Non-canonical metabolic pathways in the malaria parasite detected by isotope-tracing metabolomics

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Non-canonical metabolic pathways in the malaria parasite detected by isotope-tracing metabolomics

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Abstract

The malaria parasite, Plasmodium falciparum, proliferates rapidly in human erythrocytes by actively scavenging multiple carbon sources and essential nutrients from its host cell. However, a global overview of the metabolic capacity of intraerythrocytic stages is missing. Using multiplex 13C-labelling coupled with untargeted mass spectrometry and unsupervised isotopologue grouping, we have generated a draft metabolome of P. falciparum and its host erythrocyte consisting of 911 and 577 metabolites, respectively, corresponding to 41% of metabolites and over 70% of the metabolic reaction predicted from the parasite genome. An additional 89 metabolites and 92 reactions were identified that were not predicted from genomic reconstructions, with the largest group being associated with metabolite damage-repair systems. Validation of the draft metabolome revealed four previously uncharacterised enzymes which impact isoprenoid biosynthesis, lipid homeostasis and mitochondrial metabolism and are necessary for parasite development and proliferation. This study defines the metabolic fate of multiple carbon sources in P. falciparum, and highlights the activity of metabolite repair pathways in these rapidly growing parasite stages, opening new avenues for drug discovery.

Keywords: haloacid dehalogenase; mass spectrometry; metabolite repair; Plasmodium; SHMT

Subject Categories: Metabolism; Microbiology, Virology & Host Pathogen Interaction

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Introduction

Considerable progress has been made in reducing the incidence of malaria over the last decade, although the decline in malaria cases has stalled in recent years and resistance to frontline antimalarials is on the rise (WHO, 2019). Identifying new antimalarials with novel targets therefore remains a priority, and significant investment has been made in expanding the drug development pipeline with novel classes of antimalarials (Antonova-Koch et al, 2018; Hooft van Huijsduijnen & Wells, 2018). Metabolic enzymes and metabolite transporters are direct or indirect targets of most of the existing antimalarials and current lead compounds (Cowell et al, 2018; Ross & Fidock, 2019). However, the total number of enzymes/transporters that have been rigorously validated as drug targets remains small. A detailed understanding of the metabolism of the different developmental stages of the malaria parasite, Plasmodium falciparum, and the host cells within which they live is therefore necessary for informing the development of new antimalarial therapies. Plasmodium falciparum progresses through a number of different developmental stages during its life cycle in the Anopheles mosquito and its human host. Infection in humans is initiated by infective sporozoites that develop asymptomatically in the liver, resulting in the release of thousands of merozoites that initiate repeated cycles of infection and asexual replication in erythrocytes (i.e. red blood cells (RBCs)) that cause the clinical symptoms associated with malaria. The P. falciparum intraerythrocytic developmental cycle (IDC) takes approximately 48 h and involves the development of the metabolically-active trophozoite and schizont stages, followed by cell division of individual parasites into 16–32 new merozoites. The massive expansion of parasite biomass during development is fuelled by the uptake and catabolism of glucose, as well as a number of other essential nutrients (e.g. amino acids, purines and vitamins) that are either directly scavenged from the RBC or derived from breakdown of RBC haemoglobin and other proteins (Roth, 1990; Atamna et al, 1994; Liu et al, 2006; Olszewski et al, 2009). Considerable progress has been made in delineating key salvage and metabolic pathways involved in P. falciparum asexual development, which has formed the basis for genome-scale models of parasite metabolism (Fatumo et al, 2009; Plata et al, 2010; Bazzani et al, 2012; Tymoshenko et al, 2013). Despite these advances, > 40% of the protein-encoding genome remains unannotated and a significant fraction of annotated metabolic genes have yet to be assigned to specific metabolic pathways or reactions. In the phylum Apicomplexa, many genes have also been repurposed to fulfil non-canonical
functions, impeding genomic reconstructions and a systematic understanding of the total metabolic capacity of the pathogen (Oppenheim et al., 2014; Ke et al., 2015). Finally, enzyme promiscuity and side reactions can result in the production of unanticipated and novel metabolites that can have important roles in regulating cellular metabolism (Linster et al., 2013; Bommer et al., 2019; Dumont et al., 2019), further complicating predictions of enzyme function based on gene homology.

Defining the observable metabolic capacity of key developmental stages of *P. falciparum* and its host cell is required to verify the accuracy of genomic reconstructions and to identify unexpected metabolic pathways and gene functions. A number of approaches have been used to undertake a global analysis of the metabolic capacity of other organisms. For example, a system-wide reverse genetics approach was used to identify the metabolic function or indirect metabolic impact of each gene within *E. coli* (Fuhrer et al., 2017). The emergence of genome-wide disruption libraries in *P. falciparum* and *P. berghei* makes this approach theoretically possible (Bushell et al., 2017; Zhang et al., 2018). However, these studies have highlighted the essentiality of many metabolic enzymes in *Plasmodium* spp., limiting the effectiveness of this approach. Even viable but slow-growing mutants generated through such approaches are likely to be difficult to compare directly to parental parasites at a metabolic level. The converse approach – acquiring untargeted mass spectrometry data and verifying the “observed” metabolome – has not yet been fully exploited because of the lack of well-established pipelines for data filtering and metabolite identification. Current liquid chromatography–mass spectrometry platforms allow detection of >10,000 mass-to-charge (m/z) features, yet a significant majority (>90%) of these features correspond to background noise or degeneracy (Creek et al., 2011; Mahieu & Patti, 2017; Wang et al., 2019). The absence of autonomous methods for controlling the false discovery rate has hampered the compilation of an accurate metabolome for most organisms to date.

Here we use stable-isotope resolved metabolomics to prioritise m/z features corresponding to metabolites actively synthesised by *P. falciparum* or the host RBC (Huang et al., 2014; Sevin et al., 2017). Previous work has demonstrated the ability of this approach to define the extent of active 13C-glucose metabolism in RBCs (Srivastava et al., 2017), and here we expand this approach to ten biologically relevant 13C-substrates in *P. falciparum*-infected RBCs. Filtering for actively-labelled metabolites enabled >95% of m/z features to be removed, and the remaining m/z features were then identified and the active metabolome defined. This approach led to the identification of 577 metabolites in uninfected human RBCs and 911 metabolites in *P. falciparum*-infected RBCs corresponding to 41% coverage across the predicted metabolome of *P. falciparum* (the summation of all expected metabolites from all known pathways inferred from a genomic reconstruction irrespective of enzyme gaps). The pattern of stable-isotope labelling for each metabolite allowed us to further infer metabolic reactions corresponding to 70.5% coverage across predicted reactions in *P. falciparum*, with the mis-match between metabolite and reaction coverage largely due to a subset of metabolites participating in many reactions. Defining the “observed” metabolome without constraining the results to the expected composition inferred from genomic reconstructions revealed 89 metabolites and 92 reactions not predicted from genomic reconstructions. These studies have highlighted unanticipated complexity in *P. falciparum* metabolism, including the presence of active metabolite damage and repair systems in rapidly dividing parasite stages.

