Validation of *N*-myristoyltransferase as an antimalarial drug target using an integrated chemical biology approach

Authors: Megan H. Wright^{1,2}, Barbara Clough³, Mark D. Rackham¹, Kaveri Rangachari³, James A. Brannigan⁴, Munira Grainger³, David K. Moss³, Andrew R. Bottrill⁵, William P. Heal^{1,†}, Malgorzata Broncel¹, Remigiusz A. Serwa¹, Declan Brady⁶, David J. Mann^{2,7}, Robin J. Leatherbarrow^{1,2,#}, Rita Tewari⁶, Anthony J. Wilkinson⁴, Anthony A. Holder³, Edward W. Tate*,1,2

Affiliations:

- 1. Department of Chemistry, Imperial College London, London SW7 2AZ, UK.
- 2. Institute of Chemical Biology, Imperial College London, London SW7 2AZ, UK.
- 3. Division of Parasitology, MRC National Institute for Medical Research, The Ridgeway, Mill Hill, London, NW7 1AA, UK.
- 4. York Structural Biology Laboratory, Department of Chemistry, University of York, York YO10 5DD, UK.
- 5. Protein and Nucleic Acid Chemistry Laboratory, University of Leicester, Hodgkin Building, Lancaster Road, Leicester LE1 9HN, UK.
- 6. Centre for Genetics and Genomics, School of Biology, Queens Medical Centre, University of Nottingham, Nottingham, NG2 7UH, UK.
- 7. Division of Molecular Biosciences, Department of Life Sciences, Imperial College London, London SW7 2AZ, UK.
- † Current address: Department of Chemistry, Kings College London, London SE1 1UL, UK.
- # Current address: Liverpool John Moores University, Liverpool L1 2UA, UK.

Abstract

Malaria is an infectious disease caused by parasites of the genus *Plasmodium* that inflicts approximately one million deaths *per annum* worldwide. Chemical validation of new antimalarial targets is urgently required in view of rising resistance to current drugs. One such putative target is the enzyme *N*-myristoyltransferase (NMT), which catalyzes *N*-myristoylation of protein substrates. Here we report an integrated chemical biology approach to explore protein myristoylation in the major human parasite *P. falciparum*, combining chemical proteomic tools for identification of the myristoylated and glycosylphosphatidylinositol-anchored proteome with selective small molecule NMT inhibitors. We demonstrate that NMT is an essential and chemically tractable target in malaria parasites both *in vitro* and *in vivo*, and show that selective inhibition of *N*-myristoylation leads to catastrophic and irreversible failure to assemble the inner membrane complex, a critical subcellular organelle in the parasite life cycle. Our studies provide the basis for development of new antimalarials targeting NMT.

Malaria is an infectious disease caused by parasites of the genus *Plasmodium*, and is among the most serious in the developing world, resulting in an estimated 200 million cases and 1.2 million deaths in 2010¹. The majority of cases and mortality are due to P. falciparum in sub-Saharan Africa, whilst P. vivax is widespread and causes persistent infections². Resistance has been recorded against all frontline therapies, and the need for new drugs is widely recognized³. N-Myristoyltransferase (NMT, EC 2.3.1.97) catalyzes transfer of the lipid myristate (C14:0) from myristoyl coenzyme A (Myr-CoA) to specific substrate proteins in eukaryotes⁴, and has been explored preclinically as a target in fungal⁵ and trypanosome infections⁶, and suggested as a potential target in malaria and leishmaniasis⁷. Protein *N*-myristoylation occurs predominantly co-translationally at an N-terminal glycine following removal of initiator methionine (Fig. 1a); although an N-terminal glycine is essential, there is no panspecies consensus sequence for myristoylation⁴. *In silico* analyses suggest that up to 2% of proteins in *P. falciparum* may be substrates for NMT (**Supplementary Table S1**), although lack of suitable training sets for *Plasmodium* myristoylation results in inconsistent predictions⁸. To date, only two proteins have been shown to be N-myristoylated in their native context in P. falciparum – calcium-dependent protein kinase-1 (CDPK1)9 and glideosome-associated protein 45kDa (GAP45)10 – and there is indirect evidence for a small number of other substrates 10, 11, 12, 13. N-Myristoylation regulates membrane localization and protein complex assembly in eukaryotes¹⁴, and selective inhibition of NMT is hypothesized to lead to pleiotropic effects on the parasite through multiple downstream substrates. *Plasmodium* species possess a single NMT gene¹⁵ and partial knockdown of NMT expression in the rodent malaria parasite P. berghei was found to result in rapid depletion of parasitemia in vivo, demonstrating that NMT is essential in this murine model of

malaria¹⁶. However, target validation in human malarial parasites is particularly challenging due to their complex life cycle, the limited range of genetic tools available, and the very low efficiency of genetic modification¹⁷. Despite intensive research efforts, conditional genetic knockout of an essential gene in *P. falciparum* has not been demonstrated to date, and thus chemical genetic knockdown approaches based on small molecule inhibitors are a particularly powerful and versatile strategy for target validation.

Here we present an integrated chemical biology approach to understand inhibition of *N*-myristoylation in *P. falciparum* parasites that enables both the study of protein myristoylation and the chemical dissection of NMT as a drug target in this genetically intractable organism. These studies greatly expand knowledge of both *N*-myristoylated and glycosylphosphatidylinositol (GPI) anchored proteins in *Plasmodium*, and provide the first compelling evidence that NMT is a druggable target in the symptomatic blood stage of the most important human malaria parasite. Using these tools in combination with quantitative whole proteome analyses and cellular studies, we demonstrate that NMT inhibition results in a catastrophic and irreversible failure to form critical parasite subcellular structures, followed by rapid parasite cell death. Furthermore, we define potent starting points for the development of NMT inhibitors as potential antimalarials that are active against human malaria parasites *in vitro*, and *in vivo* in an animal model of malaria.

