The Prenylated Proteome of *Plasmodium falciparum* Reveals Pathogen-specific Prenylation Activity and Drug Mechanism-of-action*

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*Plasmodium* parasites contain several unique membrane compartments in which prenylated proteins may play important roles in pathogenesis. Protein prenylation has also been proposed as an antimalarial drug target because farnesyltransferase inhibitors cause potent growth inhibition of blood-stage *Plasmodium*. However, the specific prenylated proteins that mediate antimalarial activity have yet to be identified. Given the potential for new parasite biology and elucidating drug mechanism-of-action, we performed a large-scale identification of the prenylated proteome in blood-stage *P. falciparum* parasites using an alkyne-labeled prenyl analog to specifically enrich parasite prenylated proteins. Twenty high-confidence candidates were identified, including several examples of pathogen-specific prenylation activity. One unique parasite prenylated protein was FYVE-containing coiled-coil protein (FCP), which is only conserved in *Plasmodium* and related Apicomplexan parasites and localizes to the parasite food vacuole. Targeting of FCP to this parasite-specific compartment was dependent on prenylation of its CaaX motif, as mutation of the prenylation site caused cytosolic mislocalization. We also showed that PRab5b, which lacks C-terminal cysteines that are the only known site of Rab GTPase modification, is prenylated. Finally, we show that the THQ class of farnesyltransferase inhibitors abolishes FCP prenylation and causes its mislocalization, providing the first demonstration of a specific prenylated protein disrupted by antimalarial farnesyltransferase inhibitors. Altogether, these findings identify prenylated proteins that reveal unique parasite biology and are useful for evaluating prenyltransferase inhibitors for antimalarial drug development. 

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Protein prenylation is the covalent attachment of hydrophobic prenyl chains to cysteine residues of protein substrates. This post-translational lipid modification often mediates membrane association important for the cellular function of prenylated proteins. During symptomatic infection, *Plasmodium* parasites reside inside mature red blood cells. Although the uninfected red blood cell lacks nucleus, mitochondria, and ER/Golgi and consists only of a cell membrane, *Plasmodium*-infected red blood cells contain numerous unique membranes both present within the parasite and induced by the parasite in the host cell. These include a specialized parasite organelle called the food vacuole in which host hemoglobin is digested, a parasitophorous vacuole that surrounds the parasite, and new membrane structures in the red cell cytoplasm. *Plasmodium* proteins modified with palmitoyl and/or myristoyl lipids have important functions at several of these new membranes (1, 2). It seems likely that prenylated *Plasmodium* proteins will also play critical roles within these unique parasite membranes. Protein prenylation is catalyzed by three classes of protein prenyltransferases: farnesyltransferase (FT)1, geranylgeranyltransferase 1 (GGT1), and Rab geranylgeranyltransferase (RabGGT). These enzymes differ in the addition of either 15-carbon farnesyl or 20-carbon geranylgeranyl chains and their recognition of substrate proteins. Both FT and GGT1 recognize and modify a 4-amino acid Cysteine-aliphatic-aliphatic-terminal residue (CaaX) motif at the C-terminus of protein targets, whereas RabGGT exclusively recognizes a family of Rab GTPases and modifies two C-terminal cysteines. *Plasmodium* prenylases encode at least a putative FT and RabGGT. In *P. falciparum*, two CaaX proteins which share homology with known prenylated mammalian proteins were shown to be prenylated, and 11 Rab GTPases are annotated in the genome (3–6). However, large-scale, unbiased approaches will

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1 The abbreviations used are: FT, farnesyltransferase; GGT1, geranylgeranyltransferase; RabGGT, Rab geranylgeranyltransferase; CaaX, cysteine-aliphatic-aliphatic-terminal residue residue; THQ, tetrahydroquinoiline; AlkFOH, alkyne-modified farnesyl probe; CuAAC or click chemistry, copper-catalyzed alkyne-azole cycloaddition; PI3P, phosphatidylinositol 3-monophosphate.
be required to accelerate the identification of nonconserved prenylated parasite proteins that have unique functions in pathogenesis.

Given the diverse and often essential functions of prenylated proteins, protein prenylation is also a target for therapeutic intervention. Inhibitors of human FT and GGT1 have undergone extensive investigation for treatment of numerous cancers, the premature aging syndrome progeria, and hepatitis delta infection, resulting in multiple clinical trials (7). The mechanism-of-action of these inhibitors and their efficacy against specific diseases is highly dependent on the cellular functions of specific CaaX protein substrates. The prenylation of oncogenic Ras GTPases by FT and GGT1 is the desired target in cancer treatment, whereas disrupting the farnesylation of nuclear lamins underlies the efficacy of FT inhibitors in treatment of progeria.