**Results**

**Global stable-isotope labelling filters for metabolites actively synthesised in uninfected and *P. falciparum*-infected human erythrocytes**

*Plasmodium falciparum* trophozoite-infected red blood cells (iRBCs) or uninfected RBCs (uRBCs) were metabolically labelled for 5 h in parallel cultures containing different 13C-labelled compounds. Metabolites were extracted and analysed in parallel by GC-MS, LC-MS polar and LC-MS apolar analytical platforms to maximise coverage of different metabolites classes. All mass-to-charge (m/z) features were extracted and untargeted isotopologue grouping performed to identify m/z features that correspond to metabolites that were differentially labelled between iRBCs and uRBCs (Fig 1A). Metabolites were provisionally identified based on METLIN database matching and their identities subsequently confirmed based on comparison with authentic standards, MS/MS matching, stable-isotope incorporation pattern and exact mass. The resulting list of metabolites and their respective labelling patterns were compiled into the “observed” metabolome of *P. falciparum* at the trophozoite stage (Dataset EV1).

As an example of the workflow, polar LC-MS analysis of 13C-glucose labelled iRBC extracts revealed that 859 of the original 33,691 m/z features detected in unlabelled iRBCs exhibited decreased intensity following 13C-glucose labelling (Fig 1B), indicating that they likely correspond to mono-isotopic masses (i.e. the unlabelled species) that decrease as metabolites become enriched for 13C atoms. These m/z features were then ranked by fractional enrichment and the exact mass of the unlabelled feature queried against the METLIN metabolite database (Fig 1C). Of the 859 m/z features, 410 returned putative matches within a 10 ppm tolerance for M-H precursors and are annotated as “presumptive” features. Presumptive features exhibited a broad range of fractional enrichments, ranging from 0.01 to 0.999, indicating no significant bias in annotating putatively labelled metabolites via stable-isotope enrichment. Highlighted is the presumptive feature m/z 275.0167 and its isotopologue group (Fig 1D), matching to three possible metabolites (2-carboxyarabinitol 5-phosphate, 2-carboxyarabinitol 1-phosphate and 6-phosphogluconate; all Δ2 ppm). MS/MS spectral matching, together with the predominance of 6 mass isotopomer in 13C-glucose labelled cells, confirmed the identity of this metabolite to be 6-phosphogluconate (Fig 1E). This procedure resulted in the identification of 232 polar metabolites that were significantly labelled with 13C-glucose and was repeated for all labelled substrates and MS platforms.

The observed metabolome of uninfected and *P. falciparum*-infected human erythrocytes

The consolidated list of all 13C-labelled metabolites detected on the three MS platforms was then incorporated into the observable metabolome for uRBC and iRBC (Fig 2A). To capture additional metabolites that were not labelled with any of the 13C-substrates tested, all m/z
Figure 1.
features from unlabelled uRBC and iRBC extracts were compared with the expected exact mass of all metabolites in the predicted *P. falciparum* and RBC metabolomes (Huthmacher et al., 2010). Putative matches were then verified as described above. The metabolic network reconstruction reported by Huthmacher and colleagues contains predictions of both host and parasite metabolic activity, incorporates literature and manual curation for higher accuracy predictions, and is structured for matching enzyme classification reactions and PlasmoDB IDs. The reconstruction contains 566 metabolites and 349 reactions in human erythrocytes and 1,622 metabolites and 998 reactions in *P. falciparum* (Fig 2A).

Following manual curation and verification, we compiled an “observable” metabolome of iRBCs and uRBCs, which comprised 911 and 577 metabolites, respectively. All metabolites were collapsed into unique KEGG IDs and compared to the predicted metabolome of each cell type (396 and 299 metabolites for iRBC and uRBC, respectively). 255 observed metabolites (with unique KEGG IDs) matched the predicted metabolome of iRBCs, corresponding to 41% coverage of the predicted metabolome, whereas 152 observed metabolites matched the predicted metabolome of uRBCs (36.3% coverage). Core metabolic pathways were statistically over-represented in the observable metabolomes of both iRBC and uRBC (Fig 2B), with consistently higher metabolite coverage across each pathway for iRBC (Dataset EV2).

Metabolites predicted from genome annotations but not observed in iRBC samples could reflect: (i) incorrect annotation of the genome (e.g. the annotated branched-chain amino acid degradation pathway is likely missing and the sole annotated enzyme in the pathway, BCKDH, is known to fulfil an alternative function; Oppenheim et al., 2014), (ii) down-regulation of the metabolic pathway during trophozoite development (e.g. *de novo* fatty acid biosynthesis is known to be down-regulated in the presence of exogenous fatty acids in the intraerythrocytic stages; Yu et al., 2008) or (iii) technical issues such as low metabolite abundance, sequestration by host/parasite proteins or incompatibility with the applied extraction or MS methods for detection (e.g. haem and ubiquinone biosynthesis). Pathway enrichment analysis of predicted metabolites that were not observed yielded no statistically-enriched pathways (Dataset EV2).

We were interested in defining which metabolites were uniquely detected in iRBCs or uRBCs. iRBCs contained 102 unique KEGG IDs corresponding to 339 metabolites that were not observed in uRBCs, with phospholipid and CoA biosynthetic pathways most enriched (unadjusted *P* = 0.017 and 0.033, respectively). These pathways are not active in human RBCs, but are required for the rapid growth of *P. falciparum* asexual stages (Fig 2B). Interestingly, five metabolites were only detected in uRBCs. These included N-acetylmuramylpentapeptide, sucrose, 5-formyl-tetrahydrofolate, glucosamine and pyridoxine-5-P. Pyridoxine-5-P is an essential vitamin/cofactor required for the activity of multiple enzymes in both cell types. The absence of detectable pyridoxine-5-P in iRBCs may reflect the sequestration of this cofactor by parasite enzymes (Fig 2C).

A significant number of observed metabolites did not match to the predicted metabolomes of iRBCs and uRBCs (141 and 147, respectively). These unpredicted metabolites did not statistically over-represent any conventional metabolic pathways (Dataset EV2). For example, 

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**Figure 1. Strategy for defining the metabolic capacity of *P. falciparum*-infected red blood cells (iRBCs) and uninfected red blood cells (uRBCs) using untargeted stable-isotope labelling.**

A Purified trophozoite-stage iRBCs and matched uRBCs were labelled with one of ten 13C-substrates (listed in purple). All mass-to-charge (m/z) features identified by LC-MS were extracted and isotope groups grouped. 13C-labelled and 13C-labelled isotope groups were compared to identify m/z features in each cell type and that were differentially labelled between iRBC and uRBC. Putative metabolite identities were then confirmed with authentic standards, MS/MS spectral matching, 13C-labelling profile and exact mass. The observed metabolic network of each cell type was then constructed using the draft metabolome and 13C-labelling information and compared to the predicted *P. falciparum* metabolome reported by Huthmacher et al. (2010).