Results

Myristoyl-CoA analogue YnMyr-CoA is a biomimetic substrate for *Plasmodium* NMT

Lipid probes bearing a bioorthogonal tag such as a terminal alkyne are powerful tools for analysis of protein lipidation in living systems 18, 19. The myristate surrogate tetradec-13-ynoic acid (YnMyr) can be transferred from YnMyr-CoA thioester to target proteins by recombinant NMT²⁰, in bacterial co-expression systems²¹ or in mammalian cells²². Subsequent chemoselective ligation through copper(I)-catalyzed [3+2] azide-alkyne cycloaddition (CuAAC) imparts a range of useful functionality to tagged proteins, including a fluorophore and/or affinity label (Fig. 1b)^{20, 22, 23, 24}. We determined the structure of NMT from P. vivax (PvNMT, which shares >80% identity with P. falciparum NMT (PfNMT))²⁵ in complex with YnMyr-CoA (PDB 2YNC): Supplementary Fig. S1 and Table S2), revealing that YnMyr-CoA occupies an extended groove that runs across one face of the enzyme, with the YnMyr chain occupying the fatty acyl binding pocket (**Fig. 1c**). The terminal *sp* hybridized carbon atoms are accommodated without steric clashes and the hydrophobic contacts and native interactions of the thioester carbonyl with backbone amide and polar side chain moieties are preserved (Fig. 1c)²⁶. This structure is the first reported example of a tagged analogue of a post-translational modification precursor in complex with its cognate transferase, and together with kinetic data (Supplementary Fig. S2) indicates that a terminal alkyne moiety is fully compatible with enzymatic transfer of YnMyr to substrate proteins.

YnMyr tags *N*-myristoylated and GPI-anchored proteins in *P. falciparum* blood stage parasites

We next investigated the capacity of YnMyr to tag proteins in the asexual stage of the parasite, which invades and replicates within red blood cells (RBCs). Intracellular P. falciparum 3D7 parasites (schizonts) were treated with YnMyr at concentrations up to 50 µM and allowed to develop over 5 h; no changes were observed in morphology or life cycle progression in YnMyr-treated parasites. Parasite proteins were isolated and CuAAC ligation to tri-functional capture reagent AzTB (azido-TAMRA-PEG-Biotin; Supplementary Fig. S3)²⁷ enabled direct in-gel fluorescence detection of protein tagging following SDS PAGE (Fig. 2a). YnMyr tagging was concentration-dependent with no detectable background, readily out-competed by excess myristate, and inhibited by the protein synthesis inhibitor cycloheximide (CHX, **Supplementary Fig. S4**), consistent with biomimetic co-translational modification of target proteins. The biotin label in AzTB further permitted quantitative immobilization of fluorescently labeled proteins on streptavidin-coated beads (Supplementary Fig. S5). The known N-myristoylated proteins CDPK1 and GAP45^{10, 28} were selectively pulled down from parasites cultured in the presence of YnMyr, and resistance of labeling to base and hydroxylamine indicated the presence of N-acyl (amide linked) YnMyr (Fig. 2a and Supplementary Fig. S6). Control experiments cross-comparing metabolic incorporation of the longer chain palmitate probe YnPal (heptadec-14-ynoic acid) confirmed that YnMyr was not significantly incorporated into sites of S-palmitoylation, which occurs through a thioester linked to cysteine side chains (Supplementary Fig. S7). However, a substantial fraction of the fluorescently labeled bands were base sensitive suggesting the presence of ester-linked YnMyr (Fig. 2a). Ester-linked incorporation of myristate at the GPI

anchor inositol is characteristic of *Plasmodium* species²⁹, and we found that the abundant GPI-anchored surface protein MSP1 was indeed tagged by YnMyr in a base-sensitive manner (**Fig. 2a**). Taken together, these data suggest that YnMyr is a high-fidelity myristate mimetic in live parasites, and can be used as a probe for both *N*-myristoylated and GPI-anchored proteins under standard culture conditions.

Chemical proteomic profiling of *N*-myristoylated and GPI-anchored proteins in *P. falciparum*

We next performed de novo identification of YnMyr targets in P. falciparum by LC-MS/MS based chemical proteomics after AzTB ligation, pull-down and protein digest, using both gel-based and gel-free approaches. Tryptic peptides were analyzed by a standard proteomics workflow, and proteins found with very high (>99 %) confidence and >4-fold enrichment compared to controls were considered for further analysis; proteomic methods are described in detail in Supplementary Methods. N-Myristoylated target proteins were mapped to specific regions of the gel following gel-based proteomics (Fig. 2b; Supplementary Fig. S8 and Table S6), identifying ADP-ribosylation factor 1 (ARF1) as the major component of the strongest fluorescent band. A novel antibody directed against PfARF1 further confirmed this assignment (Fig. 2c). Base-treatment prior to pull-down was used to dissect the contributions of GPI-anchored and N-myristoylated proteins; identification of MSP1 and predicted GPI-anchored proteins was dramatically reduced or eliminated by base treatment, whilst putative *N*-myristoylated proteins were unaffected (**Fig. 3a**; Supplementary Tables S3, S4 and S5)^{30, 31}. YnMyr-tagged peptides were not detected in these analyses, due to the presence of the biotin label added during CuAAC ligation that anchors the peptide to the resin during tryptic digest. Indeed, direct determination of the site of modification has been recognized as a significant

unsolved challenge in metabolic tagging of lipidated proteins³² due to poor release of the biotinylated peptide from the streptavidin resin and the large size of the resultant label. We therefore designed a new tetra-functional capture reagent (AzKTB, **Fig. 3b**) which bears a short trypsin-cleavable peptide sequence between the azide module and the TAMRA/PEG-biotin labels. Tryptic on-bead digest releases the YnMyr-modified peptide bearing a hydrophilic zwitterionic moiety, permitting the site of YnMyr modification to be determined unambiguously by tandem MS (**Supplementary Fig. S9**). Using this new approach, 17 N-terminal YnMyr-Gly motifs were detected (**Fig. 3a** and **Supplementary Table S7**), further confirming the fidelity of the YnMyr probe for NMT substrates in live parasites.