To combat growing antimalarial drug resistance, FT inhibitors have been proposed as a new class of antimalarials based on several important attributes. First, these compounds potently inhibit the growth of blood-stage \textit{Plasmodium} parasites (8, 9). Second, they show minimal to no effects on mammalian cell growth at the therapeutic doses required for parasite growth inhibition (8). Finally, the drug development process benefits enormously from the vast medicinal chemistry and biological knowledge accrued over decades of pharmaceutical research on FT inhibitors for cancer chemotherapy. The tetrahydroquinoline (THQ) series of FT inhibitors, which show low \textit{nm} efficacy against blood-stage \emph{Plasmodium falciparum}, have been investigated most extensively as antimalarials. Studies on their mechanism-of-action showed that THQ compounds block farnesylation activity in \emph{Plasmodium} cell lysates and selectively inhibit farnesylation of unidentified cellular proteins (8). Notably, resistance to THQ compounds was associated with mutations in the parasite’s putative FT \( \beta \) subunit (10, 11). These studies validated \emph{P. falciparum} FT as the drug target but left unanswered questions about the specific CaaX protein(s) that mediate parasite growth inhibition.

To perform the first unbiased and large-scale identification of the prenylated proteome in \emph{P. falciparum}, we took a chemical biology approach using metabolic labeling with an alkyn-modified farnesyl probe (AlkFOH) combined with mass spectrometry proteomics. Key questions we sought to answer were: What are the protein substrates of prenyltransferrases in \emph{Plasmodium}? How is protein prenylation in the parasite distinct from that in mammalian cells? Which prenylated proteins are important for mediating the antimalarial activity of FT inhibitors? Though the prenylated proteome is significantly reduced in \emph{P. falciparum} compared with humans, we provide evidence for parasite-specific prenylation activity, including a novel prenylated protein and a novel site of modification. Furthermore, we use one of these newly identified prenylated proteins to investigate the mechanism-of-action of antimalarial farnesyltransferase inhibitors.

**EXPERIMENTAL PROCEDURES**

\emph{P. falciparum In Vitro Culture}—All experiments described were performed in W2 (MRA-157) and Dd2\textsuperscript{aap} (MRA-843) strain parasites, or with transgenic parasites generated in this study. Parasites were grown in human erythrocytes in RPMI 1640 media supplemented with 0.25% Albumax II (Gibco Life Technologies; Carlsbad, CA), 2 g/L sodium bicarbonate, 0.1 mm hypoxanthine, 25 mm HEPES (pH 7.4), and 50 \( \mu \)g/L gentamycin, at 37 °C, 5% \( \text{O}_2 \), and 5% \( \text{CO}_2 \). Whole blood was obtained from the Stanford Blood Center. For comparison of growth between different treatment conditions, cultures were carried simultaneously and handled identically with respect to media changes and addition of blood cells.

**Metabolic Labeling and Cu\textsuperscript{1+} Catalyzed Alkyn-Azide Cycloaddition**—Early trophozoite parasites were labeled for 6 h with 1 \( \mu \)M AlkFOH, unless otherwise noted. After labeling, parasites were washed and saponin pellets were prepared. Pellets were stored at −80°C. AlkFOH incorporation was monitored by in-gel fluorescence. Parasite pellets were lysed in 4% SDS, 50 mm TEA pH 7.4, and 150 mm NaCl. Lysate was immediately reacted with AzTAMRA (Life Technologies; Carlsbad, CA) under Cu\textsuperscript{1+} catalyzed alkyn-azide cycloaddition (CuAAC) conditions (12). Final concentrations were 1 mm TCEP (made fresh), 1 mm CuSO\(_4\), 100 \( \mu \)M BTBA, and 100 \( \mu \)M azide. The reaction was incubated for 1 h at room temperature. Protein was precipitated using methanol/chloroform/water (4/1/5.3 v/v/v) and washed three times with 1 ml cold methanol.

**Mass Spectrometry Sample Preparation**—Samples for mass spectrometry were generated by labeling 200 ml of 4% HCT, 10% early trophozoites (per condition) for 6 h with 1 \( \mu \)M AlkFOH or mock treated with DMSO. Parasites grown for each condition were mixed and separated immediately prior to labeling. Saponin pellets were prepared and stored at −80°C. Parasite pellets were lysed in 4% SDS, 50 mm TEA pH 7.4, 150 mm NaCl with protease inhibitor minitablets (Pierce) and Benzonase nuclease (Sigma-Aldrich; St. Louis, MO). At least 3 mg/ml protein underwent CuAAC with AzBiotin (Click chemistry tools) for 5 h at room temperature. CuAAC conditions were the same as described above. Cell lysate was diluted to 1 mg/ml for the reaction. In order to help prevent SDS contamination, after the 5 h incubation, reactions were diluted 10 times with 50 mm TEOA pH 7.4, 150 mm NaCl and precipitated using methanol and chloroform (methanol/chloroform/water at ratios of 4/1/5.3 v/v/v) overnight at −20 °C. Samples were centrifuged and washed 3 times with cold methanol. Protein pellets were dried for 1 h then solubilized in 1 ml 1% SDS, 50 mm TEOA pH 7.4, 150 mm NaCl using bath sonication. Samples were incubated with 100 \( \mu \)l high capacity streptavidin beads (Pierce) for 1 h, washed 3 times with 1% SDS, 50 mm TEOA pH 7.4, 150 mm NaCl. To help remove SDS, samples were washed 6 times with 6 \( \mu \)l urea in 50 mm TEOA, pH 7.4, 150 mm NaCl. Samples were then washed 3 times with 25 mm ammonium bicarbonate (ABC) then resuspended in 25 mm ABC. Samples were reduced using 10 \( \mu \)m DTT and alkylated using 500 \( \mu \)m iodoacetamide. Samples were digested using sequence-grade trypsin (Promega) overnight at 37 °C. Acetonitrile was added to the digestion buffer at a final concentration of 35%. The final solution containing the released peptides was removed from the beads and dried. Peptides were desalted using C18 stage tips and resuspended in 0.1% formic acid before being injected into the LC/MS-MS.