B All m/z features detected from iRBC extracts plotted as the log2 ratio of their abundance detected in 13C-glucose versus 13C-glucose conditions. 859 m/z features (from a total 33,691 m/z features identified) were significantly altered in 13C versus 13C samples of iRBC. Highlighted in black is a single m/z feature (m/z 275.0167).

C The 859 significant m/z features (x-axis; putative M*+* species) were ranked by their fractional enrichment (y-axis) and the observed mass queried against the METLIN metabolite database. m/z features that returned a hit were classified as presumptive features (blue) and retained for validation. Presumptive m/z features displayed a wide range of fractional enrichments, indicating no systematic bias.

D Changes in the isotope group distribution in 13C-glucose and 13C-glucose labelled samples of m/z 275.0167. Data are presented as the mean 13C-fractional enrichment ± SEM from six biological replicates.

E MS/MS fragmentation of m/z 275.0167 matched the reference spectrum of 6-phosphogluconate, consistent with the exact mass and labelling pattern, confirming the metabolite identity. This approach was repeated for all presumptive m/z features.
to 100-fold and 78-fold, respectively, in trophozoite-stage-infected RBCs (Roth, 1990; Atamna et al., 1994), highlighting the lack of correlation between metabolite abundance and corresponding metabolic fluxes. The reduced pool size of glycolytic/PPP intermediates in iRBC may increase the sensitivity of parasite pathways to subtle changes in exogenous glucose levels by aligning substrate...
levels more closely to the respective $k_m$'s of rate-controlling enzymes (Bennett et al., 2009; Park et al., 2019).

The metabolic activity network of *P. falciparum* reveals a plethora of metabolic damage-repair systems

To further define the unpredicted metabolic activity of *P. falciparum* and the host RBC, all metabolites labelled with the different $^{13}$C-substrates were mapped to pathways to identify all observable reactions from both iRBC and uRBC. In all cases, the number of metabolites labelled with each tracer was higher in iRBCs compared to uRBCs, as was the complexity of the corresponding sub-networks (Fig 3A). Strikingly, the majority of the detectable metabolome of uRBCs was unlabelled, consistent with the loss of many enzymes and metabolic pathways in mature erythrocytes (Srivastava et al., 2017). In contrast, most metabolites detected in iRBC were labelled with one or more $^{13}$C-substrates (Dataset EV1), indicating a high level of redundancy and metabolic complexity in these intracellular parasite stages. Metabolites in iRBC that did not label with any $^{13}$C-substrate tested (183 compounds) mainly consisted of vitamins (e.g. pyridoxine and riboflavin), purines and specific lipid classes (e.g. sphingomyelins) for which the parasite is known to be auxotrophic and dependent on salvage from the host cell or media in the case of *ex vivo* culture.

We then matched labelled substrates to predicted reactions from the genome reconstruction. Either the substrate and/or the product of 87.4% of metabolic reactions predicted from the reconstructions was labelled with one or more $^{13}$C-tracers in iRBC (Fig 3B). The discordance between metabolite and reaction coverage (42.6% and 87.6% for iRBC, respectively) is partly due to a small minority of metabolites participating in a large number of reactions (Fig 3C). Removing the top ten detectable metabolites (ATP through to NADPH) from the analysis still gave 70.5% coverage across the theoretical reaction landscape of iRBCs. 104 reactions had no substrate/product detection and corresponded to metabolites that were incompatible with MS detection (low abundance or poorly ionised), such as intermediates of haem biosynthesis, superoxide metabolism and dolichol production (Dataset EV2).

We reconstructed the $^{13}$C-label tracing through the predicted metabolic network and incorporated additional reactions to accommodate labelled metabolites not predicted from the genome reconstruction (Appendix Figs S2–S6). These additional steps were based on the closest conversion from known metabolites and consistency of the $^{13}$C/$^{12}$C fraction label observed between putative substrate/product pairs. The $^{13}$C-glutamine reconstruction indicated operation of canonical pathways, such as the TCA cycle, pyrimidine biosynthesis and glutathione metabolism, but also $^{13}$C-labelling into unpredicted metabolites, indicating the presence of unannotated reactions (Fig 3D; highlighted in green). These unexpected reactions did not correspond to complete biosynthetic pathways, but were either additional steps to predicted pathways (e.g. conversion of y-aminobutyric acid into succinate semialdehyde) or potential metabolic side reactions leading to the production of non-canonical metabolites, such as 2-hydroxyglutarate, that need to be excreted or catabolised to prevent metabolic dysregulation (Lu et al., 2012; Intlekofer et al., 2017; Dumont et al., 2019).

Metabolic side products constituted the largest group of new and unanticipated metabolites detected in the $^{13}$C-network reconstructions. Of the 92 reactions that were not predicted, 36 corresponding to metabolic side or repair reactions. Using the network reconstructions, we refined the observed metabolites that were unpredicted to 89, corresponding to the highest confidence assignments (consistency with reaction network) (Dataset EV4). These “damaged” metabolites included methionine sulfoxide (which can be a marker of oxidative stress; Mashima et al., 2003), unconventional glutathione adducts (e.g. methyl- and succinyl-glutathione) and pipecolate which arises via lysine degradation.

The $^{13}$C-bicarbonate labelling network was generally in agreement with the predicted reactions (Appendix Fig S4) with the exception of labelling into the unconventional metabolite orotidine. Orotidine was also labelled with $^{13}$C-glucose and $^{13}$C-glutamine, consistent with synthesis via a side reaction of pyrimidine biosynthesis. No reaction has been reported for this metabolite, but it is possibly an overflow metabolite, generated via dephosphorylation of orotidine monophosphate. A number of non-phosphorylated intermediates of central carbon metabolism (mannose, galactose) were also labelled in $^{13}$C-glucose-fed iRBC, indicating synthesis and subsequent dephosphorylation of the cognate sugar phosphates (Dataset EV4). These data suggest that unanticipated metabolites can be generated by specific or promiscuous phosphatases. Candidate enzymes include members of the haloacid dehalogenase (HAD) family of enzymes that have been shown to regulate metabolic fluxes *in vivo* (Guggisberg et al., 2014; Guggisberg et al., 2018; Dumont et al., 2019).