NMT substrates in wild type parasites, encompassing a diverse range of functions including motility, protein transport, parasite development and phosphorylation pathways (**Fig. 3c**). As expected, many substrates have predicted or known localization at membranes or in membrane-bound compartments (**Supplementary Fig. S10**). Across the proteomic datasets five proteins had previously been annotated as PfNMT substrates *via* direct (GAP45, CDPK1) or indirect (GRASP1¹³, Armadillo repeats-only protein ARO³³, ARF1³⁰) experimental evidence. New proteins were identified that are *N*-myristoylated in other eukaryotes, including several ARF/ARL proteins and the 26S proteasome regulatory subunit 4 (P26s4), which is highly enriched in YnMyr-tagged samples³⁴. Interestingly, studies in yeast have shown that loss of *N*-myristoylation of the yeast P26s4 homologue results in aberrant localization of the entire proteasome complex³⁴ and the proteasome has recently been described as a promising drug target in *P. falciparum*³⁵. Other substrates with important functions include Protein Kinase A regulatory subunit PKAr³⁶, the small

GTPase Rab5b³⁷ and CDPK4, a kinase essential in gametogenesis and disease transmission³⁸. We also identified the small inner-membrane complex (IMC) proteins ISP1 and ISP3; along with GAP45 these proteins have important roles in regulating formation of the IMC, a subcellular organelle that is critical for assembling the molecular motor apparatus required for RBC invasion³⁹.

Identification and structural characterization of potent small molecule inhibitors of *P. falciparum* and *P. vivax* NMTs

The diversity of function of the identified *N*-myristoylated proteome in *P. falciparum* indicates that specific NMT inhibition could substantially reduce parasite viability. To support this hypothesis, we sought to identify potent chemical inhibitors to enable selective and tunable chemical genetic knockdown of NMT activity in live parasites. As noted above, such tools are particularly powerful when applied to *P. falciparum* since there is currently no proven genetic method for inducible knockdown of an essential gene in this parasite. Through further development of our recently-reported Plasmodium NMT inhibitors 40 combined with screening known inhibitors of NMT from other organisms we identified two distinct PfNMT/PvNMT inhibitor series 1 and 2 (Fig. 4a). Series 1 compounds were first reported as inhibitors of T. brucei NMT, and compound **1a** is effective on trypanosomes in vitro and in vivo⁶; however, the causative agents of malaria and trypanosomiasis have entirely different mechanisms of infection and disease pathology. Series 2 is derived from a P. falciparum NMT inhibitor based on a benzothiophene scaffold, 40 the structure-guided development of which will be reported elsewhere in due course. X-ray crystallography was used to determine the binding mode of the most potent compounds 1a and 2a in ternary complexes with PvNMT and a non-hydrolyzable myristoyl-CoA analogue (PDB 2YND and 2YNE; Supplementary Table S2). Both inhibitors bind at the peptide

binding pocket, with **1a** exhibiting a similar binding mode to that observed in NMTs from other species (**Fig. 4b** and **Supplementary Fig. S11**). A salt bridge is formed between the positively charged piperazine (**1a**) or piperidine (**2a**) nitrogen and the C-terminal carboxylate of the enzyme, which plays an essential catalytic role during myristate transfer⁴. Despite this common point of contact, the trajectory of these two chemotypes through the binding site is distinct. Substitution of the trimethyl pyrazole (**1c**, **2b**) substantially reduces potency, likely due to reduced H-bonding capacity, whilst sulfonamide alkylation in **1b** yielded a more subtle decrease (**Fig. 4a**). Although **1a-c**⁴¹ are more potent against *Homo sapiens* NMT (HsNMT1) than against the *Plasmodium* enzymes, RBCs lack *de novo* protein synthesis and do not express NMT, suggesting that *in vitro* culture would not be compromised⁴². In contrast, **2a-b** possess promising selectivity over HsNMT1, with **2a** being the most potent PfNMT and PvNMT inhibitor reported to date.

NMT inhibitors act on-target to kill *P. falciparum* blood stage parasites and reduce parasite burden in an *in vivo* model of malaria

With inhibitor and substrate chemical probes for PfNMT characterized at the structural, biochemical and proteomic level, we sought to determine whether **1a-c** and **2a-b** act on-target in parasites in culture and to assess NMT as a potential drug target in blood stage *P. falciparum*. Analysis of EC₅₀ values for parasite growth inhibition in synchronized intraerythrocytic *P. falciparum* cultures revealed that translation of potency from enzyme to cell (EC₅₀/K_i^(app)) varied significantly across series (**Fig. 5a**). Uninfected RBCs remained morphologically normal (**Supplementary Fig. S12**). Since protein *N*-myristoylation occurs co-translationally we reasoned that inhibition of NMT in parasites could be assessed directly by treatment with inhibitor combined with pulse tagging with YnMyr. When tagging

experiments were repeated in the presence of **1a** a highly specific dose-dependent depletion of fluorescence was observed for all bands previously identified as NMT substrates, with no change in GPI-anchored proteins (Fig. 5b; Supplementary Figs. **S13** and **S14**). Parallel experiments with the tagged methionine surrogate azidohomoalanine (AHA⁴³, **Supplementary Fig. S3**) showed that **1a** does not directly affect protein synthesis (Supplementary Fig. S15). Selective inhibition was further confirmed for specific substrate proteins by Western blot against CDPK1 and GAP45, and using novel antibodies directed against ARF1 and ARO (Fig. 5b; Supplementary Figs. S16 and S17). Total abundance of each protein analyzed was unaltered by inhibitor treatment over this short pulse tagging period; however, on pull-down the amount of each YnMyr-tagged substrate recovered was dosedependently inhibited, whilst MSP1 remained unaffected. Quantification of the 20 kDa ARF1 band by fluorescence intensity or Western blot gave 346 nM as the inhibitor concentration required for 50% reduction in YnMyr tagging (here termed TC_{50}) for **1a**, closely matching the TC_{50} values calculated for other specific substrates by quantification of Western blots (Fig. 5c and Supplementary Fig. S18).