**Mass Spectrometry Experimental Design and Statistical Rationale**—Five biological replicates of each sample were analyzed by online capillary nanoLC-MS/MS. Samples were separated on an in-house made 20 cm reversed phase column (100 \( \mu \)m inner diameter, packed with ReproSil-Pur C18-AQ 3.0 \( \mu \)m resin (Dr. Maisch GmbH)).
the TAMRA channel. If further analysis by immunoblot was needed, protein was then transferred onto nitrocellulose membrane using Trans-Blot Turbo Transfer System (BioRad; Hercules, CA). The membrane was blocked for 1 h or longer at room temperature in 0.1% Casein (Casein, Hammersten Affymatrix) dissolved in 0.2× PBS. Membranes were probed overnight with 1:5000 Living Colors mouse monoclonal αGFP (Clonetech) or 1:2,000 rabbit polyclonal αAldolase (Abcam; Cambridge, UK) in 50% 0.1% Casein in 0.2× PBS and 50% 1×TBST. Secondary antibodies were probed for an hour or more at room temperature in the same buffer. Secondary antibodies used were either IRDye 680LT goat-α-mouse or IRDye 800CW donkey-α-rabbit when appropriate. Western blots were imaged using a LI-COR Odyssey Imaging System.

Immunoprecipitation. GFP-FCP, GFP-FCP(C322S) and Rab5b-GFP were purified from transgenic parasites using the GFP-TrapA immunoprecipitation kit (Chromotek; Planegg, Germany). Five ml cultures of mixed stage parasite between 2–5% parasitemia at 4%HCT were treated with BMS-386914 or lonafarnib for 1 h, then 1 μM AlkFOH was added to the media and parasites were labeled for 6 h. Saponin pellets were lysed in RIPA buffer with benzonase and protease inhibitor minitablets (Pierce) and incubated on ice for 20 min, with periodic vortexing. Whole cell lysates were diluted 5× with GFP-Trap dilution buffer and incubated with 25 μl of a 50% slurry of GFP-Trap agarose beads for 1 h at RT. Beads were washed three times in 500 μl wash buffer using a 27 G needle and syringe to remove any excess buffer. Protein was reacted with AzTAMRA on bead in 50 μl 1× PBS as described above. After a 1 h click reaction, beads and protein were precipitated using MeOH/ chloroform/water and washed twice with 1 ml MeOH to remove any excess AzTAMRA. Pellets were dried and resuspended in 2× NuPage running buffer, then analyzed by SDS-PAGE and Western blotting as described above.

Live Epifluorescence Microscopy—Transgenic parasites were stained with 2 μg/ml Hoechst 33342. Epifluorescent images were collected using a Nikon Eclipse Ti-E using μManager software for acquisition. Images were processed using ImageJ. Single z-stack images were collected. Images were scored as showing either cytosolic localization, food vacuole localization, or a localization that had a weak cytosolic signal with brighter rings surrounding the food vacuole, which we termed "other." Examples of scored images are shown in supplemental Fig. S2.

RESULTS

An Alkyny-modified Prenyl Lipid Analog Specifically Labels Parasite Proteins—To enable large-scale identification of prenylated proteins, we performed metabolic labeling with an alkyny-modified farnesol analog, AlkFOH, in Plasmodium-infected red blood cells. Previously AlkFOH has been successfully used to identify prenylated proteins in mammalian cells (18). Mature red blood cells lack prenyltransferase activity, therefore incorporation of AlkFOH into proteins is due exclusively to the activity of protein prenyltransferases expressed by the infecting parasite (19). P. falciparum parasites were cultured with varying amounts of AlkFOH to determine its effect on parasite growth. Parasites could tolerate up to 10 μM AlkFOH with minimal growth inhibition (Fig. 1A). Parasites treated with 1 μM AlkFOH across the entire parasite replication cycle showed no growth defect nor any other observable phenotype, and 1 μM AlkFOH was sufficient to robustly label proteins in the parasite lysate (Fig. 1B and 1C). The most prominent bands incorporating the AlkFOH label were ob-
served at ~25 kDa and correspond to the expected size of Rab GTPases. Incorporation of AlkFOH into parasite proteins is specific, as protein labeling was competed away with increasing amounts of geranylgeraniol in vivo (Fig. 1C). Thus, this alkyne-modified prenyl lipid analog is a suitable tool for proteomic identification of prenylated proteins in blood-stage P. falciparum parasites.