**Haloacid phosphatases regulate parasite metabolic flux**

We wanted to explore the role of two uncharacterised HAD phosphatases given the important role this protein family plays in regulating metabolic flux and metabolite repair. We targeted PF3D7_1118400 (annotated as HAD4) and PF3D7_0803200 (annotated as a member of the HAD superfamily, hereon referred to as Lipin) for inducible disruption. Both endogenous loci were targeted via Cas9-mediated double-stranded break towards the centre of each gene and the 3’-half replaced with a recondensed sequence flanked by two loxP sites (Wilde et al., 2019). The 3’-end was also HA-tagged and a glmS ribozyme introduced in the 3’-UTR. Transfection of this construct into a 3D7 parasite line expressing an integrated copy of the dimerisable cre-recombinase then enabled inducible excision of the gene upon rapamycin addition or transcript degradation when glucosamine was added. Transfections were performed with line-arrised rescue DNA template and integration confirmed by PCR (Appendix Fig S7). HA-tagged proteins of the expected size were detected and were efficiently depleted following the addition of 100 nM rapamycin and 2.5 mM glucosamine (Fig 4A). A transgenic line with lactate dehydrogenase 1 (LDH1) under inducible disruption was used as a positive control. Disruption of Lipin significantly impaired parasite growth, whereas loss of HAD4 had no effect (Fig 4B). Loss of LDH1 led to specific metabolic changes, with increased abundance of pyruvate (the enzyme’s substrate) and reduced abundance of lactate (Fig 4C), demonstrating the validity of the approach for inferring enzyme function.

Loss of Lipin was associated with elevated levels of ceramides and lyso-phosphatidic acid (LPA) lipid species in iRBC (Fig 4D and Dataset EV5). Sphinganine-1-P and sphingosine-1-P were also elevated, suggesting that this enzyme dephosphorylates diverse lipid.
Figure 3. Reconstruction of the active metabolic networks in *P. falciparum* trophozoite stages.

A The sum of metabolites labelled with each $^{13}$C-substrate in iRBCs and uRBCs. Dark grey indicates metabolites that were $^{13}$C-labelled in both cell types, whereas light grey represents metabolites $^{13}$C-labelled in either iRBCs or uRBCs.

B The number of predicted metabolic reactions in each cell type, the number of reactions with an observed substrate, an observed product, and the number of predicted reactions with no observed substrate or product (unobserved). The same analysis is reproduced following removal of the top ten cofactors present in the observed metabolome.

C Predicted reactions were deconstructed into individual metabolites and were ranked according to their frequency across all predicted reactions. A small subset of metabolites participate in a large number of reactions. The inset includes ranked metabolites with a frequency (y-axis) $\geq$ 20 from all predicted reactions. Black indicates metabolites present in the observed metabolome.

D Reconstruction of the $^{13}$C-glutamine observed metabolic network. Green metabolites correspond to observed metabolites that are not predicted from the genomic reconstruction. Black arrows indicate where the $^{13}$C-incorporation is consistent with predicted reactions. Green arrows indicate where the observed labelling pattern is inconsistent with predicted reactions, and proposed reactions are annotated in Dataset EV4.
substrates \textit{in vivo}. This phenotype is consistent with the function of Lipin domain-containing proteins in other eukaryotes, in regulating intracellular pools of phosphatidic acid and diacylglycerols, and flux of these lipids into other pathways of bulk lipid synthesis (phospholipid, triacylglycerol). Interestingly, infection with wild-type \textit{P. falciparum} leads to a depletion of phosphatidic acid and accumulation of diacylglycerol in iRBC (Gulati \textit{et al}, 2015). This likely reflects the need for increased phospholipid biosynthesis in rapidly growing intraerythrocytic stages, the sequestration of host lipids as triacylglycerols (TAGs) within the parasite, and the pivotal role of phosphatidic acid/DAG signalling in microneme secretion and invasion (Bullen \textit{et al}, 2016). The pleiotropic role of Lipin in these different pathways likely accounts for the severe growth defect observed when Lipin expression was inducibly reduced in this study.

HAD4 disruption led to an accumulation of several nucleotides and glycolytic intermediates which feed into lipid metabolism.
Figure 4. Inducible disruption of haloacid dehalogenase 4 and a putative Lipin.

A Anti-HA Western blots for each protein targeted for disruption with a detectable band corresponding to the expected product size. 100 nM rapamycin and 2.5 mM glucosamine were added to induce protein depletion for either one cycle (LDH1) or three cycles (Lipin and HAD4). Anti-PFβiP was used as the loading control.

B Growth of DiCre-ΔD7 transgenic lines following LDH1, HAD4 and Lipin depletion with 100 nM rapamycin and 2.5 mM glucosamine was assessed relative to uninduced parasite cultures. The untransfected DiCre-ΔD7 parasite line was used as a negative control. Data are presented as the mean ± standard error of the mean (SEM) from three independent replicates.

C Untargeted LC-MS analysis of LDH1 depletion (one cycle of rapamycin and glucosamine treatment) indicated minimal but selective metabolic changes. m/z feature intensities are plotted as the log, ratio of treated/untreated and the Benjamini–Hochberg-corrected P values across six biological replicates plotted as –log10(P). Below, LDH1 substrate and product (pyruvate and lactate, respectively) abundance plotted as the ratio of treated/untreated from biological replicates (mean ± SEM) and the parental DiCre line presented as the negative control.

D Loss of Lipin (PF3D7_0503200) leads to accumulation of various lipid species (three cycles of rapamycin and glucosamine treatment). Data are presented as the mean log, fold change Lipin-depleted (treated) enriched trophozoite-stage iRBCs to untreated controls from three to six biological replicates (± SD). The schematic depicts the proposed lipid phosphatase activity of Lipin.

E Loss of HAD4 leads to increases in intracellular levels of several nucleotides and intermediates in lower glycolysis. Data are represented as the mean log, fold change of enriched trophozoite-stage iRBCs treated for three cycles compared to untreated controls from three/four biological replicates. Three nucleotide–nucleoside pairs are depicted in the top inset panels, and HAD4 disruption leads to accumulation of the nucleotide monophosphate and depletion of the nucleoside. The schematic depicts lower glycolysis and triose-phosphate interconversion, with the corresponding metabolite levels following HAD4 disruption depicted in the lower right inset.

F Phosphatase activity against the generic substrate pNPP of wild-type HAD4, catalytically inactive HAD4S21G and a no-enzyme (No Enz) control, presented as the mean ± SEM (three biological replicates).

G Inhibition of HAD4 phosphatase activity by inorganic phosphate. Phosphatase activity against the generic substrate pNPP as the mean ± SEM from three independent experiments. The inorganic phosphate IC_{50} of HAD4 is 21 ± 0.21 mM.

H Substrate specificity of HAD4. Substrates are divided into nucleotide phosphates (black) and other metabolites such as sugar phosphates and vitamins (white). Presented is the enzyme activity from three independent experiments mean ± SEM. Substrate abbreviations are listed in methods.
generates precursors for de novo biosynthesis of the major phospholipid species (phosphoethanolamine and phosphocholine; Witola et al., 2008). However, 13C-serine labelling suggested that ethanolamine-P may also be directly converted to glycerol-ethanolamine-P rather than being generated via the canonical pathway of PS and PE lipid breakdown (Fig 6A). The identity of the serine phosphate decarboxylase involved in synthesis of ethanolamine-P or the enzyme involved in generating glycerol-ethanolamine-P remains to be discovered (Rontein et al., 2003; Liu et al., 2018).

