children and pregnant women (Snow et al., 2005; World Health Organization, 2014). Malaria is caused by infection with single-cell protozoan parasites from the genus *Plasmodium*. Five *Plasmodium* species infect humans (*Plasmodium falciparum*, *vivax*, *malariae*, *ovale*, and *knowlesi*). Ninety percent of clinical cases are due to infection with either *P. falciparum* or *Plasmodium vivax* (World Health Organization, 2014). *P. falciparum* is associated with the highest mortality (~80% of all malaria-related deaths) but *P. vivax* infection is prevalent and associated with high morbidity (Rodgeron and Carter, 2008; Anstey et al., 2009).
The geographic overlap between *P. falciparum* and *P. vivax* endemic areas is significant, especially in tropical regions. Thus, new anti-malarial drugs should target both species.

The development of resistance to antimalarial drugs has been a recurring problem. Chloroquine (CQ) was the mainstay of antimalarial chemotherapy until CQ resistance developed worldwide (Wellens and Plowe, 2001). In 2006, the WHO recommended Artemisinin-based Combination Therapies (ACT) as first-line treatment for *P. falciparum* infection. Unfortunately, resistance to current ACT regimens is expanding in Southeast Asia (Dondorp et al., 2011; Arney et al., 2014; Hastings et al., 2015; Straimer et al., 2015). The fact that resistance to a three day ACT treatment course emerged in as little as a decade after the large scale introduction of ACTs as first line therapy underscores the importance of identifying new drug targets that take advantage of weaknesses in *Plasmodium* biology.

One potential target for the development of novel antimalarial drugs is the purine salvage pathway (Downie et al., 2008; Cassera et al., 2011; Frame et al., 2015a). Similar to other protozoa, *Plasmodium* species can perform de novo pyrimidine synthesis but are incapable of de novo purine synthesis (Manandhar and Van Dyke, 1975; Gero and O’Sullivan, 1990; Downie et al., 2008; Frame et al., 2015a). Therefore, *Plasmodium* parasites must import purines from the host cytoplasm. Imported purines are processed via the purine salvage pathway enzymes to form the purines required for RNA synthesis, DNA replication, and metabolism. Hence, the purine import and processing pathways are potential targets for antimalarial drug development (Downie et al., 2008; Ducati et al., 2013; Frame et al., 2015a).

*Plasmodium* parasites use equilibrative nucleoside transporters (ENT) to import purines (Landfear et al., 2004; Downie et al., 2008). Genomic sequence analysis of *P. falciparum* (3D7) and *P. vivax* (SalI) (www.PlasmoDB.org) shows that both species possess four putative ENT homologues: PFENT1-4 and PVENT1-4 (Martin et al., 2005; Kirk and Lehane, 2014). *P. falciparum* ENTs have been studied more extensively. Multiple genetic, biochemical, and functional experiments show that PFENT1 is the principle route for purine uptake into the *P. falciparum* parasites. PFENT1 is localized to the parasite plasma membrane and transports both purine and pyrimidine substrates (Carter et al., 2000a; Parker et al., 2000; Rager et al., 2001; Riegelhaupt et al., 2010a). Genetic knockout of the PFENT1 gene (pfent1Δ) is lethal if the parasites are grown in concentrations of purines present in human blood, <10 μM (Traut, 1994; El Bissati et al., 2006; El Bissati et al., 2008; Frame et al., 2015b). However, PFENT1-knockout parasites survive when grown in culture with supra-physiological purine concentrations (>100 μM) (El Bissati et al., 2006; El Bissati et al., 2008; Frame et al., 2015b). Thus, a secondary low affinity and/or low capacity purine transport pathway must be present, at least in the pfent1Δ parasites. The molecular basis for this secondary purine uptake pathway is unknown but may involve the AMP uptake pathway (Cassera et al., 2008) or possibly, PFENT4 (Frame et al., 2012).

To test whether chemical inhibition of PFENT1 would be lethal to *P. falciparum* parasites, we identified PFENT1 inhibitors using a yeast-based, high-throughput screen (HTS) (Frame et al., 2015b). We screened 64,500 compounds and identified 171 hits. Nine of the highest activity compounds that represent six distinct chemical scaffolds were characterized in depth. They blocked [3H]adenosine uptake into PFENT1-expressing yeast and into erythrocyte-free trophozoite stage parasites with 5–50 nM IC_{50} values and killed chloroquine-sensitive and -resistant *P. falciparum* parasites with 5–50 μM IC_{50} values (Frame et al., 2015b). These results provide strong support for the hypothesis that inhibition of purine uptake is a potential target for the development of novel antimalarial drugs.

Because of the extensive geographic overlap between *vivax* and *falciparum* malaria, an effective antimalarial drug should treat infection by both parasites. In the current work, we sought to characterize *P. vivax* ENT1 (PvENT1) functionally and determine whether the PfENT1 inhibitors also inhibit PvENT1. Based on its genomic sequence, PvENT1 is a 47.3 kDa, 416 amino acid protein. PvENT1 shares ~75% amino acid sequence identity with PFENT1. However, unlike its *P. falciparum* homologue, the pvent1 gene is only moderately AT rich (57%; vs. 72% pfent1). Although no crystal structures are available for any known ENTs, glycosylation scanning and modeling algorithms support an 11-transmembrane segment topology with a cytoplasmic N-terminus and an extracellular C-terminus (Sundaram et al., 2001; Valdés et al., 2009). Hydropathy plots suggest that PfENT1 has a similar transmembrane topology to PFENT1. In the current work, we expressed PvENT1 in *Saccharomyces cerevisiae*. We showed that a similar group of purine and pyrimidine nucleobases and nucleosides competed with radioactive uridine or adenine uptake into PfENT1-expressing yeast, but the measured IC_{50} were distinct from those previously reported for PFENT1-expressing yeast (Frame et al., 2015b). We determined the sensitivity of PvENT1 to the recently identified PfENT1 inhibitors. All nine PfENT1 inhibitors block PfENT1 with similar efficacy. Furthermore, the five known PfENT1 non-synonymous single nucleotide polymorphisms (SNP) have minimal impact on inhibitor potency.