The Prenylated Proteome Reveals a Limited Number of Prenylated Targets—Having successfully introduced an alkyne label into prenylated proteins, we used Cu1-catalyzed cycloaddition (click chemistry) to append an azido-biotin tag, affinity purified the labeled proteins, and identified them by mass spectrometry. Labeling began during the trophozoite stage when the protein prenyltransferases are maximally expressed (20). Five biological replicates were performed, and protein abundance was quantified by the total MS1 area of all matched peptides. The average abundance of proteins in AlkFOH- versus mock-labeled samples was plotted against average MS1 area measured for the same protein in DMSO mock-treated samples. Proteins which were detected only in AlkFOH labeled samples are shown on the left of the graph (ND = not detected). All data points are shown. Rab GTPases are marked in green, whereas proteins with a C-terminal CaaX box are shown in magenta. CaaX proteins which have previously been shown to be prenylated are designated with an asterisk. Results are based on five biological replicates.

**Fig. 1.** AlkFOH specifically labels prenylated proteins in *P. falciparum*. A, *P. falciparum* growth was measured in the presence of varying concentrations of AlkFOH for 72 h. Growth relative to parasites treated with DMSO was plotted against AlkFOH concentration (μM, log scale, based on 3 biological replicates). B, Growth of parasites, treated with either 1 μM AlkFOH or DMSO, were monitored over the course of two replication cycles. Growth relative to parasites treated with DMSO was plotted against the number of reinvasion cycles (each cycle is 48 h; based on 3 biological replicates). C, Early trophozoites were labeled with 1 μM AlkFOH for 6 h with 0, 1, 10, or 100 μM GGOH or treated only with DMSO. Whole cell lysates were reacted with AzTAMRA under CuAAC conditions and resolved by SDS-PAGE, and in-gel fluorescence was measured.

**Fig. 2.** Proteomic identification of prenylated proteins in *P. falciparum* reveals a limited number of substrates. Average MS1 area measured in AlkFOH-labeled samples is plotted against average MS1 area measured for the same protein in DMSO mock-treated samples. Proteins which were detected only in AlkFOH labeled samples are shown on the left of the graph (ND = not detected). All data points are shown. Rab GTPases are marked in green, whereas proteins with a C-terminal CaaX box are shown in magenta. CaaX proteins which have previously been shown to be prenylated are designated with an asterisk. Results are based on five biological replicates.
The Prenylated Proteome of *Plasmodium*

**TABLE I**

| CaaX box proteins | Annotation | Motif | PrePS prediction | PRENbase cluster |
|-------------------|------------|-------|------------------|------------------|
| PF3D7_0322000     | peptidyl-prolyl cis-trans isomerase (CYP19A) | CGEL | FT– GGT1– | Predicted conserved |
| PF3D7_0910600     | SNARE protein, putative (YKT6.1) | CCSIM | FT+++ GGT1+++ | Known conserved |
| PF3D7_1113100     | protein tyrosine phosphatase (PRL) | CHFM | FT– GGT1– | Known conserved |
| PF3D7_1319100     | Conserved DUF544 protein, unknown function | CTIM | FT+++ GGT1+++ | Predicted conserved |
| PF3D7_1324700     | SNARE protein, putative (YKT6.2) | CCSLY | FT+ GGT1– | Known conserved |
| PF3D7_1428700     | Conserved protein, unknown function | CNFM | FT– GGT1– | Unknown Not conserved |
| PF3D7_1437900     | HSP40, subfamily A, putative (ERdj3) | CAQQ | FT+ GGT1– | Known conserved |
| PF3D7_1460100     | FYVE and coiled-coil domain-containing protein (FCP) | CNIM | FT+ GGT1++ | Unknown Not conserved |

**Rab GTPases**

| Gene ID | Annotation | Motif | PrePS prediction |
|---------|------------|-------|------------------|
| PF3D7_0106800 | Ras related protein, Rab5c | KKCC | RabGGT+++ | Known conserved |
| PF3D7_0211200 | Ras related protein, Rab5a | KGCC | RabGGT+++ | Known conserved |
| PF3D7_0512600 | Ras related protein, Rab1b | KKCC | RabGGT+++ | Known conserved |
| PF3D7_0807300 | Ras related protein, Rab18 | NCAC | RabGGT+++ | Known conserved |
| PF3D7_0903200 | Ras related protein, Rab7 | SRCC | RabGGT+++ | Known conserved |
| PF3D7_1144900 | Ras related protein, Rab6 | KCLC | RabGGT+++ | Known conserved |
| PF3D7_1231100 | Ras related protein, Rab2 | FSCC | RabGGT+++ | Known conserved |
| PF3D7_0513800 | Ras related protein, Rab1a | FCSC | RabGGT+++ | Known conserved |
| PF3D7_1320600 | Ras related protein, Rab11a | NKCC | RabGGT+++ | Known conserved |
| PF3D7_1340700 | Ras related protein, Rab11b | VKCC | RabGGT++ | Known conserved |
| PF3D7_1310600 | Ras related protein, Rab5b | Noneb | RabGGT– | Known conserved |

**H. sapiens Rab GTPase**

| Gene ID | Annotation | Motif | PrePS prediction |
|---------|------------|-------|------------------|
| Q15286 | Ras related protein, Rab35 | KRCC | GGT2+++ | Known conserved |

*a* High confidence proteins were defined by (1) homology to known prenylation substrates, (2) presence of a motif, (3) present in three or more biological replicates, and (4) detected only in AlkFOH labeled samples.