Series 1 and 2 were compared for their capacity to knock down NMT activity in parasites with the aim of rationalizing their diverse enzyme to cell translation (**Fig. 5a** and **Supplementary Figs. S19** and **S20**). An excellent correlation was observed between TC_{50} and parasite viability (EC_{50}) across all compounds tested ($R^2 = 0.97$, **Fig. 5a**), establishing a direct link between in-cell NMT inhibition and efficacy, and resolving the differences in translation between members of series 1 and 2. In particular, compounds 1a and 2a have very similar effects on protein myristoylation in the cell despite their significantly different K_i values, resulting in similar EC_{50} values for parasite killing (**Fig. 5a**). These data demonstrate that **1a-c** and **2a-b**

possess the same on-target mode of action, and establish the chemical tractability of NMT in blood stage human malaria parasites.

We also examined the impact of NMT inhibition on parasite burden *in vivo* in the rodent malaria parasite P. berghei. Both $\mathbf{1a}$ and $\mathbf{2a}$ maintained reasonable potency against recombinant PbNMT ($K_i = 90$ and 14 nM, respectively), but initial experiments found that $\mathbf{1a}$ induced toxicity *in vivo* at sub-therapeutic concentrations. However, when $\mathbf{2a}$ was administered to infected mice a significant ($p \le 0.01$, n = 3) reduction in parasitemia was observed compared to controls (PBS vehicle) over the course of 3 days treatment (50 mg/kg intraperitoneally, twice daily) (**Fig. 5d**). Furthermore, $\mathbf{2a}$ caused no acute toxicity and animals survived without visible symptoms past day 3, whereas all control mice needed to be euthanized due to acute symptoms of malaria. This first demonstration of *in vivo* activity of a chemical inhibitor of *Plasmodium* NMT is encouraging, particularly given that the potency and pharmacokinetic properties of $\mathbf{2a}$ are suboptimal (Supplementary Table S8).

NMT inhibition results in formation of non-viable parasites lacking critical components required for development and RBC invasion

Having confirmed on-target action of NMT inhibitors in parasites, we sought to use these tools to investigate how NMT inhibition leads to parasite cell death. During asexual reproduction, *P. falciparum* undergoes a 45-48 hour life cycle within infected RBCs (iRBCs), maturing from ring stage, to trophozoite, and then schizont, before differentiation into merozoites that egress and reinvade RBCs (**Fig. 6a**). We examined the effects of NMT inhibition on life stage progression, monitoring growth at 1 μ M **1a**, a dose that causes approximately 80% inhibition of protein myristoylation in parasites (**Fig. 5**). NMT inhibitor or DMSO vehicle (control) was added to highly

synchronized ring stage parasites 2 hours post-invasion (PI), and the parasite population was monitored up to 45 hours PI. Whilst control parasites progressed as expected over 45 hours to form multinucleate schizonts, **1a**-treated parasites became trapped in a morphologically distinct life stage (here termed a 'pseudoschizont'). Nuclear staining (DAPI) revealed that at 45 hours all (n = 50, p < 10^{-30}) multinucleate parasites contained 4-7 (5.6 ± 1.1) nuclei in the presence of **1a**, compared to ca. 20 (19.6 ± 2.7) nuclei in control schizonts (**Fig. 6a** and **Supplementary Fig. S21**). The clarity and severity of this phenotype is striking given that **1a** was applied at a concentration that does not completely eliminate NMT activity in cells, and is consistent with the observed high sensitivity of parasites towards NMT inhibition (**Fig. 5**). At 66 hours PI, DMSO-treated parasites had egressed and reinvaded RBCs as expected, developing into new ring stages and resulting in an increase in parasitemia. However, in the presence of **1a** parasitemia dropped by >85%, showing that NMT inhibition results in a substantial loss of parasite viability (**Supplementary Fig. S22**).

In order to characterize the distinguishing features of pseudoschizonts, we performed a quantitative proteomic analysis in pseudoschizonts (inhibitor-treated) and schizonts (control) at 45 hours PI. The relative abundance of 1181 parasite proteins was determined through label-free quantification, revealing a conspicuous clustering amongst strongly affected proteins with known function (**Fig. 6b** and **Supplementary Table S9**). Multiple markers of the Inner Membrane Complex (IMC) were greatly reduced in abundance in inhibited cells (e.g. MyoA, MTIP, GAP45, ISP3), as were a set of late-expressed proteins specific to apical (secretory) organelles in developing merozoites; these included ARO, PTRAMP and key RBC binding proteins and antigens (e.g. EBA140/175/181, RH2a). Western blot analyses

of GAP45, MTIP, CDPK1 and ARO further confirmed marked depletion of these proteins in pseudoschizonts (**Fig. 6c**). Interestingly, there was relative accumulation of a set of soluble secreted proteins, including several members of the serine repeat antigen family (SERA) which under normal conditions are exported and processed during schizont development.

The IMC is comprised of flattened membrane compartments lying directly under the plasma membrane, where one of its roles is assembly of the molecular motor complex (including MTIP, MyoA and GAP45) that drives invasion of RBCs. In the intracellular parasite, assembly of the IMC occurs during schizogony and can be visualised with protein markers such as GAP45⁴⁴. In *Plasmodium* nuclear division precedes cell division, and the IMC initially appears as small ring-like structures, which then extend around developing merozoites during schizont segmentation (cell division). This transition was recapitulated in immunofluorescence images of DMSOtreated parasites at 35 and 45 hours PI, whereas in the inhibitor treated parasites the IMC structure was not visible at either time point (Fig. 6d). Given the established importance of the IMC in parasite development and invasion, we hypothesized that NMT inhibition-induced loss of the IMC would have catastrophic and irreversible consequences for parasite viability. Indeed, when inhibited parasites (1 µM 1a) were analyzed 10 hours later (at 55 hours PI), iRBCs bearing pseudoschizonts were largely eliminated even when inhibitor was removed at 45 hours PI, whereas control parasites egressed and reinvaded new RBCs to form new ring stages over the same timeframe (**Fig. 6e**).