2. Materials and methods

2.1. Purine auxotrophic yeast (ade2Δ)

All genetically modified yeast were created using the *S. cerevisiae* BY4741 strain that also had a deletion of the *FLU1* gene (MATa; his3Δ 1; leu2Δ 0; met15Δ 0; ura3Δ 0; fui1Δ::KanMX4) (Winzeler et al., 1999). To disrupt de novo purine synthesis in *S. cerevisiae*, we replaced the yeast *ADE2* gene (Chr. 15; phosphoribosyl-aminimidazole carboxylase) with the *hphNT1* (hygromycin B phosphotransferase) selectable marker using homologous recombination. Briefly, using two-step PCR, we created 45 nt 5′ and 3′ ade2Δ homology arms flanking the *hphNT1* gene (which was amplified from the pFA6a-hphNT1 plasmid; gift from Dr. Ian Willis). The following sets of primers were used: 1st-stepF: 5′-GACAAAACATCAAGTATGCGTACGCTGCAGGTCGACGGATCCCC-3′; 1st-stepR: 5′-GTATATCAATAAACTTATA-3′; 2nd-stepF: 5′-ACAAAATCATCAAGTATGCGTACGCTGCAGGTCGACGGATCCCC-3′; 2nd-stepR: 5′-GTATATCAATAAACTTATA-3′. The underlined portion corresponds to the sequence of yeast chromosomal DNA flanking the *ADE2* gene. Each 100 μL PCR reaction contained 50 ng DNA, 1x PfuUltra Reaction Buffer, 250 μM of each dNTP (dATP, dCTP, dGTP, dTTP), 0.2 μM of each primer (F/R), 2% (v/v) PfuUltra Fusion HS DNA Polymerase (Agilent). The PCR conditions were: 1 cycle—95 °C (2 min); 26 cycles—95 °C (30 s), 60 °C (30 s), 72 °C (30 s); 1 cycle—72 °C (2 min), 4 °C (hold). The PCR product was verified using ethidium bromide agarose gel electrophoresis, excised, and column purified. The ampiclon was transformed into yeast (see below) and plated on YPD + 500 μg/mL hygromycin B selection plates at 30 °C. The resulting yeast strain was MAFa; his3Δ 1; leu2Δ 0; met15Δ 0; ura3Δ 0; fui1Δ::KanMX4, ade2Δ::hphNT1. Disruption of the ade2Δ gene was confirmed by PCR. The purine auxotrophic yeast was pigmented red and displayed a retarded growth phenotype under purine starved conditions (Kokina et al., 2014). Single colonies of the purine auxotrophic yeast were picked and expanded for transformation with the pCM1895 constructs.

The purine auxotrophic yeast strain with the *ADE2* gene deletion (ade2Δ::hphNT1) was used in all experiments in this paper and was
either transformed with an episomal expression plasmid containing the PvENT1 gene or an empty vector (EV) lacking the PvENT1 gene. All subsequent references to “yeast” in the Results section refer to this purine auxotrophic strain.

2.2. PvENT1 DNA plasmid constructs

We synthesized a yeast codon-optimized version of the PvENT1 gene (Gene Designer® DNA2.0) based on the inferred amino acid sequence of PVX_083260 gene sequence in Plasmodb (http://www.plasmodb.org/plasmo/) (DNA sequence available on request). The construct contained a 5’ BamHI restriction site, the 1251 bp codon-optimized gene, and a 3’ EcoRI restriction site. The construct was sub-cloned into the yeast pCM189 plasmid (Gari et al., 1997) using BamHI/EcoRI restriction enzymes and T4 DNA ligase. The endogenous EcoRI site upstream of the tetracycline regulatable element in pCM189 was deleted using site-directed mutagenesis prior to cloning steps. The plasmid is a low copy number, tetracycline regulated promoter drives PvENT1 expression. The final construct generated was pCM189tetOff-PvENT1 (Gari et al., 1997). The empty vector (EV) construct pCM189-EV lacked only the transporter gene. All sequences were verified by DNA sequencing (Genewiz, Inc.).

PvENT1 SNP mutant constructs were generated using the QuikChange site-directed mutagenesis kit (Agilent Technologies) according to the manufacturer’s recommendations. Point mutations were verified by DNA sequencing.