The 13C-serine tracing network also included intermediates in one-carbon folate metabolism. Entry of serine into these pathways is mediated by the enzyme, serine hydroxymethyltransferase (SHMT), which converts serine to glycine with the transfer of one-carbon to tetrahydrofolate. *P. falciparum* encodes two SHMT isoforms: a well-characterised cytosolic isoform (SHMT-C) and a mitochondrial isoform (18% identity to SHMT-C; Maenpuen et al., 2009; Pang et al., 2009; Read et al., 2010). SHMT-C participates in general one-carbon folate metabolism, necessary for dTTP production for DNA synthesis and formylation reactions (Fig 6B). *P. falciparum* lacks the mitochondrial enzymes involved in folate metabolism found in other eukaryotes and no in vitro activity could be detected for SHMT-M (Pang et al., 2009), leaving the role of the SHMT-M isoform undefined.

To investigate the function of SHMT-M (PF3D7_1456100) and mitochondrial one-carbon metabolism in more detail, we confirmed the localisation of HA-tagged SHMT-M to the mitochondrion (Fig 6C) (Read et al., 2010; Pornthanakasem et al., 2012). Conditional disruption of SHMT-M in asexual parasite stages was associated with changes in intracellular levels of many metabolites (Fig 6D and E), in sharp contrast to the limited metabolic changes observed when the essential enzyme LDH1 was depleted (Fig 4C). In particular, loss of SHMT-M was associated with significant changes in intermediates in glycolysis, the pentose phosphate pathway and nucleotide metabolism (Fig 6F; Dataset EV6), suggesting that SHMT-M is fundamental for central carbon metabolism. To differentiate between the possibility that these changes were directly connected to SHMT-M function versus a generalised death phenotype, protein turnover rates were assessed using a recently developed pulse-SILAC method in SHMT-M-depleted parasites (Yang et al., 2019). Parasites were labelled with 13C15N-isoleucine for 5 h and the turnover of parasite proteins assessed by measuring the ratio of labelled (newly synthesised) and unlabelled peptides (nascent) in corresponding tryptic peptides (Fig 7A). Similar rates of protein turnover (R2 = 0.92) were observed before or after knock-down of SHMT-M (Fig 7B; Dataset EV7), indicating that parasite protein homeostasis was unperturbed when SHMT-M is disrupted and that the metabolic phenotype is unlikely to be caused by decreased cell viability. Significantly, nuclear-encoded mitochondrial proteins and one of the three mitochondrial ribosomes (cytochrome b) also exhibited similar rates of turnover before and after SHMT-M knock-down, suggesting that the metabolic phenotype does not reflect marked changes in mitochondrial protein import or translation. This finding indicates
Figure 6. The architecture and function of the serine metabolic network.

A. 13C-serine tracing into the metabolic network of iRBC. Black arrows indicate reactions predicted from genomic reconstruction, whereas green arrows indicate reactions that are not predicted but best match the stable-isotope labelling pattern observed. Proposed reactions are annotated below. Metabolites in grey were not observed via mass spectrometry, and green indicates a metabolite observed but not predicted. Highlighted in orange is the glycine cleavage system, which is predicted to be active but has not been verified.

B. One-carbon folate metabolism in P. falciparum. The parasite encodes all the enzymes necessary for a complete cytosolic pathway but only appears to possess a single enzyme – serine hydroxymethyltransferase – in the mitochondrion. Highlighted in white are the conventional enzymes and reactions of one-carbon metabolism in the mitochondria of other eukaryotes.

C. HA-tagged SHMT-M detection via immunofluorescence microscopy. DAPI was used to visualise the nucleus, and MitoTracker CMXros used to illuminate the mitochondrion. Anti-HA signal partially colocalises with the mitochondrion.

D. Addition of 100 nM rapamycin and 2.5 mM glucosamine for three growth cycles (+) leads to depletion of the HA-tagged SHMT-M.

E. Depletion of SHMT-M leads to significant changes in intracellular levels of many metabolites in iRBC. m/z features are plotted as the log2 ratio of treated/untreated intensities and the Benjamini–Hochberg-corrected P values across six biological replicates plotted as −log10(P).

F. Pathway enrichment analysis of the m/z features identified as significantly different when SHMT-M is depleted. Depicted are the Gene Set Enrichment Analysis (GSEA) P values and Mummichog P values generated via Metaboanalyst 3.0, with the combined P value from each pathway enrichment approach represented by colour.
Figure 7. SHMT-M is required for mitochondrial function, mETC maintenance and asexual growth.

A. *P. falciparum*-infected cultures were incubated for 5 h with $^{13}$C$^{15}$N-isoleucine (heavy). Following protein extraction and trypsin digestion, each isoleucine-containing peptide exists as a heavy (H) and light (L) form, corresponding to newly translated and nascent pools, respectively.

B. The log$_{2}$ H/L ratio reflects the turnover (summation of synthesis and degradation) for each protein detected between the control (untreated) and SHMT-M-depleted conditions (three growth cycles of rapamycin and glucosamine treatment). Data represent the mean log$_{2}$ H/L ratio across three biological repeats.

C. One-carbon folate intermediates are plotted as the log$_{2}$ ratio of their abundance detected in treated (100 nM rapamycin/2.5 mM glucosamine treatment for three growth cycles) compared to untreated in the DiCre parental line (C) and SHMT-M (SM) (mean ± SEM).

D. Pyrimidine biosynthetic intermediates accumulate when SHMT-M is disrupted compared to the DiCre parental line (C) following rapamycin/glucosamine treatment (mean ± SEM). Pyrimidine biosynthetic disruption is linked to dihydroorotate dehydrogenase (DHODH) inhibition and blockage of the mitochondrial electron transport chain (mETC). Ubiquinone (Q) is the necessary electron carrier for the mETC.

E. Asexual growth when SHMT-M is depleted. Parasitaemia was determined across seven replication cycles (x-axis) via flow cytometry and presented as the mean ± SEM from three independent experiments. Decylubiquinone (Dub) was added at 10 µM to determine if the growth defect observed was due to impaired ubiquinone synthesis or recycling.
SHMT-M is not necessary for mitochondrial genome replication (via dTTP synthesis) or tRNA formylation (Fig 6B), in contrast to the function of SHMT2 in mammalian cells (Ducker et al, 2016; Minton et al, 2018; Morscher et al, 2018).

Significantly, knock-down of SHMT-M was associated with changes in one-carbon folate intermediates (Fig 7C), indicating a direct function in one-carbon metabolism. Significant changes were also observed in intracellular levels of pyrimidine biosynthetic intermediates (Fig 7D) which mimicked the metabolic phenotype observed following inhibition of the mitochondrial electron transport chain (mETC) (Ke et al, 2015; Cobbold et al, 2016). Loss of SHMT-M could lead to loss of ubiquinone – which requires a source of one-carbon units – and is crucial for mETC function. However, attempts to rescue the growth defect of SHMT-M-deficient parasites by supplementation with the ubiquinone analogue, decylubiquinone (Ke et al, 2011), were unsuccessful (Fig 7E), suggesting that impaired ubiquinone synthesis is not the primary cause of the growth defect.