Discussion

Here we have established a novel chemical approach to study N-myristoylation and inhibition of NMT in the most important human malarial parasite, and demonstrated that NMT is an essential and chemically tractable target in *P. falciparum*. Metabolic chemical tagging has enabled the first proteome-wide survey of both N-myristoylated and GPI-anchored proteins in P. falciparum. Previously, identification of PfNMT substrates relied on painstaking generation of mutant parasites or specific antibodies and radiolabeling that are impractical for high-throughput studies. In contrast, our chemical approach identifies substrates directly across the proteome in their natural context in live parasites, without the need for genetic manipulation and substrate overexpression. In combination with a new reagent for the identification of protein lipidation sites by metabolic tagging, this technology provides high confidence assignment of over 30 NMT substrates; the majority are identified for the first time in this study, and have roles in a wide range of essential processes. These include protein trafficking and secretion (ARF1, Rab5b, ARO), ion channel regulation (PKAr), organelle biogenesis (GRASP1), motility (CDPK1, GAP45), inner membrane complex (IMC) integrity (ISP1, ISP3, GAP45), development (CDPK4), and protein homeostasis (P26s4). Combining PfNMT inhibitors with a chemical proteomic toolbox (YnMyr, AHA, AzKTB) backed up by extensive characterization (biochemical, structural, in vitro, in vivo) provides a powerful and self-consistent validation of ontarget specificity. This study is also the first to apply AHA to track *de novo* protein synthesis in *Plasmodium*; notably, metabolic tagging proceeds efficiently without the need for defined media, and we anticipate that AHA will be a powerful alternative to radiolabeling for monitoring protein synthesis in malaria parasites.

Whilst work on fungi and trypanosomes suggests NMT may be an interesting potential target in infectious disease, *Plasmodium* parasites are evolutionarily

specialized, and infect their host and mosquito vector through unique and highly complex mechanisms; the specific consequences of PfNMT inhibition were thus unknown prior to the work reported here. Our chemical approach offers a unique opportunity to examine the phenotype arising from perturbation of the pathways in which NMT substrates are involved, particularly since knockout of an essential gene such as PfNMT is currently intractable in human malaria parasites. In particular, we discovered that PfNMT inhibition induced an unexpected, irreversible and catastrophic failure to assemble the IMC during early schizogony in asexual blood stage parasites. This was accompanied by loss of essential components of the myosin motor that drives RBC invasion (MTIP, MyoA), as well as the absence of proteins critical for parasite engagement of new RBCs (EBA140/175/181, RH2a). A strong reduction in proteins synthesized late in intracellular development, including characteristic markers of organelles assembled at the apical end of developing merozoites (rhoptries/ARO, micronemes/EBA family), provides strong evidence that inhibited parasites are non-infectious³³. Marked retention and failed processing of key proteins exported during late schizogony (SERA family, GBP, MSP9) further illustrates a failure to progress to merozoites. Interestingly, parasites commit to early rounds of nuclear division but are unable to proceed through cell division, supporting recent evidence for a key role of the IMC in development in addition to invasion⁴⁴. GAP45, ISP1 and ISP3 are NMT substrates identified here that are candidates for driving this IMC-related phenotype. N-Myristoylated GAP45 acts as a scaffold that spans the space between the plasma membrane and the IMC⁴⁵; it also has an essential role in organizing the acto-myosin motor that drives invasion of RBCs⁴⁴. We recently demonstrated that *P. berghei* ISP1 and ISP3 are highly expressed in invasive stages, and localize to the anterior apical end of the parasite where IMC

organization is initiated³⁹. Whilst deletion of ISP3 in *P. berghei* is compensated by up-regulation of ISP1, NMT inhibition may result in simultaneous inactivation of both proteins through loss of the membrane anchor. Loss of the IMC is sufficient to explain the irreversible and cidal mode of action of PfNMT inhibition during early schizogony, and failure to develop into infective merozoites. However, the comparative whole-proteome dataset obtained here shows that this is far from the only phenotype arising from PfNMT inhibition, and provides a rich resource that can be mined to initiate future studies of IMC formation, protein trafficking and assembly of the invasion apparatus.

Chemical knockdown of PfNMT effectively combines the benefits of a selective enzyme inhibitor with a defined multi-target, pleiotropic downstream mode of action. We have shown here that *P. falciparum* is highly sensitive to a global drop in *N*-myristoylation, and structural characterization of the binding modes of inhibitors at the protein substrate binding pocket of *Plasmodium* NMT provides the potential for future improvements in potency and selectivity through structure-guided design. Remarkably, despite unoptimized pharmacokinetic (PK) properties and modest *in vitro* potency, NMT inhibitor **2a** significantly reduced parasitemia in the rodent model *P. berghei* with no acute toxicity, providing further evidence for the druggability of NMT in malaria. This *in vivo* proof of principle suggests that compounds with improved potency and PK properties could form the basis for an antimalarial drug discovery program⁴⁶.

In future studies, the chemical tools described here may be applied to examine the role and druggability of PfNMT and *N*-myristoylation in other stages of the parasite, including in the sexual blood stages and liver stages that are important targets for blocking transmission. Our data suggest that NMT inhibitors could be highly effective

against these life stages; for example CDPK4, identified here as an NMT substrate, is a promising target for kinase inhibitors that block development of sexual stage parasites³⁸, whilst invasion of liver cells requires an intact IMC⁴⁷. The chemical tools described here also provide a novel approach to study the role of the IMC in parasite development, given the striking effects of NMT inhibition on this structure. Future studies could usefully focus on proteins lacking functional annotation that are strongly changed in abundance in pseudoschizonts. Finally, contribution to parasite cell death could be explored for substrates identified here that have no annotated function, and our chemical tools will facilitate discovery of novel biology underlying these proteins.