2.3. Yeast transformation with PCR amplicon and pCM189 constructs

Yeast was transformed with plasmid or PCR DNA using the standard lithium-acetate/DMSO (8% v/v) method (Hill et al., 1991). For each transformation 10 mL of yeast culture was grown overnight in YPD media until the cell density reached ~2 × 10^6 cells/mL. The culture was pelleted at 3500 × g, 5 min (room temperature; RT) and the yeast pellet was washed once with cold lithium acetate buffer (LiOAc: 100 mM LiOAc, 10 mM Tris pH 8; using 2 PBS vol/culture vol). The final pellet was resuspended with 100 μL of LiOAc to which 10 μL of salmon-sperm DNA (2 mg/mL) and 0.5–1 μg of DNA (PCR or construct) was added. The solution was incubated at RT for 5 min 280 μL of 50% (w/v) PEG (polyethylene glycol 3350) was added. The PEG-LiOAc suspension was incubated at 30 °C for 45 min. DMSO was added to the mixture and the sample was heat shocked at 42 °C for 15 min. Transformed samples were pelleted at 16,300 × g, 30 s (RT) and resuspended in appropriate media before being transferred onto selection plates. For the creation of ade2∆::hphNT1 yeast, the final pellet was first resuspended in 10 mL YPD, allowed to recover for 3 h, pelleted again, and resuspended into a plating volume containing 500 μM hygromycin B. The final strains used in experiments had the following genotype: MATα; his3Δ1; leu2Δ0; met15α; ura3Δ0; flul1α::KanMX4, ade2Δ::hphNT1 containing either pCM189-EV or pCM189-PvENT1 constructs.

2.4. Yeast growth media and determination of cell density

Yeast cells were grown in YPD media: 1% (w/v) yeast extract, 2% (w/v) peptone, and 2% (w/v) glucose. Purine auxotrophic yeast were maintained in synthetic defined media (SDM) containing 2% (w/v) dextrose, 0.5% (w/v) ammonium sulfate, 0.17% (w/v) yeast nitrogen base (US Biologicals, #Y2030), 0.02% (w/v) nutritional dropout mix (US Biologicals, cat# D9542; lacking uracil, adenine, histidine, and tryptophan), 40 mg/L tryptophan, and 40 mg/L histidine. Supplemental adenine (300 μM) or adenosine (1 mM) was added for culture maintenance of the EV and PvENT1 expressing yeast, respectively. Solid growth media contained the same SDM formulation supplemented with either 1 mM adenine or 10 mM adenosine in 2% (w/v) agar.

2.5. Yeast growth and bioscreen assays

Purine auxotrophic yeast growth was assessed on both solid and liquid media where adenine or adenosine was the sole purine source. Briefly, the EV- and PvENT1-expressing yeast were grown in SDM-adenine media to mid-log phase, harvested at 3400 × g for 5 min (RT), and washed three times in sterile water. The final yeast cell pellet was diluted to 4 × 10^5 cells/mL. 96-well TC-treated plates were preloaded with 100 μL of purine serially diluted in 2 × SDM lacking purine. 100 μL of cells were added to the plates and resuspended three times. The cells were incubated at 30 °C and evaluated for growth (OD_{600} after 17–21 h. Biological replicates (n ≥ 4) were done on different days. For growth on solid media, yeast were grown to a density of 10^6 cells/mL serially diluted in water (1:5), inoculated onto SDM-agar plates (3000 cells/spot) and incubated at 30 °C for 2–3 days.

Bioscreen growth curve analysis for PvENT1 WT and SNP mutants was done using a Bioscreen C machine (Growth Curves USA, Piscataway, NJ). Briefly, cultures were grown in the appropriate yeast media to mid-log phase, washed as previously noted, and then diluted to ~2 × 10^6 cells/mL (OD_{600} = 0.018) either in 1 mM adenine or 10 mM adenosine media. Three aliquots of 150 μL culture (per strain/per condition) were added to a 100-well plate. Growth at 30 °C was monitored every 15 min at OD_{600} for 72 h.

2.6. Inhibition of yeast growth with small molecule compounds

We evaluated the viability of purine auxotrophic yeast grown in the presence of various inhibitors. Briefly, 80 μL of exponentially growing PfENT1-expressing yeast (~400,000 cells) were added to each well of a 384-well microplate (black, clear/flat-bottom, polystyrene; Corning #3712). The cells were grown in 1 mM adenosine—this concentration was picked based on growth that allowed maximum signal detection without saturating the culture/machine detection. Compound was serially diluted 3- or 4-fold in DMSO, and 0.8 μL was added to each well and resuspended three times. The final DMSO concentration was ~1% (v/v). The plates were incubated at 30 °C for 15–17 h. Growth was evaluated at OD_{600}, values were normalized to maximum growth with DMSO only and maximum death values for each compound. The normalized data was fit to a variable slope concentration-response model with no constraints to determine the concentration that gave half-maximal growth inhibition (IC_{50}) using Prism 6.02 software (GraphPad). Biological replicates (n ≥ 4) were done on different days.

2.7. Radiolabel uptake time course experiments

PvENT1-expressing yeast were grown to mid-log phase in SDM + 1 mM adenosine overnight at 30 °C. Cells were harvested by centrifugation at 3500 × g for 1 min, RT. Cells were washed three times in yeast PBS (150 mM NaCl, 10 mM KH_{2}PO_{4}, 40 mM K_{2}HPO_{4}, 11 mM glucose, pH 7.4). The yeast pellet was resuspended in PBS to a final concentration of 2 × 10^6 cells/mL. 96-well microplate wells

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were loaded with 100 μl of 100 nM \(^{[3]H}\)adenosine (2,8-\(^{3}H\)adenosine; 35 Ci/mmol) or 500 nM \(^{[3]H}\)uridine (2,8-\(^{3}H\)uridine; 8.5 Ci/mmol). All radiolabels were purchased from Moravek Biochemicals. 100 μl of yeast cell suspension was added to the radiolabels at the appropriate times to generate the time-course: 0, 0.5, 1, 2.5, 5, 10, 15, 20, 30, 45, and 60 min. At the conclusion of the experiment, cells were harvested onto glass fiber filters (Filtermat A, GF/C; Perkin Elmer) using a TomTec 96-well cell harvester system (#96-3-469). Filters were dried, sealed in plastic bags containing Betaplate Scint LSC fluid (Perkin Elmer). Samples were counted using a microplate scintillation counter (1450 MicroBeta Trilux, Perkin Elmer). Total counts are represented as CPM/10^6 cells (mean ± SD) from n ≥ 3 independent experiments. The data was fit using a linear-regression model with no constraints (Prism 6.02).