*b* No cysteine residues found in last 5 amino acids of predicted protein sequence.

riched in the AlkFOH-labeled samples, suggesting that our method can specifically identify prenylated proteins.

We used both proteomic and biological criteria to segregate the list of identified proteins. Medium-confidence hits were selected if they satisfied criteria based on proteomic results: (1) identified by >1 unique peptide, (2) present in 3 out of 5 biological replicates, and (3) detected in AlkFOH-labeled samples and absent in DMSO controls. The full list of medium-confidence prenylated proteins are listed in supplemental Table S1. High-confidence prenylated proteins also fulfilled biological criteria, either containing cysteine residues within four amino acids of the C-terminal or had homology to known prenylated proteins. These criteria were based on established features of substrate recognition by the three classes of protein prenyltransferases.

A total of 20 high-confidence prenylated proteins were identified (Table I). We assigned the proteins to “prenylated protein clusters” identified in PRENbase, a database of known and predicted protein prenylation substrates (19). Each cluster is comprised of protein homologs with known or predicted prenylation motifs that are evolutionarily-conserved and likely functionally related. Thus, prenylated parasite proteins that cannot be assigned to a PRENbase cluster are more likely to have new, pathogen-specific functions that have not been described previously. A majority of the identified proteins clustered with homologs for which prenylation is both validated and highly conserved among eukaryotes, including 10 Rab GTPases and 4 CaaX proteins. Two candidates, a DUF544 protein of unknown function (PF3D7_1319100) and a peptidyl-prolyl cis-trans isomerase (PF3D7_0322000), are conserved eukaryotic proteins containing CaaX motifs whose prenylation are predicted but have not been previously validated in any organism (21). Significantly, two prenylated proteins could not be assigned to a PRENbase cluster: a FYVE and coiled-coil domain-containing protein, FCP (PF3D7_1460100) and a peptidyl-prolyl cis-trans isomerase (PF3D7_0322000), which lacks any C-terminal cysteines and human Rab35, a host red blood cell protein.

**Localization of a Novel Parasite Protein FCP is Dependent on Prenylation**—In our proteomic dataset, FCP stood out as a novel, pathogen-specific protein that has not been previously identified in any organism. FCP was identified by both proteomic and biological criteria and was shown to be present in three or more biological replicates and absent in DMSO controls. A majority of the identified proteins clustered with homologs for which prenylation is both validated and highly conserved among eukaryotes, including 10 Rab GTPases and 4 CaaX proteins. Two candidates, a DUF544 protein of unknown function (PF3D7_1319100) and a peptidyl-prolyl cis-trans isomerase (PF3D7_0322000), are conserved eukaryotic proteins containing CaaX motifs whose prenylation are predicted but have not been previously validated in any organism (21). Significantly, two prenylated proteins could not be assigned to a PRENbase cluster: a FYVE and coiled-coil domain-containing protein, FCP (PF3D7_1460100) and a peptidyl-prolyl cis-trans isomerase (PF3D7_0322000), which lacks any C-terminal cysteines and human Rab35, a host red blood cell protein.

**Localization of a Novel Parasite Protein FCP is Dependent on Prenylation**—In our proteomic dataset, FCP stood out as a unique protein that is conserved in *Plasmodium spp.* and related Apicomplexan parasites that cause human diseases. In addition, there is evidence that FCP may be essential for blood-stage *Plasmodium* growth (22). Therefore, we further characterized the function of its CaaX prenylation. FCP was previously shown to localize to the *Plasmodium* food vacuole, site of host hemoglobin digestion (22). Unexpectedly, this localization did not require the binding of phosphatidylinositol.
3-monophosphate (PI3P) by the FYVE domain but was dependent on the C-terminal 44 amino acids of FCP (22). Based on these previous findings, we hypothesized that prenylation of its CaaX motif may be responsible for membrane association and localization of FCP to the food vacuole. To test this hypothesis, we generated P. falciparum strains overexpressing N-terminal GFP-tagged wild-type or C322S mutant FCP. Cys322 is the modified cysteine residue in the CaaX motif, and its mutation to serine is expected to block prenylation of FCP. Indeed GFP-FCP purified from parasites treated with AlkFOH was covalently modified with this prenyl analog (visualized by reaction with fluorophore-azide), whereas labeling was not observed for GFP-FCP(C322S), consistent with loss of the prenyl modification when the CaaX motif is mutated (Fig. 3A). Live fluorescence microscopy also showed that GFP-FCP and GFP-FCP(C322S) differed in their cellular localization (Fig. 3B). GFP-FCP showed fluorescence overlapping or surrounding parasite hemozoin, consistent with its previously reported food vacuole localization. However, GFP-FCP(C322S) fluorescence appeared throughout the cell consistent with cytosolic mislocalization. Additional representative images are shown in supplemental material (Supplemental Fig. S1).