Taken together, our results provide new insights into the scope and role of protein *N*-myristoylation in *Plasmodium*, and show for the first time that NMT is a chemically tractable target in the symptomatic stage of the most important human malarial parasite.

Methods

This section contains key experiments only. An extended experimental section is provided in the Supplementary Methods. Procedures used in this study were approved by the United Kingdom Home Office and carried out in compliance with European Directive 86/609/EEC for the protection of animals used for experimental purposes and in accordance with the United Kingdom "Animals (Scientific Procedures) Act 1986".

Chemical probes

Synthesis of YnMyr, YnPal, AzTB and YnTB has been described previously^{20, 27}.

AHA was purchased from Iris Biotech. Synthesis of other compounds is described in the Supplementary Methods.

Parasite culture

Parasites were cultured as described previously and schizont stages purified using magnetic columns (MACS),⁴⁸ before returning to culture for 2-4 h and synchronizing the newly invaded ring-stage parasites with 5 % D-sorbitol,⁴⁹ one cycle prior to use.

YnMyr tagging and labeling

For tagging, assays were set up using parasites at 40 h post invasion (based on initial Giemsa-stained smear observations) and diluted to ~10 % parasitemia. A haematocrit of 4 % was used to increase the total number of parasites harvested. To each culture the required volume of YnMyr, YnPal, AHA or CHX was added from stocks, along with any inhibitors. DMSO controls were set up in parallel. Cultures were re-incubated (37 °C) for a further 5 h. Parasite cultures were pelleted and the red cells lysed in 1 × culture volume of 0.15 % saponin in PBS. Pellets were washed

twice in the same buffer to remove red cell proteins. Parasite proteins were extracted using 1 % Triton X-100 in 10 mM Na₂PO₄ pH 8.2 with protease inhibitors (EDTA-free, Roche) and incubated overnight at 4 °C with rotation. Extracts were pelleted and the concentration of protein in the supernatant determined by DC protein assay (Bio-Rad).

CuAAC bioorthogonal ligation and pull-down of labeled proteins was carried out essentially as previously described²¹; full details are given in Supplementary Methods, along with details of in-gel fluorescence imaging and immunoblot analysis.

Proteomic identification of YnMyr tagged proteins by LC-MS/MS

Following CuAAC ligation and pull-down, samples were reduced, alkylated and digested on-bead with trypsin, as described in the Supplementary Methods. LC-MS/MS was carried out using an RSLCnano HPLC system (Dionex, UK) and an LTQ-Orbitrap-Velos mass spectrometer (Thermo Scientific). Data were further processed using Mascot (version 2.2.04, Matrix Science Ltd.) and Scaffold (version 3.6.02, Proteome Software). Protein identifications were required to contain at least 2 peptides exceeding 95 % probability giving a calculated protein false discovery rate of 0.0 %. Further experimental details and a description of data analysis are given in the Supplementary Methods.

Proteomics analysis of inhibited parasites

Equal protein quantities in whole cell lysate of parasites (45 h PI) cultured in presence of NMT inhibitor (1 μ M **1a**) or DMSO vehicle were processed by FASPTM, ⁵⁰ subjected to tryptic digest and analyzed by nanoLC-MS/MS (n = 4). Protein identification (>99% confidence) and label-free quantification in MaxQuant produced

a list of 1900 parasite proteins; data were filtered in Perseus v1.4.1.3 for completeness (minimum 3 quantifications per protein per group) yielding 1181 proteins for which relative abundance was determined using a two-tailed t-test (FDR = 0.05, 250 randomizations).

Supplementary Methods

Synthesis and characterization of compounds, protein crystallography, parasite assays, *in vivo* experiments, pharmacokinetics analysis, immunofluorescence and immunoblot analysis and further details of CuAAC labeling, gel analysis and LC-MS/MS proteomics is described in the Supplementary Methods. Substrate kinetic and inhibition assays for NMT were carried out essentially as described previously;^{25,} details are given in the Supplementary Methods.

Accession codes

Primary accessions (Protein Data Bank): 2YNC (PvNMT-YnMyrCoA); 2YND (PvNMT-NHMCoA-1a); 2YNE (PvNMT-NHMCoA-2a).

Referenced accessions (Protein Data Bank): 1IID (ScNMT-NHM-GLYASKLA), 4A95 (PvNMT-NHM), 2WUU (LdNMT-NHM), 2P6E (ScNMT-MyrCoA), 2WSA (LmNMT-MyrCoA-1a), 3IWE (HsNMT1-MyrCoA-1a).

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Author Contributions

M.H.W. designed and executed the chemical proteomics experiments, performed experiments on parasite samples and analyzed data. M.G., D.K.M., K.R. and B.C. produced synchronized parasites and performed parasite culture. D.K.M, B.C. and K.R. produced protein specific antibodies and K.R. and B.C. performed immunofluorescence microscopy under the guidance of A.A.H.. M.D.R., E.W.T. and R.J.L. designed series 2, and M.D.R. synthesized compounds **2a-b**. J.A.B. prepared recombinant proteins, performed crystallization experiments and determined X-ray structures, in collaboration with A.J.W. A.R.B. assisted M.H.W. with aspects of proteomic data generation and analysis. W.P.H. and R.A.S. synthesized compounds

1a-c. M.B., E.W.T. and R.A.S. designed and synthesized reagent **AzKTB**. R.A.S., E.W.T. and M.H.W. performed nanoLC-MS/MS experiments, proteomic identification of modified peptides and whole proteome analyses. R.T. and D.B. RT and DB performed *in vivo* analysis using the rodent malaria model. E.W.T. conceived and designed experiments, analyzed data and directed the overall collaboration. M.H.W., A.A.H. and E.W.T. co-wrote the manuscript, with comments and contributions from all the authors.

Additional information

Supplementary information and chemical compound information are available in the online version of the paper. Reprints and permissions information is available online at www.nature.com/reprints. Correspondence and requests for materials should be addressed to E.W.T.

Competing Financial Interests Statement

The authors declare no competing financial interests.