2.8. Radiolabel-uptake inhibition by purines and pyrimidines

PvENT1-expressing yeast were grown and prepared as described above. To measure the concentration dependent inhibition of radiolabel uptake, 96-well plates were preloaded with 50 μl of 200 nM \(^{[3]H}\)adenosine for pyrimidines or 1 μM \(^{[3]H}\)uridine for purines in PBS. 50 μl of purine or pyrimidine (serially diluted 3- or 4-fold in buffer) was added to each well and resuspended. 100 μl yeast (2 × 10^8 cells/mL) were added to each well and incubated at RT for 15 min. At the end of each experiment, cells were harvested and radioactivity was counted. Biological replicates (n ≥ 3) were done on different days.

2.9. \(^{[3]H}\)adenosine uptake inhibition by compounds

Compounds identified as PfENT1 inhibitors were purchased from Chembridge. Validation of purchased compounds by mass spectrometry and NMR were published previously (Frame et al., 2015b). To measure the concentration dependence of compound inhibition of \(^{[3]H}\)adenosine uptake, 96-well plates were preloaded with 100 μl of 100 nM \(^{[3]H}\)adenosine in PBS. 0.5 μl of compound (serially diluted 3- or 4-fold in DMSO, as described above) was added to each well and resuspended. 100 μl yeast (2 × 10^8 cells/mL) were added to each well and incubated at RT for 15 min. At the end of each experiment, cells were harvested and radioactivity was counted. Biological replicates (n ≥ 3) were done on different days.

3. Results

3.1. Expression of P. vivax ENT1 in yeast

The amino acid sequences of PvENT1 and PfENT1 are 75% identical. Since PfENT1 is a low affinity, broad-spectrum transporter of nucleosides and nucleobases, we hypothesized that PvENT1 would also recognize and transport these substrates. To study PvENT1, we expressed PvENT1 in S. cerevisiae yeast. We used a yeast codon-optimized version of the PvENT1 gene because species-specific differences in codon usage can affect gene expression levels (codon usage bias) (Hershberg and Petrov, 2009; Downie et al., 2010; Frame et al., 2012, 2015b). We used purine auxotrophic S. cerevisiae generated by deletion of the ADE2 gene, a critical enzyme in the yeast’s de novo purine synthetic pathway. Purine auxotrophic ade2Δ yeast grow poorly when purine starved and display a characteristic red color phenotype (Dorfan, 1969; Kokina et al., 2014). Adenine can enter ade2Δ yeast cells via the endogenous FCY2 nucleobase transporter and rescue growth (Schmidt et al., 1984). Yeast do not encode an endogenous adenosine transporter. Thus, growth of the purine auxotrophic yeast with adenosine as the sole purine source can only occur if adenosine can enter the cell through the PvENT1 transporter.

Yeast transformed with either an empty vector (EV) or PvENT1 construct displayed similar concentration-dependent growth in liquid culture media containing adenosine as the sole purine source (Fig. 1A) (EC50: 39 ± 2 μM EV vs. 27 ± 2 μM PvENT1). In contrast, PvENT1-expressing yeast grew in adenosine media (growth EC50: 394 ± 85 μM), while EV yeast failed to grow (Fig. 1A). Similar growth phenotypes were observed for the yeast cells grown on solid media (data not shown). Growth rescue with adenosine as the
sole purine source demonstrated that the yeast cells produced functional PvENT1 protein. We assume that the PvENT1 protein trafficked to the yeast plasma membrane but we did not explicitly demonstrate this.

Next, we examined whether PvENT1-expressing yeast could transport two radiolabeled substrates, adenosine and uridine. PvENT1-expressing yeast cells displayed time-dependent, linear accumulation of both [3H]adenosine and [3H]uridine (Fig. 1B and C, circles). As expected, EV transformed yeast not expressing PvENT1 did not accumulate either [3H]adenosine or [3H]uridine (Fig. 1B and C, squares). Since yeast lack an endogenous adenosine transporter and the endogenous FUT1 uridine transporter was deleted (fut1Δ), uptake of both radiolabeled substrates is only possible through PvENT1. The uptake of both purines and pyrimidines is consistent with the broad substrate specificity of the transporter.

3.2. PvENT1 mediated transport of purines and pyrimidines

We investigated whether other purines and pyrimidines could affect PvENT1-mediated uptake of radiolabeled adenosine or uridine. To avoid the potential for competition between the test substrate and the radiolabeled substrate at cytoplasmic metabolic enzymes (Kirk et al., 2009; Riegelhaupt et al., 2010a), we used the inhibition of [3H]uridine uptake to characterize purine uptake and inhibition of [3H]adenosine to characterize pyrimidine uptake. Thus, the radiolabel is metabolized by different cytoplasmic enzymes than the competing cold substrate. The only potential site of competition between the test substance and radioactive substrate is at the transporter. For these experiments, we assume that inhibition of radiolabel uptake is due to the competitive transport of the purine or pyrimidine test substance by the transporter. Unfortunately, due to financial constraints we did not have radiolabeled versions of all of the test substances to demonstrate that they were all transported into the yeast via PvENT1.