We further investigated whether SHMT-M might be in involved in generating one-carbon intermediates for cytoplasmic pathways by labelling iRBC with 2,3,3-D-serine and quantifying the amount of label exported into the cytosol for dTTP synthesis. Negligible amounts of dTTP + 1 isotope were detected after 2,3,3-D-serine labelling, indicating minimal export of one-carbon units from the mitochondrion into the cytosol (Appendix Fig S8A), consistent with a lack of a formate transporter at the mitochondrial membrane. SHMT-M is also likely not required for the mitochondrial glycine cleavage system, as 13C-glycine labelling contributes < 0.1% of the total dTTP pool (Appendix Fig S8B). Taken together, these data suggest that P. falciparum SHMT-M may be required to fuel down-stream fluxes, such as the conversion of 5,10-methyl-THF to 10-formyl-THF which can be a major source of NADPH production in eukaryotic cells (Fan et al, 2014). A key role in maintaining mitochondrial redox balance would account for the impact of SHMT-M loss on mETC activity, pyrimidine biosynthesis and parasite viability.

Discussion

We have developed a new approach for measuring de novo synthesised and salvaged metabolites in asexual stages of P. falciparum, achieving over 70% coverage of all metabolic reactions predicted from P. falciparum genome-wide metabolic reconstructions (after exclusion of the top 10 most promiscuous metabolites). We show that asexual parasite stages constitutively salvage a wide range of different carbon sources and nutrients from the host which are assimilated by overlapping anabolic and catabolic pathways, likely contributing to the robustness of parasite metabolism in the RBC. These analyses allowed the detection of over 80 additional reactions and metabolites that are not predicted from gene annotations. Many of these new reactions correspond to additional steps in predicted pathways, or apparent promiscuous-enzyme activity and damage-repair pathways. It is increasingly clear that damaged metabolites have an impact on normal metabolic function – via competitive inhibition or allosteric regulation of key enzymes in central carbon metabolism – and organisms require effective means to detoxify or repair them (Linster et al, 2013; Bommer et al, 2019; Dumont et al, 2019). Exploring the function and impact of the 89 unpredicted metabolites identified in this study will likely reveal an increasingly complex metabolic arrangement and damage-repair enzymes necessary to maintain parasite metabolism. For example, the 13C-lysine network included pipecolate which is normally produced during lysine degradation (Appendix Fig S2). P. falciparum lacks the genes for a complete lysine degradation pathway but does possess a putative saccharopine dehydrogenase. The subsequent intermediate, aminoadipate semialdehyde, spontaneously cyclises to piperidine-6-carboxylate which competes for pyrroline-5-carboxylate reductase (Fuji et al, 2002; Struys & Jakobs, 2010; Linster et al, 2013). A pyrroline-5-carboxylate reductase is encoded in the genome of P. falciparum and we observed conversion of 13C-arginine into 13C-proline consistent with its activity. Curiously, the pyrroline-5-carboxylate reductase itself appears to be essential in P. falciparum (Zhang et al, 2018), but essentiality of arginine/proline interconversion seems questionable given the excess of each amino acid liberated during haemoglobin catabolism. It is plausible that detoxification of piperidine-6-carboxylate is a necessary function for maintaining parasite metabolism and a primary function of the enzyme (Linster et al, 2013).

This dataset comprehensively defines the observable metabolome of uninfected and P. falciparum-infected erythrocytes and by using multiple isotopic substrates, enabling the activity of over 70% of predicted reactions to be monitored. We selected the trophozoite stage of asexual development, primarily to capture the greatest number of active reactions at a single stage of development. Investigating other asexual stages may yield additional reactions as well as the process of host metabolome reorganisation. Here we identified five unique metabolites present in uninfected human erythrocytes. Whether these metabolites are actively depleted by the parasite in iRBC remains unclear. Moreover, this work provides a baseline for understanding how environmental factors alter parasite metabolism. For example, phosphate levels in human plasma vary during the day (Lederer, 2014), fluctuate with diet (Goretti Penido & Alon, 2012), and disease severity (Lewis, 1987; Davis et al, 1991; Suen et al, 2020). Future studies investigating how parasite metabolism responds to fluctuations in plasma phosphate will be of great interest. In addition, the growing evidence that lipid metabolism influences gametocyte commitment necessitates a deeper understanding of how plasma lipids fluctuate during disease progression and how the parasite responds to these triggers and initiates gametocyte commitment (Branucci et al, 2017).

Our attempt to validate the draft metabolome – and the associated reaction network captured via stable-isotope labelling – led us to target several uncharacterised enzymes for inducible disruption. Each of the disrupted genes/proteins contained predicted enzymatic domains, are not part of complete pathways, or are predicted to be essential from genome-wide disruption studies (Bushell et al, 2017; Zhang et al, 2018). Serum hydroxymethyltransferase is a key enzyme in folate recycling, and the cytosolic version has been well characterised (Maenpuen et al, 2009; Pang et al, 2009; Read et al, 2010). In contrast, the function of the mitochondrial-targeted SHMT has not been defined (Pang et al, 2009), and Plasmodium spp. lack many of the accessory enzymes needed for mitochondrial folate metabolism. In other eukaryotes, SHMT-M and mitochondrial folate metabolism is important for providing methyl-groups for tRNA modification and effective mitochondrial protein translation and also for providing the precursors for mitochondrial DNA replication (Anderson et al, 2011; Morscher et al, 2018). Here we demonstrate that mitochondrial protein turnover is unaffected when SHMT-M is
depleted, consistent with earlier studies suggesting that mitochondrial tRNAs are modified in the cytosol prior to import to the mitochondrion (Pino et al., 2010). We also show that ubiquinone does not require methyl-group donation via SHMT-M. It remains unclear how dNTPs are sourced for mitochondrial DNA replication, or how methyl-groups are generated and maintained within the mitochondrion for ubiquinone biosynthesis. While metabolite profiling of knock-down parasites confirmed that SHMT-M has a direct role in mitochondrial folate metabolism, significant changes in the levels of many other metabolites from unrelated pathways were also detected. These results strongly suggest that Plasmodium SHMT-M has gained additional functions that are necessary for mETC activity and mitochondrial function.