Figure legends

Figure 1: YnMyr-CoA is an effective substrate mimic for *P. falciparum* and *P. vivax N*-myristoyltransferases. (a) Co-translational *N*-myristoylation by NMT and Myr-CoA at an N-terminal glycine. (b) YnMyr tag & label strategy: YnMyr is fed to parasites in culture and transferred to substrate proteins by metabolic incorporation. Proteins are isolated and ligated by copper (I) catalyzed azide-alkyne cycloaddition (CuAAC) to a multifunctional capture reagent, followed by enrichment and downstream analysis. The cleavable linker is an optional module in the capture reagent that enables direct identification of the modification site by MS/MS. (c) The structure of YnMyr-CoA in PvNMT was determined by X-ray crystallography to 1.75 Å resolution (PDB 2YNC). At left: overall structure of PvNMT showing YnMyr-CoA (cyan), with a peptide substrate (PDB 1IID) modeled in (green) following least-squares superposition of the respective protein backbone atoms in PyMOL (DeLano Scientific). At right: key interactions made by YnMyr-CoA in the PvNMT acyl-CoA binding site, showing accommodation of the YnMyr chain in the hydrophobic acyl binding pocket and interactions of the thioester carbonyl.

Figure 2: YnMyr tags *N*-myristoylated and GPI-anchored proteins in blood stage *P. falciparum*. (a) YnMyr was fed to parasites in culture, ligated by CuAAC to AzTB and a portion subjected to base (0.2 M NaOH) or hydroxylamine (1 M NH₂OH) treatment to differentiate amide and ester-linked YnMyr. Samples were analyzed by in-gel fluorescence (top) or enriched by pull-down on streptavidin beads and analyzed by Western blot (WB; bottom). MSP1 is detected as the processed ~45 kDa GPI-anchored C-terminal fragment. (b) Summary of N-terminal glycine-containing proteins found by gel-based proteomics following base treatment and pull-down on streptavidin beads. Proteins are grouped by the gel slice in which they were

predominantly found and dominant proteins in each slice are highlighted in bold. (c) Confirmation of the major fluorescent band as ARF1 by Western blot against a novel anti-ARF1 antibody, analyzed before and after pull-down on streptavidin beads; a shift to higher molecular weight following addition of AzTB is observed only for tagged ARF1 (marked by arrow), which precisely overlays with the fluorescent band both before and after pull-down.

Figure 3: Chemical proteomic discovery and direct detection of modification site for *N*-myristoylated and GPI-anchored proteins in blood stage *P*. *falciparum*. (a) Assigned spectra profile (vertical log scale) of proteins identified from a representative gel-free analysis +/- base-treatment of samples, showing those proteins predicted or known to be GPI-anchored (blue bars), those containing an N-terminal glycine indicative of *N*-myristoylation (red bars), and those for which N-terminal YnMyr was detected directly by MS/MS (following ligation to AzKTB); other proteins are shown in white. (b) Structure of capture reagent AzKTB. (c) Functional annotation of 31 potential NMT substrates: all substrate proteins carry a base-insensitive modification, an N-terminal glycine, and evidence of N-terminal YnMyr and/or bioinformatic prediction of *N*-myristoylation.

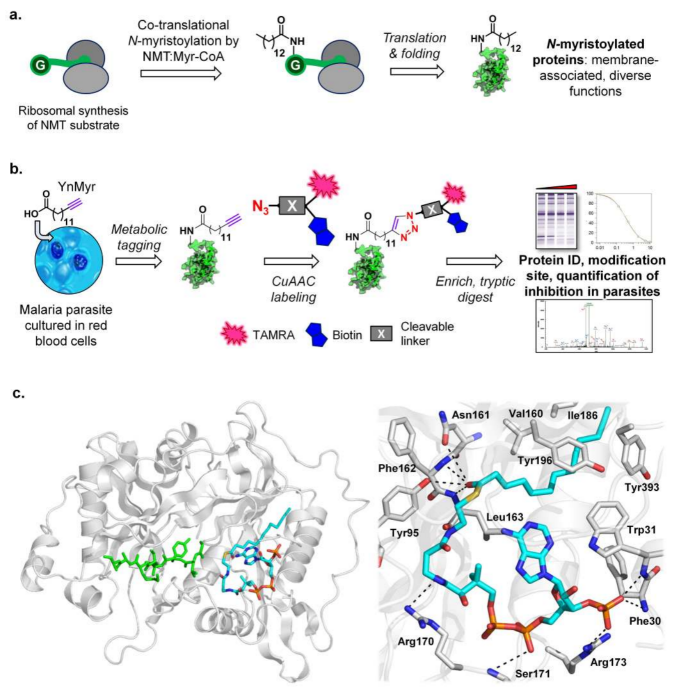
Figure 4: Compounds 1a-c and 2a-b bind in the *Plasmodium* NMT peptide binding pocket and inhibit the recombinant enzyme. (a) Structures and $K_i^{\text{(app)}}$ (apparent inhibitor dissociation constant) against recombinant PfNMT and PvNMT, and the host enzyme HsNMT1, for series 1a-c and 2a-b; IC₅₀ values were determined by a fluorogenic assay⁵¹ and differences in substrate K_m accounted for using the Cheng-Prusoff equation to give a value for apparent K_i . Errors are quoted \pm standard error ($n \ge 3$). (b) Structures for 1a and 2a in PvNMT were determined by X-ray crystallography with a non-hydrolyzable myristoyl-CoA (NHM) cofactor to 1.89 or

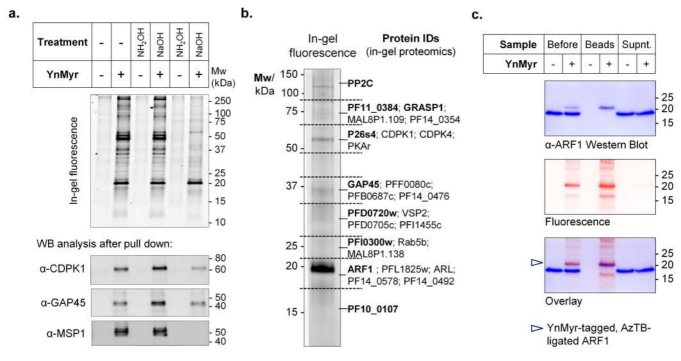
1.72 Å resolution (PDB 2YND and 2YNE), respectively. Left: interactions made by **2a** (green) in the peptide binding site showing a water-mediated hydrogen bond to Ser319, an ionic bond to the C-terminal acid (Leu410) and interactions with aromatic residues Tyr334 and Tyr211. NHM is shown in cyan. Right: overlay of binding mode of **1a** (yellow) and **2a** (green) in PvNMT, following alignment of the protein backbone in PyMOL (DeLano Scientific).