We quantified the observed effects as the IC50 of the test substance to inhibit the radiolabeled substrate uptake into the yeast. It should be noted that for these experiments, the ratio of the tritiated substrate concentration used in the uptake competition experiments to the test substance Km was between 10−3 and 10−4. Thus, by the Cheng–Prusoff equation the IC50 ≈ Km where Km is the inhibition constant of the test substance for the transporter (Cheng and Prusoff, 1973).

We evaluated 11 purines for their ability to inhibit [3H]uridine uptake into PvENT1-expressing yeast. These included nucleosides, nucleotides, and pyrimidines. Experiments were conducted at 15 min, which was within the linear phase of radiolabel uptake (Fig. 1C). Fig. 2A illustrates the experimental data for determination of IC50 values for the inhibition of radiolabel uptake by the purines inosine, hypoxanthine and xanthine. In this case, similar results were obtained whether the radiolabel substrate was a purine (adenosine) or a pyrimidine (uridine) implying that in both cases the inhibition of radiolabel uptake is due to competition at the PvENT1 transporter and not at a cytoplasmic metabolic enzyme. IC50 values for all purines tested are shown in Table 1. The nucleosides inosine and guanosine had the lowest IC50 values, in the low µM range, while their nucleotides were 50 and 300 times higher, respectively (Table 1). Adenosine had an IC50 value of 142 µM while the IC50 for its nucleobase adenine was one order of magnitude higher (Table 1).

Similarly, we evaluated 11 pyrimidines for their ability to inhibit the uptake of 50 nM [3H]adenosine. Fig. 2B illustrates the experimental data for determination of IC50 values for the inhibition of [3H]adenosine uptake into PvENT1-expressing yeast by members of the uracil series of pyrimidines. IC50 values for all pyrimidines tested are shown in Table 1. The nucleoside thymidine had the lowest IC50 of all pyrimidines tested (183 µM) followed by its nucleobase thymine (564 µM) (Table 1). The nucleobase cytosine caused slight inhibition of [3H]adenosine uptake while the remaining members of the family had negligible effects. The uracil family of pyrimidines had IC50 values in the low mM range (Fig. 2B, Table 1). 5-Fluorouridine (5-Furd), a toxic uridine analog, had an IC50 value of 1.7 mM.

None of the purine or pyrimidine nucleotides tested inhibited the uptake of [3H]adenosine or [3H]uridine. This implies that nucleotides are not substrates for PvENT1. Overall, our data show that PvENT1 recognizes both purine and pyrimidine substrates in the form of nucleosides and nucleotides.

Substitution of choline chloride for NaCl in the [3H]adenosine uptake media had a minimal impact on 15 min [3H]adenosine uptake (data not shown). This indicated that purine uptake was not Na+ coupled in PvENT1.

3.3. Effect of hENT1 inhibitors and cytotoxic purines on PvENT1-mediated purine uptake

Mammalian ENTs are categorized based on their sensitivity to the inhibitor NBMPR (Griffiths et al., 1997; Baldwin et al., 2004). Human hENT1 is inhibited by sub-nanomolar NBMPR concentrations, whereas hENT2 requires µM-mM concentrations for complete inhibition. Similarly, hENT1 is inhibited by the coronary vasodilator dipyridamole in the 5–20 nM concentration range (Griffiths et al., 1997; Baldwin et al., 2004). NBMPR did not inhibit [3H]adenosine uptake into PvENT1-expressing yeast at concentrations up to 20 mM. Dipyridamole inhibited the transporter with an IC50 ~ 40 µM, about three orders of magnitude higher than the IC50 for its nucleobase adenine was one order of magnitude lower than the IC50 for its nucleobase adenine.
for hENT1 inhibition (Table 2).

To explore determinants of purine recognition, we also examined the ability of two purine analogs to inhibit uptake of \([3H]\)uridine. The immunosuppressive drug 6-mercaptopurine (6-MP) had comparable efficacy in hENT1 inhibition (Table 2).

### 3.4. Effect of small molecule PfENT1 inhibitors on PvENT1

Using a yeast-based high throughput screen (HTS), we recently identified small molecule inhibitors of PfENT1 (Frame et al., 2015b). Of the 171 hits identified, we validated the efficacy of nine of the highest activity compounds, representing six different chemotypes (Fig. 3A) using a series of yeast- and parasite-based assays. In the current work, we explored the ability of these nine inhibitors to block PvENT1 using two different assays: 1) inhibition of \([3H]\)adenosine uptake into PvENT1-expressing yeast (PvENT1-expressing yeast), and 2) inhibition of the growth of PvENT1-expressing purine auxotrophic yeast with adenosine as the sole purine source.