Rab5b Prenylation Does Not Require C-terminal Cysteine Residues—The identification of Rab5b in the prenylated proteome was also not predicted as it lacks C-terminal cysteines that are the known site of modification by RabGGT in all other organisms. Instead Rab5b was previously shown to be N-myristoylated and palmitoylated, and these lipid modifications were believed to substitute for prenylation to mediate Rab5b membrane localization at the parasite food vacuole, parasite plasma membrane, and parasitophorous vacuole (1, 2, 6, 23, 24). However, Rab5b was strongly identified in our proteomic data set with 53% sequence coverage. Identified peptides mapped across the entirety of the full-length sequence of Rab5b. As Parasites expressing Rab5b-GFP were treated with either DMSO or AlkFOH and Rab5b-GFP was immunoprecipitated, AlkFOH incorporation was visualized by a click reaction with AzTAMRA (top panel) whereas Rab5b-GFP loading was visualized by an αGFP immunoblot (bottom panel). The nucleus was stained using Hoechst. Additional images are shown in supplemental material (Supplemental Fig. S1).

The THQ Class Farnesyltransferase Inhibitor BMS-386914 Disrupts FCP Prenylation and Localization—Although the mechanism-of-action of antimalarial FT inhibitors depends on the cellular functions of CaaX protein substrates whose prenylation is disrupted, no specific CaaX protein substrate has...
been linked to FT inhibition. In addition, it would be ideal to have an assay that correlates cellular prenylation activity with antimalarial growth inhibitory activity. Therefore, we determined the effect of FT inhibitors on cellular prenylation of FCP. The THQ compound BMS-386914 has been shown to inhibit the enzymatic activity of \textit{P. falciparum} FT with an IC$_{50}$ of 0.9 nM and blood-stage \textit{Plasmodium} growth with an EC$_{50}$ of 5 nM, whereas lonafarnib showed IC$_{50} > 250$ nM and EC$_{50} = 3$ $\mu$M (8). Against \textit{P. falciparum} W2 parasites, we determined the EC$_{50}$ of BMS-386914 for growth inhibition was 25 nM and that of lonafarnib was 1.2 $\mu$M. We observed the amount of AlkFOH labeling of GFP-FCP in the presence of 1x and 10x EC$_{50}$ concentrations of each drug. Consistent with its mechanism-of-action inhibiting PfFT, BMS-386914 blocked prenylation of FCP at similar concentrations in which parasite growth inhibition is observed (Fig. 5A). Furthermore, BMS-386914 inhibition of FCP prenylation also caused its cytosolic mislocalization, like that observed in the GFP-FCP(C332S) mutant (Fig. 5B and supplemental Fig. S2). Quantification of GFP-FCP localization from three independent biological replicates, n>50 for each condition. Additional scored images are shown in the supplemental material (supplemental Fig. S2).

In contrast, parasite growth inhibition by lonafarnib does not disrupt FCP prenylation, suggesting that lonafarnib’s antimalarial activity is not because of FT inhibition.

**DISCUSSION**

Our findings indicate that \textit{Plasmodium} parasites contain markedly few prenylated proteins, only 20 high-confidence identifications out of 5398 protein-coding genes representing 0.3% of the proteome. Other protein lipid modifications, such as S-palmitoylation and N-myristoylation, were reported to be more abundant in \textit{P. falciparum} parasites using similar metabolic labeling and chemoproteomic approaches (1, 2). Though the low number of identifications is surprising, we believe our data set is an accurate estimate of prenylated proteins, and biological factors likely account for the low abundance of cellular prenylation in blood-stage \textit{Plasmodium}. \textit{Plasmodium} lacks homologs to several major families of prenylated proteins found in mammalian cells, including Rho/Rac GTPases, nuclear lamins, and G$_\gamma$ subunits of heterotrimeric G-proteins (25). Other protein families have been significantly reduced. For example, \textit{Plasmodium} encodes just 11 Rab GTPases compared with 66 in humans (6, 26). There are also fewer numbers of potential CaaX substrates with only 95 \textit{Plasmodium} proteins containing the required cysteine residue at position –4 from the C-terminus, and only a small subset of these are expected to be recognized as a CaaX motif and

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**Fig. 5.** BMS-386914 inhibits FCP prenylation and causes mislocalization to the cytosol. A, GFP-FCP was immunoprecipitated from AlkFOH labeled parasites treated with either 1x EC$_{50}$ or 10x EC$_{50}$ BMS-386914 and lonafarnib. AlkFOH labeling was detected by reaction with AzTAMRA fluorescent tag (top panel) whereas GFP loading was visualized through an aGFP immunoblot (bottom panel). B, Live microscopy of GFP-FCP expressing parasites treated with 10x EC$_{50}$ BMS-386914 or lonafarnib. The nucleus was stained using Hoechst.

C, Quantification of GFP-FCP localization in untreated, BMS-386914 (10x EC$_{50}$) and lonafarnib (10x EC$_{50}$) images. Based on three biological replicates, n>50 for each condition. Additional scored images are shown in the supplemental material (supplemental Fig. S2).
prenylated. Even the prenylation site of the most well-known prenylated protein, Ras GTPase, is not conserved in the Plasmodium homolog. The single P. falciparum Ras GTPase (PF3D7_0616700) lacks C-terminal cysteines and was not identified among medium- or high-confidence candidates, though its transcript expression level is similar to PF3D7_1428700, the lowest expressing gene among the high-confidence candidates. Thus PRRs prenylation may no longer be required despite being highly conserved in other eukaryotes. Finally, the limited number of Plasmodium prenylated proteins is consistent with lower estimates of prenylated protein abundance in lower eukaryotes compared with mammalian cells (27). These biological factors predict that protein prenylation is drastically reduced in Plasmodium spp. parasites, consistent with our results.