We also performed a preliminary characterisation on two putative haloacid dehalogenases. Lipin was shown to dephosphorylate a subset of lipid species and to be necessary for normal parasite growth. The P. berghei ortholog is also predicted to be essential (Bushell et al., 2017) but no data are available at other lifecycle stages. It will be of interest to determine the precise lipid substrate specificity of this phosphatase and determine whether the enzyme is necessary for bulk lipid regulation or is necessary for specialise lipid signalling (Gulati et al., 2015; Bullen et al., 2016). In contrast, HAD4 appears to dephosphorylate a range of nucleotides, with a preference for nucleotide monophosphates. Despite the lack of a growth defect in asexual RBC stages and the dispensability of this protein in P. berghei liver stage development (Stanway et al., 2019), HAD4 may contribute to optimal metabolic flux and nucleotide homeostasis by repressing monophosphate accumulation. In particular, lack of HAD4 led to an accumulation of orotidine phosphate and a reciprocal decrease in the non-canonical orotidine (Fig 4E), suggesting HAD4 may regulate pyrimidine biosynthetic flux via dephosphorylation of a key intermediate. Lastly, AMR1 appears to be necessary for isoprenoid biosynthesis in agreement with previous work demonstrating its importance to apicoplast biogenesis (Tang et al., 2019). Defining the temporal profile of apicoplast impairment will aid in identifying the precise role of AMR1 and the function of other essential proteins involved in apicoplast biogenesis. Collectively, these data show that comprehensive metabolite profiling coupled with multiplex 13C-labeling can be used to detect subtle, as well as major perturbations in different metabolic mutants and is a powerful tool for functionally defining the large number of poorly annotated genes in these protists.

Materials and Methods

Cultivation and stable-isotope labelling

Plasmodium falciparum-infected RBCs were cultivated in RPMI 1640 GlutaMAX supplemented with 500 µM hypoxanthine, 22 µg/ml gentamicin, 0.2% (w/v) n-glucose, 25 mM HEPES and albumax II (0.5% w/v). Cultures were routinely synchronised with 5% sorbitol (w/v) to maintain a 12-hour developmental range. Transfectant parasite cultures were maintained with 5% human serum and cell density determined using a Neubauer Haemocytometer. Following a one-hour recovery in fresh RPMI 1640 at 37°C, 1 × 10^6 cells were centrifuged and the media replaced with 5 ml of RPMI containing unlabelled substrates or 11 mM 1^3C_6-glucose, 2 mM 1^3C_5 glutamine, 1.1 mM 1^3C_6 arginine, 1 mM 1^3C_4 serine, 1 mM 1^3C_5 proline, 5 mM 1^3C_2 acetate, 23 mM 1^3C_1 bicarbonate, 1 mM 1^3C_5 lysine, 10 mM 1^3C_3 glycerol or 1^3C-amino acid mix at 1 mg/ml (and a 1^2C-amino acid mix 1mg/ml used as a control). iRBC and matched uRBC samples were incubated for 5 h under standard culturing conditions.

Metabolite extraction

An aliquot of culture (1 × 10^6 cells/sample) was transferred to a microcentrifuge tube and centrifuged at 14,000 g for 30 s. The medium was aspirated, and the cell pellet was resuspended in ice-cold PBS (1 ml) to quench cell metabolism and transferred equally into two microcentrifuge tubes. Following centrifugation (14,000 g, 30 s), the PBS was aspirated and 200 µl of 80% acetonitrile (containing 1 µM 1^3C_5^15N_1 aspartate or 5 µM 1^3C_5 valine as the internal standard) was added and rapidly mixed for polar LC-MS analysis. Samples were centrifuged to remove precipitated protein (14,000 g, 5 min) and the supernatant transferred to a glass mass spectrometry vial (containing an insert) and stored at −80°C until LC-MS analysis. For GC-MS and LC-MS lipid analysis, the quenched cell pellet was suspended in 100 µl chloroform and vortex-mixed, prior to addition of 400 µl of methanol:H_2O (3:1 v/v) with further vigorous mixing. Samples were centrifuged for 5 min at 14,000 g and the supernatant transferred to a fresh microcentrifuge tube. H_2O (200 µl) was added to generate a biphasic mixture which was vortex-mixed and then centrifuged for 1 min at 14,000 g, and the top aqueous layer was collected for derivatisation and GC-MS analysis. The lower organic phase was collected for lipid analysis.

For GC-MS analysis, samples were dried by vacuum centrifugation, then resuspended in 90% methanol (100 µl) and transferred to a glass mass spectrometry insert. Samples were dried by vacuum centrifugation, washed with 100 µl methanol and dried. Samples were resuspended in 20 µl of methoxyamine (20 mg/ml) prepared in pyridine, sealed and incubated overnight with shaking at ambient temperature. The next day, 20 µl of BSTFA was added to each sample and incubated for 1 h prior to GC-MS analysis.

Lipid extraction was carried out on the remaining lower organic phase. The organic layer was transferred to a fresh microfuge tube and dried down under nitrogen flow. Dried samples were stored at −80°C until ready for lipid LC-MS analysis.

LC-MS acquisition

Polar metabolite detection was performed on an Agilent 6550 Q-TOF mass spectrometer operating in negative mode. Metabolites were separated on a SeQuant ZIC-pHILIC column (5 µM, 150 × 4.6 mm, Millipore) using a binary gradient with a 1200 series HPLC system across a 45-min method using 20 mM ammonium carbonate (pH 9) and acetonitrile as outlined in Cobbold et al., (2016). Two independent replicates of the metabolite profiling following AMR1 and Lipin depletion were performed using the same ZIC-pHILIC chromatography on a Thermo Q-Exactive operating in both positive and negative mode (rapid switching) as described previously (Creek et al., 2016). Lipid extracts were analysed on an Agilent 6550 Q-TOF using the reverse phase chromatography outlined by Bird et al (2011).
GC-MS analysis was performed using methods previously described (Saunders et al., 2011). Metabolites were separated using a BDS capillary column (J&W Scientific, 30 m × 250 μM × 0.25 μM) on a Hewlett Packard 6890 system (5973 EI-quadrupole MS detector). The oven temperature gradient was 70 °C (1 min); 70°C to 295°C at 12.5°C/min, 295°C to 320°C at 25°C/min; 320°C for 2 min. MS data were acquired using scan mode with a m/z range of 50–550, threshold 150 and scan rate of 2.91 scans/second. GC retention time and mass spectra were compared with authentic standards analysed in the same batch for metabolite identification.

Draft metabolome compilation

Raw Agilent.d files were converted to mzXML with MSconvert and analysed using the X13CMS R package (Huang et al., 2014). XCMS centWave peak detection was performed with a 10 ppm mass tolerance, following obiwave retention time correction. The getIsoLabelReport function was performed with P < 0.01 (Welch’s t-test), to identify features that significantly labelled under each 13C-labelling condition in both iRBC and uRBC cell types (isotope mass tolerance = 15 ppm). The getIsoDiffReport function then compared the significantly labelled features between iRBC and uRBC using P < 0.01 (Welch’s t-test). IsoDiffReport m/z features were removed if > 1,200 m/z, and the putative M0 species queried to the METLIN database with a 10 ppm mass tolerance, excluding all toxicants and adducts, except M-H, from the search. Only m/z features with a putative METLIN match were retained for further analysis. m/z features were further curated to remove mis-annotated in-source fragments and isotopologues, and where 13C-enrichment was < 1%. m/z features with a retention time of < 250 s were removed from the polar metabolite analysis (assuming these lipid-like species were also detected in the lipid C18 LC-MS analysis). This processing was performed for all 13C-labelling conditions across both C18 lipid and pHILIC polar LC-MS data and a single metabolite list compiled.