### 3.5. Effect of PfENT1 non-synonymous SNPs on purine IC50 values and sensitivity to inhibitors

The substrate and inhibitor profiles described above are for the PfENT1 amino acid sequence encoded by the gene from the El Salvador I isolate (Carlton et al., 2008). To date, genomic sequencing of 170 P. vivax field isolates has identified six unique non-synonymous SNPs in the PfENT1 gene (PvENT1). Two of the SNPs, M99I and Q367K, always appear together (http://www.plasmodb.org/plasmo/showRecord.do?name=GeneRecordClasses.GeneRecordClass&project_id=PlasmoDB&source_id=PVX_083260, accessed 6/30/15). The SNPs, strain and protein locations are: D23E (Thailand VKTS-52, N-terminus), M99I/Q367K double mutant (Brazil strain; TM 3 and TM10-11 loop), Q178K (Columbia_30103103280, TM5-6 loop), L188M (PNG58 and others, TM6), and N329S (N. Korea strain; TM 9). We evaluated the functional significance of the PfENT1 polymorphisms using the yeast expression system.

Yeast expressing WT and each of the SNP mutant PfENT1s grew at a similar rate with adenosine as the sole purine source (Fig. 4A left panel and B). This was expected because the yeast express an endogenous adenosine transporter. With adenosine as the sole purine source, the M99I/Q367K double mutant PfENT1-expressing yeast grew at rates comparable to WT (Fig. 4A right panel and C). The D23E, Q178K, and L188M mutants also had comparable growth rates (data not shown). This implied that these yeast all expressed PfENT1 to a similar extent and with similar transport capacity. The Q178K and L188M mutations increased the hypoxanthine IC50 values by 6- to 26-fold. In contrast, none of the adenosine IC50 values were significantly different than WT, although the Q178K IC50 value was 3 times higher (Table 4).

The growth behavior of yeast expressing the N329S mutation was distinctly different from WT and the other SNPs. The N329S yeast grew on adenosine as the sole purine source implying that the purine salvage pathway was intact. In contrast, with adenosine as the sole purine source, the N329S-PfENT1-expressing yeast behaved similar to the EV-transformed yeast—the growth rate was minimal even in the presence of 10 mM adenosine (Fig. 4C). This phenotype was observed with N329S-PfENT1-expressing yeast from several independently transformed colonies and from two independent transformations. However, if the N329S-expressing yeast were
grown in media with adenine as the purine source, we could measure [3H]adenosine uptake. While the IC50 values for inhibition of [3H]adenosine uptake by three purines were similar to WT PvENT1 (Table 4) the amount of uptake at 15 min was 5–10-fold lower than the amount of uptake into WT-expressing yeast. Thus, the N329S mutant transported [3H]adenosine, but either the rate of transport per transporter or the number of N329S transporters in the plasma membrane was lower than for WT. Differences in the media pH and composition in the uptake and growth assays did not affect N329S function. The [3H]adenosine uptake experiments were done at pH 7.4 whereas the growth media had a pH ~4.2. [3H]adenosine uptake in 15 min for both WT- and N329S-expressing yeast was comparable in PBS at pH 4.2 and 7.4, the absolute amounts were different for WT and N329S (n = 5, data not shown). Furthermore, the amount of [3H]adenosine taken up in 15 min was comparable whether the buffer was PBS or SDM media (n = 4, data not shown). Thus, neither a difference in the effect of media pH on the N329S mutant or the media composition explains the failure of N329S-expressing yeast to grow with adenosine as the sole purine source.

We determined whether the PvENT1 non-synonymous SNP mutations altered the sensitivity to the PfENT1 inhibitor compounds (Fig. 3A). For most of the compounds, the PvENT1 SNPs had IC50 values comparable to WT PvENT1 in the [3H]adenosine uptake inhibition assays. For the D23E mutant, the IC50 values for compounds 3, 6 and 7 were ~3.5 fold higher than WT (p < 0.05, data not shown), but the mutant was more sensitive than WT to compounds 1 and 4. The N329S mutant induced a 2.6 fold increase in the IC50 of compound 2 relative to WT. No SNP caused a similar effect across all nine compounds. The results demonstrated that the SNP mutations do not confer resistance to the majority of the PfENT1 inhibitors identified and characterized in our initial screen.

Table 2
IC50 Values for inhibition of [3H]Adenosine uptake by human hENT1 inhibitors and cytotoxic purine derivatives. Effect of hENT1 inhibitors (NBMPR, nitrobenzylthioinosine; dipyridamole) and toxic purine analogs, 6-mercaptopurine and tubercidin (7-deaza-adenosine) on uptake inhibition of 50 nM [3H]adenosine into PvENT1-expressing yeast. Inhibition was evaluated over a range of concentrations up to 20 mM. IC50 values are represented as the mean ± SD from n = 3 separate experiments. NE signifies no observable effect.

| Compound          | [3H]Adenosine uptake inhibition IC50 (µM) |
|-------------------|------------------------------------------|
| NBMPR             | 40 ± 21                                   |
| dipyridamole      | 158 ± 31                                  |
| 6-mercaptopurine   | NE                                       |
| Tubercidin        | 7 ± 1                                     |

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Despite 75% amino acid sequence identity between PfENT1 and PvENT1, PvENT1 displays functional differences from PfENT1 in terms of substrate interaction profiles. PvENT1 is more selective for purines than for pyrimidines. Compared to PfENT1, the IC_{50} values for inosine and guanosine were lower than for adenosine, a profile similar to the Leishmania donovani LdNT2 transporter (Carter et al., 2000b). Inosine had the lowest IC_{50} for inhibition of \[^{3}H\]uridine uptake, followed by guanosine and adenosine/hypoxanthine. Hypoxanthine and adenosine are endogenous purine substrates for PfENT1 (and for isolated P. falciparum parasites). PvENT1 had about 5-fold lower IC_{50} values for hypoxanthine and adenosine, ~100 \mu M and ~140 \mu M, respectively, than PfENT1, ~480 \mu M for hypoxanthine and ~650 \mu M for adenosine (Riegelhaupt et al., 2010a). These differences may relate to the fact that P. vivax parasites only infect reticulocytes, immature erythrocytes, whereas P. falciparum parasites infect mature erythrocytes as well as reticulocytes. The cytoplasmic environment and purine composition is different in immature vs mature erythrocytes (Srivastava et al., 2015). It remains to be determined if the difference is a consequence of their respective preference for immature versus mature erythrocytes. Furthermore, it is unknown whether the differences in purine IC_{50} values for PvENT1 versus PfENT1 are physiologically significant.