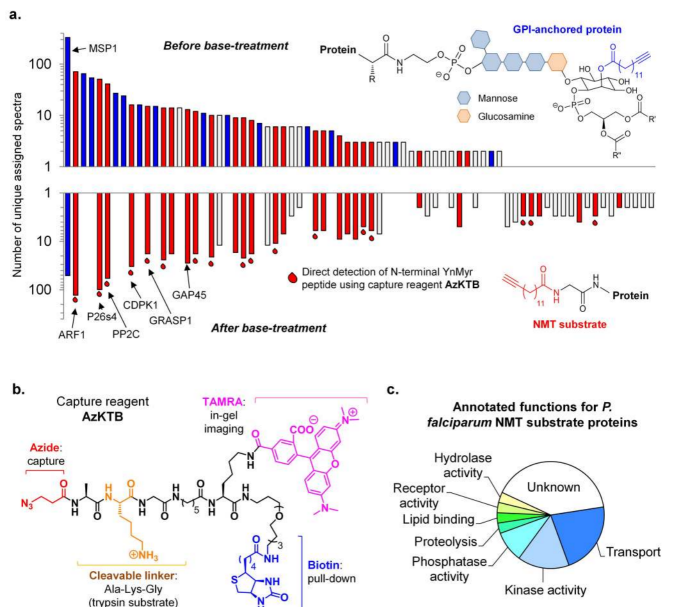
Figure 5: Inhibition of NMT in *P. falciparum* blood stage parasites leads to antiparasitic activity, and reduces parasitemia in an *in vivo* model of malaria. (a) Left: EC₅₀ (inhibitor concentration required for 50% reduction in parasitemia), $EC_{50}/K_i^{(app)}$, TC_{50} (inhibitor concentration required for 50% reduction in YnMyr tagging) and EC₅₀/TC₅₀ for **1a-c** and **2a-b** against *P. falciparum*. Right: correlation between EC₅₀ and TC₅₀ for **1a-c** and **2a-b**. K_i is also plotted. Horizontal and vertical error bars show \pm standard error ($n \ge 2$). (**b**) Quantification of NMT inhibition in live parasites. Schizonts were treated with 1a (0.13-10 µM illustrated by gold shaded gradient, or DMSO) followed by tagging with YnMyr (or DMSO). Left: analysis by ingel fluorescence following base treatment to remove GPI-anchored proteins. ARF1 band (marked) was used to calculate TC₅₀; Coomassie and α-BiP are loading controls for gel and Western blots, respectively. Right: Analysis by Western blot before and after pull-down of YnMyr-tagged; all NMT substrates show dosedependent reduction in enrichment, whilst MSP1 (GPI-anchored) does not. Full blots in Supplementary Information. (c) Quantification of normalized fluorescence and Western blot for ARF1, using GraFit 7.0 (Erithacus Software Ltd, UK). (d) Compound **2a** dosed at 50 mg/kg b.i.d. by intraperitoneal injection (i.p.) (n = 3 animals in)inhibitor-treated group; n = 2 in control group) reduces parasitemia in the *P. berghei* mouse model of malaria (one-tailed t-test; * p<0.05, ** p<0.01). An error bar is

missing in the 3 day control because one animal was euthanized due to severe symptoms of malaria.

Figure 6: NMT inhibition results in loss of the inner membrane complex and irreversible loss of parasite viability prior to parasite egress. (a) Top: P. falciparum intraerythrocytic life cycle showing approximate timings of life stages; parasite egress occurs ~45h post-invasion (PI). Bottom: mean number of nuclei per infected red blood cell (iRBC) 45h PI, +/- inhibitor 1a; error bars show ± standard deviation from mean (n = 50). (b) Label-free quantification (LFQ) of change in protein abundance plotted against significance of change for 1181 proteins in pseudoschizonts (NMT inhibition) vs. schizonts (DMSO), 45h PI; see **Supplementary Table S9** for dataset; groups of proteins are highlighted by function. (c) Western blot of parasite proteins 45h PI, cultured in the presence of NMT inhibitor (1 μM **1a**) or DMSO vehicle, showing depletion of IMC and motor proteins; BiP: loading control. (d) Indirect immunofluorescence microscopy of parasites 35h or 45h PI, cultured in presence of NMT inhibitor (1 μM **1a**) or DMSO; green: secondary antibody fluorescence; blue: DAPI (nuclear stain); all images at identical magnification, white bar = $5 \mu m$. (e) Synchronized parasites were cultured with NMT inhibitor (1 µM 1a) or DMSO vehicle to 45 h PI, whereupon medium was exchanged for medium containing either NMT inhibitor (1 µM 1a) or DMSO (45h wash-out), and parasite life stages analyzed at 55 h PI (p = 0.00081, two-tailed t-test; n = 3).







a.

	Enzyme		PVNMT	PfNMT	HsNMT1
Cpd	х	R	K _i ^(app) /nM		
1a	NMe	Н	32 ±2	66 ±8	8 ±0.5
1b	NMe	CHF ₂	42 ±4	101 ±20	23 ±4
1c	0	Н	1288 ±364	2373 ±256	760 ±74
2a	NMe	_	2.7 ±0.2	8 ±1	60 ±3
2b	NH	_	75 ±5	55 ±3	1541 ±90

b.