The efficiency and broad incorporation of modified prenyl analogs into parasite proteins also suggests that our ability to detect prenylated proteins was not limited by this approach. AlkFOH was previously shown to be recognized by all three types of protein prenyltransferases and incorporated into >100 known and novel prenylated proteins in mammalian cells (12, 18). In fact, labeling in mammalian cells was significantly less efficient than in Plasmodium parasites, requiring 50-fold higher concentration of the analog and depletion of endogenous prenyl lipids by inhibitor treatment. In our experiments, 1 µM AlkFOH was able to compete with endogenous prenyl lipids under conditions that maintained parasite viability. Under these conditions, both Rab GTPases and CaaX substrates, which are recognized by different protein prenyltransferase classes, were enriched. Labeling with an alkyne-modified geraniol (10-carbon) yielded similar identifications as those in AlkFOH-labeled samples, suggesting that the specificity of Plasmodium prenyltransferases for substrate proteins is not altered by moderate modifications of the prenyl lipid (supplemental Table S2). Independent experiments performed using alternative alkyne- and azide-modified farnesol analogs also did not identify additional prenylated targets (R.A. Serwa, M. Jones, P. Bowyer, E.W. Tate, and J.C. Rayner; personal communication). Thus, this chemoproteomic approach was robust and likely quite sensitive.

In fact, a significant advantage of our unbiased approach was the ability to identify novel prenylated proteins such as FCP and PF3D7_1428700, because these would not have been recognized as high-priority candidates based on sequence homology with previously known prenylated proteins. Bioinformatic prediction of CaaX motifs recognized by FT and GT1 is also not sufficiently sensitive, because the Prenylation Prediction Suite (PrePS) software recognized only five of the eight high-confidence CaaX protein candidates (Table I and supplemental Table S3) (28). Our high-confidence list assumes that Plasmodium FT/GT1 enzymes recognize and prenylate Caax-containing proteins, similar to other organisms. However, numerous medium-confidence candidates are neither homologs of known prenylated proteins nor have a recognizable prenylation site. One strategy to identify true novel prenylated proteins among these candidates would be to select lower abundance proteins (based on transcript expression levels) for validation, because these are less likely to be nonspecific contaminants. Importantly, though protein farnesylation and geranylgeranylation on cysteines are by far the most common prenyl modification, we cannot rule out the possibility that the AlkFOH probe is further metabolized to longer chain prenyl groups or other isoprenoid derivatives before covalent attachment to protein. Indeed protein dolicholysis (C55) has been reported in Plasmodium, based on the incorporation of radioactive label derived from farnesyl diphosphate into dolichols released from unidentified proteins (29). These alternative prenyl modifications were detected at low abundance and were labile to methyl iodide consistent with thioether attachment via cysteine residues. Lastly, our data set only includes proteins expressed and prenylated during trophozoite/schizont stages when AlkFOH labeling was performed and may miss very low abundance proteins (<10 RPKM transcript expression).

Despite the limited number of prenylated proteins in Plasmodium, we detected several examples of prenylation activity unique to the parasite that may target proteins to novel membranes in the Plasmodium-infected red cell. FCP in particular stood out as a novel parasite-specific protein, as it was the only protein in our data set that is only conserved among Apicomplexan parasites. FCP localizes to a parasite-specific compartment, the food vacuole, and we demonstrate that prenylation at its CaaX motif is necessary for this localization through both mutagenesis and FT inhibitor treatment. The Toxoplasma FCP homolog has also been localized to its vacuolar compartment, VAC, which is functionally analogous to the food vacuole (30). Furthermore, FCP function may be essential because, in a previous study, a truncated form of FCP (lacking the FYVE domain) still localized to the food vacuole but had a dominant-negative effect on parasite growth and abnormal food vacuole morphology (22). Finally, FCP is one of only 3 PI3P-binding proteins present in P. falciparum, binding PI3P via its FYVE domain (22). PI3P is normally found on early endosomes, however, in P. falciparum it is present on two parasite-specific compartments, the food vacuole and the apicoplast (31). Normally, PI3P-binding proteins are not themselves post-translationally lipidated but often interact with prenylated and palmitoylated proteins. For example, the mammalian protein early endosomal antigen (EEA1) binds PI3P and interacts with Rab GTPases (32). By contrast, FCP was previously identified in the palmitoylated proteome and is now shown to be prenylated, making it the first example, to our knowledge, of a lipidated PI3P-binding protein (2). Notably Plasmodium does not encode a recognizable EEA-1 homolog, and a lipidated, PI3P-binding, coiled-coil protein like FCP may serve a similar function as EEA-1. Considering all the evidence, FCP is likely to have an important and pathogen-specific membrane function dependent on
its prenylation. Future studies will be necessary to determine whether it is essential for growth and identify its protein interaction partners.