One of the important findings from this study was that the PfENT1 inhibitors that we identified previously are also potent inhibitors of PvENT1 with IC_{50} values in the low nM range. In the present study, we showed that the nine compounds inhibited \[^{3}H\] adenosine uptake into PvENT1-expressing yeast and inhibited

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### Table 3

Efficacy of PfENT1 inhibitors against PvENT1 using two assays of PfENT1 function. IC_{50} values for inhibition of \[^{3}H\] adenosine uptake into or growth of PfENT1-expressing yeast by nine PfENT1 inhibitors. IC_{50} values are represented as the mean ± SD from n > 3 separate experiments. Compound numbers based on Rank Order# in HTS (Frame et al., 2015b). Compound structures are in Fig. 3A. Data for IC_{50} values for compound efficacy against PfENT1 in the \[^{3}H\] adenosine uptake assay are shown for comparison.

| Compound# | PfENT1 IC_{50} for growth inhibition of purine auxotrophic yeast in adenosine containing media (nM) | PfENT1 IC_{50} for \[^{3}H\] adenosine uptake assay (nM) | PfENT1 IC_{50} for \[^{3}H\] adenosine uptake assay (nM) |
|-----------|-------------------------------------------------|-------------------------------------------------|-------------------------------------------------|
| 1         | 2.8 ± 0.8                                        | 3.0 ± 1.3                                        | 1.8 ± 0.8                                        |
| 2         | 22.4 ± 11.8                                      | 10 ± 8.4                                        | 8.4 ± 7.6                                        |
| 3         | 2.6 ± 0.9                                        | 2.4 ± 1.5                                        | 1.8 ± 1.2                                        |
| 4         | 32.3 ± 19.9                                      | 13.7 ± 7.5                                      | 12.7 ± 7.5                                      |
| 5         | 31.7 ± 7.6                                      | 22.7 ± 7.2                                      | 19.9 ± 13.7                                      |
| 6         | 9.5 ± 3.7                                        | 9.6 ± 7.4                                       | 9.6 ± 7.4                                       |
| 7         | 5.2 ± 2.6                                        | 10.9 ± 6.8                                      | 10.9 ± 6.8                                      |
| 13        | 7.5 ± 4.0                                        | 3.9 ± 1.6                                       | 3.9 ± 1.6                                       |
| 19        | 8.3 ± 0.9                                        | 38.4 ± 16.5                                     | 38.4 ± 16.5                                     |

* PfENT1 data from (Frame et al., 2015b).

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### 4. Discussion

Multidrug-resistant malaria parasites have become more common. Thus it is imperative to develop new antimalarial compounds that target novel aspects of parasite biology. Inhibition of the purine import pathway(s) is one potential target for the development of novel antimalarial drugs (Carter et al., 2001; Downie et al., 2006; El Bissati et al., 2006; Riegelhaupt et al., 2010a; Frame et al., 2015b). However, a major hurdle has been the difficulty in developing HTS assays to identify inhibitors of equilibrative nucleoside transporters. Recently, we described a yeast-based growth assay that we used to screen a 64,500 compound library. We identified inhibitors of PfENT1, the primary purine import pathway in P. falciparum parasites (Frame et al., 2015b). Nine of the highest affinity PfENT1 inhibitors, comprising six distinct chemotypes, kill P. falciparum parasites in culture (Frame et al., 2015b). Because of the considerable geographical overlap in the distributions of P. falciparum and P. vivax, to be effective, novel drugs should ideally target both species. In the present work we have characterized the P. vivax equilibrative nucleoside transporter homolog, PvENT1, in a yeast expression system. We show that it has a similar, but distinct, substrate profile compared to PfENT1. Furthermore, the PfENT1 inhibitors that we identified previously (Frame et al., 2015b) are highly efficacious inhibitors of PvENT1. This implies that it may be feasible to develop inhibitors of the primary malaria purine import transporter as potential novel antimalarial drugs that will be effective against both falciparum and vivax malaria.
auxotrophic yeast. We previously showed that these compounds are not toxic to yeast at concentrations up to hundreds of micromolar (Frame et al., 2015b). Thus, at the sub-micromolar concentrations where the compounds inhibit adenosine-dependent growth, it is likely due to inhibition of PvENT1-dependent adenosine uptake. The efficacy of the nine compounds in both assays is a function of the fundamental differences in the assays and does not represent a difference in the strength of the inhibitor’s interaction with PvENT1. The IC50 values in the growth experiments depend on the level of PvENT1 expression and thus, the amount of excess purine transport capacity relative to the purine requirements for yeast cell growth and proliferation. Thus, the IC50 values in the growth experiments do not directly reflect the affinity of the inhibitors for PvENT1, but rather the fraction of expressed PvENT1 that must be inhibited to make purine uptake limit yeast cell growth. In the inhibition of [3H]adenosine uptake assay, the measured IC50 values are a direct measure of interaction between the inhibitor and PvENT1.