We also detected atypical RabGGT activity in *Plasmodium*. Our results demonstrate that Rab5b, which has been localized to the parasite food vacuole, parasite plasma membrane, and parasitophorous vacuole, is prenylated even though it lacks C-terminal cysteines that are the only known modification site of Rab GTPases in other organisms. Instead we propose that Rab5b is prenylated on one or more internal cysteine residues. In our proteomic sample preparation, the peptide containing the modified residue remains attached to the agarose beads following on-bead trypsin digestion. Based on the sequence coverage of Rab5b, only three cysteines were not represented by identified peptides and are potential candidates for the modification sites: one at residue three near the N terminus (likely palmitoylation site) and two tandem cysteines at residues 141–142 (Fig. 4A). The dicysteine site has previously been suggested to serve as an internal prenylation site (6). Normally unprenylated Rab GTPases are bound by the Rab escort protein (REP), which presents their C-terminal cysteine residues to the RabGGT for prenylation. Given this mode of recognition, it is possible that a putative *Plasmodium* REP (PF3D7_1038100) may present internal cysteines for modification. Another alternative is that the prenyl modification occurs on a noncysteine residue but this seems unlikely as it would require a change in the chemical reaction catalyzed by the RabGGT. There is no known biosynthetic route that converts isoprenoids to other common protein modifications. Thus, the modification of Rab5b by AlkFOH cannot be explained by the presence of palmitoylation, myristoylation, or other known post-translational modifications. As is the first report of Rab prenylation on a non-conserved site, it may be unique to *Plasmodium* but may also have gone undetected in other organisms. Interestingly Ara6, a plant Rab5 homolog, also lacks C-terminal cysteines and is both N-myristoylated and palmitoylated, thus it may also be prenylated on a non-conserved site (23, 33). Both N-myristoylation and palmitoylation of Rab5b are necessary for localization to two parasite-specific compartments, the food vacuole and the parasitophorous vacuole; it remains to be determined whether prenylation is also important for its localization (2, 23, 24). In particular, mutation of the internal dicysteine Cys141 and Cys142 may reveal the nonconserved prenylation site and show whether prenylation is important for Rab5b localization.

The most surprising prenylated protein identified in our high-confidence list was a human Rab GTPase, HsRab35. First, Rab GTPases in mature red blood cells should already be prenylated, therefore the C-terminal cysteines of HsRab35 should not be available for AlkFOH labeling. Second, in *Plasmodium*-infected red cells, the only PPTs are expressed by the parasite and present in the parasite cytosol, whereas HsRab35 is expected to localize to the red blood cell compartment (19). Despite these apparent discrepancies, HsRab35 was not identified in AlkFOH-treated, uninfected red blood cells, implying that AlkFOH labeling of HsRab35 is dependent on parasite infection. Only three unique peptides were detected for HsRab35, resulting in 6% protein coverage. Indeed, six other human Rab GTPases were also detected by single unique peptides in our prenylated proteome, representing seven of the nine HsRab GTPases identified in the human red blood cell proteome (19). The sparse peptide identifications likely reflect the low abundance of human Rab GTPases, which is not surprising because the proteomic samples were prepared from saponin-lysed parasites in which host cell proteins in the red blood cell membrane and cytoplasm were specifically depleted. The fact that these human Rab GTPases were detected in the prenylated proteome at all suggests they were specifically enriched. Human Rab GTPases may be more robustly detected in proteomic samples prepared from whole *Plasmodium*-infected red cells. Unfortunately, we were unsuccessful in purifying a tagged HsRab35 introduced into resealed red cells to directly detect its parasite-dependent prenyl modification. Although the detection of a human Rab GTPase in our prenylated proteome is intriguing and hints at a potential role for parasite-dependent prenylation at novel membranes in the host cell compartment, at the moment we are unable to provide direct evidence for prenylation of host cell proteins.

Finally, in addition to identifying unique parasite prenylation activity, the *Plasmodium* prenylated proteome will bolster the investigation of FT inhibitors as antimalarials. We identified eight CaaX proteins as candidate FT substrates, some or all of which may be responsible for parasite growth inhibition caused by FT inhibitors (Table I). Further validation of these candidates will require that (1) cellular prenylation of its CaaX motif is inhibited by FT inhibitors and (2) prenylation is required for parasite growth. As seen previously, we observed the loss of several ~50 kDa AlkFOH-labeled proteins upon treatment with THQ BMS-386914 (supplemental Fig. S3) (8). A high-confidence candidate in our dataset, PF3D7_1437900 (Hsp40), may represent one of these abundant prenylated proteins that are disrupted. We also demonstrated that BMS-386914 disrupts prenylation and localization of Apicomplexan-specific protein FCP. As expected, it did not disrupt Rab5b prenylation which is most likely catalyzed by the Rab geranylgeranyltansferase (data not shown). In contrast, lonafarnib which potently inhibits human FT activity has no effect on FCP prenylation and likely causes parasite growth inhibition via an alternative non-FT *Plasmodium* target (8, 9, 34). The difference between the mechanism-of-action of FT inhibitors in mammalian cells and *Plasmodium* parasites reinforces the fact that *Plasmodium* protein prenyltransferrases are divergent from human protein prenyltransferases and can be targeted selectively as a strategy toward novel anti-malarial development (8, 9).
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**DATA AVAILABILITY**

The raw mass spectrometry data has been deposited at the Chorus Project (chorusproject.org) with the project identifier 1212.

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