Alternatively, the adenosine concentration is 1 mM in the growth assay and only 50 nM in the [3H]adenosine uptake assay, in the former case, ~10 times the adenosines. IC50 and in the latter ~2000-fold lower. The difference may represent competition between adenosine and the inhibitor for an overlapping binding site. Overall, our results provide strong evidence that the PfENT1 small molecule inhibitors also block PvENT1, in many cases with higher potency (Frame et al., 2015b). The ability of these nine compounds to inhibit both PfENT1 and PvENT1 provides further support for the hypothesis that malaria purine transporters are a reasonable target for the development of novel antimalarial drugs.

In contrast to the efficacy of the PfENT1 inhibitors against PvENT1, hENT1 inhibitors had little or no efficacy against PvENT1. PvENT1 was not sensitive to NBMPR, an inosine analog where the C6 oxygen is replaced with the more bulky S-nitrobenzyl moiety. This result is in agreement with the NBMPR-insensitivity observed for other protozoan ENTs (de Koning et al., 2005). Dipyridamole did inhibit PvENT1 with a 40 μM IC50 but this is over 1000-fold higher than its efficacy against hENT1. The Plasmodium homologs are only 17% sequence identical to hENT1 so this result was not unexpected. Since both hENT1 and the Plasmodium ENTs recognize purines with similar affinity, this suggests that dipyridamole binding affinity probably depends on residues in the transporter besides those involved in purine recognition.

We investigated the impact of known non-synonymous PvENT1 SNPs identified in sequenced field isolates. Growth of yeast expressing five of the six SNPs with adenosine as the sole purine source was similar to growth rates of yeast expressing WT PvENT1. We were surprised that yeast expressing the N329S-PvENT1 variant were unable to grow in adenosine, although they grew normally with adenine as the sole purine source, because the IC50 values for adenosine, inosine, and hypoxanthine inhibition of [3H]uridine uptake for N329S-expressing yeast were similar to the IC50 values obtained with WT-expressing yeast. We ruled out differences in the media pH or composition as potential effects. It is most likely due to a difference in N329S function or level of expression relative to WT. Further experiments will be necessary to explain these results.

Some naturally occurring PfENT1 SNPs have functional effects on purine uptake (e.g., F934L; ref. (Riegelhaupt et al., 2010b)) (Table 4). Given the small numbers of identified PfENT1 non-synonymous SNPs and that these SNPs have similar affinity for the identified PfENT1 inhibitors, currently existing SNPs do not represent a potential source of pre-existing resistance to the inhibitors.

Most protozoan species have de novo pyrimidine synthesis

### Table 4

| SNP         | Hypoxanthine IC50 (μM) | Inosine IC50 (μM) | Adenosine IC50 (μM) |
|-------------|------------------------|-------------------|---------------------|
| WT          | 141 ± 24               | 5 ± 4             | 167 ± 52            |
| D23E        | 150 ± 164              | 123 ± 113         | 94 ± 93             |
| M99I/Q367K  | 107 ± 13               | 6 ± 2             | 206 ± 28            |
| Q178K       | 530 ± 110              | 32 ± 5            | 600 ± 270           |
| L188M       | 1200 ± 780             | 130 ± 23          | 190 ± 120           |
| N329S       | 158 ± 18               | 10 ± 4            | 203 ± 31            |

**Table 4.** Purine IC50 values for inhibition of [3H]-substrate uptake by yeast expressing PvENT1 containing mutations identified in P. vivax field isolates with non-synonymous SNPs in the pvent1 gene. [3H]Hypoxanthine uptake was used to assess PvENT1 activity. *Indicates the radiolabel used was [3H]adenosine, not [3H]uridine. †Significantly different than WT (p < 0.05), Student t-test unequal variance. Values are mean ± SD.
pathways and do not incorporate host pyrimidines into DNA/RNA. Nevertheless, our experiments show that PvENT1 can bind to and probably transport various pyrimidine compounds. In the case of uridine, we showed that it is transported by PvENT1. Knowledge of the PvENT1 pyrimidine binding profile may be useful in designing drugs that target the de novo pyrimidine biosynthetic pathway and might use PvENT1 as a transport pathway for drug entry.

5. Conclusions

The parasite purine salvage pathway has been a target of various potent antimalarial compounds (e.g., deoxycoformycin, immucillin) (Webster et al., 1984; Kicikas et al., 2002; Ting et al., 2005). In this body of work, we have characterized the functional properties of PvENT1, a potential antimalarial target. While PfENT1 and PvENT1 share a high degree of amino acid sequence identity, differences in amino acids in PvENT1 result in altered affinities for purine substrates. Nonetheless, PvENT1 (and the known non- synonymous SNP variants) are inhibited by a new class of small molecule inhibitors that target the P. falciparum purine importer PfENT1. Development of these inhibitors as novel antimalarial drugs may provide an additional arsenal of antimalarial compounds that have not overlap with current chemotherapeutics.

Conflicts of interest

The authors declare the following potential competing interest(s): A patent application is pending on the yeast-based assay as a method to identify nucleoside transporter inhibitors. A second pending patent covers the use of the 171 compounds as PfENT1 inhibitors and as novel antimalarial drugs. The patents have not been licensed to date and no negotiations are currently in progress.

Author contributions

RD, IJF and MHA designed the experiments. RD and YS performed the experiments. All authors analyzed data and participated in writing and/or editing the manuscript.

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