Metabolite identification was performed either with retention time matching to authentic standards (approximately 150 in-house metabolite standards or the HMDB compound library), or MS/MS matching. Polar MS/MS data were collected with an auto preferred MS/MS method at a collision energy of 10V, 20V and 40V at a threshold of 10,000 on an Agilent 6545 Q-TOF using the same chromatography outlined above. MS/MS spectra were searched against the METLIN PCDL database via the Agilent Qualitative software package and manually curated. MS/MS spectra without a match were reacquired in positive and negative mode and manually searched against the online METLIN database. Lipid MS/MS data were collected as above and converted to (ABFconverter) and analysed with the MS/MS software package (Tsugawa et al., 2015).

Polar GC-MS data were analysed using the DeXsi software package (Dagley & McConville, 2018) and metabolite identifications confirmed using an in-house metabolite standards library and merged with the LC-MS datasets. A first iteration of the observable metabolome of *P. falciparum* was compiled, and unique KEGG IDs (acquired from MetaboAnalyst 3.0 (Xia & Wishart, 2016)) were used to compare to the predicted metabolome of *P. falciparum* reported by Huthmacher et al. (2010). Under the conditions tested, we could not confidently assign stereoisomers or precise structures (e.g. D-glucose) and assigned KEGG IDs to be consistent with the level of evidence present in the literature and with the format used by Huthmacher et al. (2010). uRBC and iRBC datasets were then re-extracted for expected m/z of metabolites predicted from Huthmacher et al. (2010), to assess the rate of false negatives but also because certain classes of metabolites were not expected to label under any of the conditions tested (e.g. vitamins and some cofactors). Putative matches were then verified using authentic standards and MS/MS matching. This iteration of the observable metabolome of iRBC and uRBC cell types was then grouped according to the level of identification: (i) match to authentic standard and MS/MS match, (ii) match to authentic standard or MS/MS match, (iii) 13C-labelling data consistent with known metabolic pathway and (iv) exact mass match (≤ 10 ppm). To capture additional metabolites that were unlabelled across all 13C-substrates, all m/z features from unlabelled uRBC and iRBC extracts were compared with the expected exact mass of all metabolites in the predicted *P. falciparum* and RBC metabolomes. Putative hits were confirmed via pure standards or MS/MS matching where possible.

Pathway reconstruction

Metabolite enrichment analysis was performed with Metaboanalyst 3.0, and pathway reconstruction was performed for each stable-isotope condition. Observed metabolites that were labelled under a given condition were compared to the theoretical reactions for both iRBC and uRBC (Huthmacher et al., 2010). A matched reaction was defined as having a labelled metabolite corresponding to either an expected product or substrate. Unpredicted reactions were reported when an identified metabolite was labelled under a given 13C-substrate but did not match to a predicted reaction. A putative reaction was proposed based on the nearest proximity to an identified metabolite with consistency in the labelling pattern.

Generation of transgenic parasites for inducible protein disruption

For each gene of interest, a PAM site was selected at the centre or towards the start of the gene which ranked the highest via CHOP-CHOP v2 (Labun et al., 2016). Guide oligos were synthesised for cloning into the BtgZ1 site of pUF-Cas9g using the In-Fusion cloning kit (Takara). The gene was modified using a rescue template containing a 5′-homology arm immediately upstream of the PAM site, an artificial intron containing a loxP site, a recodonised version of the downstream gene sequence, a 3xHA tag, which was synthesised by GeneArt. The rescue template was cloned into the pGlmS plasmid using the BglII and SpeI (Prommman et al., 2013). The 3′-homology arm was generated via PCR of *P. falciparum* genomic DNA and cloned into the pGlmS plasmid with EcoRI and Ksal. Following verification of the correct DNA sequence, both pUF-Cas9g and pGlmS plasmids were purified with a midi-prep kit (Macherey-Nagel). pGlmS plasmid was linearised overnight using BglII, BglII and PvuI. 75–150 μg of pUF-Cas9g and linearised pGlmS were transformed into 3D7 ring-stage parasites with dimersisable Cre-recombinase integrated into its genome (Wilde et al., 2019). Cultures were maintained on WR99210 and Blasticidin and complete integration was confirmed on recovered parasites via PCR.

Inducible disruption of each targeted protein was performed cultivating transfected parasites under standard conditions in the presence or absence of 100 nM rapamycin and 2.5 mM
glucosamine. Initial tests were performed to determine length of treatment required to deplete the target protein and effectiveness of the DiCre-loxP and GlnS ribozyme system independently.

**Western blotting**

*Plasmodium falciparum* trophozoites were treated with saponin (0.05% w/v) and washed with ice-cold PBS, and centrifuged for 1 min (14,000 g) three times. Pellets were then extracted with RIPA buffer and protein concentration determined with a BCA assay. Following protein extraction with RIPA buffer (containing protease inhibitors), samples were processed and analysed as outlined previously (Yang et al, 2019).

**Recombinant protein expression**

PfHAD4 (PlasmoDB ID 3D7_1118400) was amplified from *P. falciparum* genomic DNA using the following primers:

5′-CTCACCCACCCACCCACCATATGAAAGATGAACAAATATCATGT TATTATC−3′

5′-ATCATCTAATTACTATGCAAGTATCATATCTAGATCTCG−3′

The PCR product was cloned by ligation-independent cloning into vector BG1861 (Alexandrov et al, 2004), which introduces an N-terminal 6xHis tag. The catalytic mutant HAD4D29A was generated by PCR amplification with the above primers in addition to the following primers:

5′-AATTGATAACGTCCCTTGCACCATACGATATG−3′

5′-ATCGTATGGTCAACGGCAACGTTATCAAATTGTA−3′

These primers were used to introduce an A→C point mutation in the coding sequence. The HAD4D29A construct was cloned by ligation-independent cloning into a modified BG1861 vector that also includes a KFS motif in front of the 6xHis tag to increase protein expression (Verma et al, 2019). Constructs were verified by Sanger sequencing.

Constructs were transformed into BL21 (DE3) pLysS *Escherichia coli* (Life Technologies). Cells were grown at 37°C and induced with 1mM isopropyl-β-D-thiogalactoside (IPTG). Cells were harvested by centrifugation and resuspended in lysis buffer containing 10mM Tris–HCl (pH 7.5), 20 mM imidazole, 1 mM MgCl2, 200 mM NaCl, 1 mg/ml lysozyme, 1mM dithiothreitol (DTT) and cComplete Mini EDTA-free Protease Inhibitor tablets (Roche), and sonicated. Soluble protein was bound to nickel agarose beads (Gold Biotechnology), eluted in 20 mM Tris–HCl pH 7.5, 150 mM NaCl and 300 mM imidazole, and further purified by size exclusion chromatography using a HiLoad 16/600 Superdex 200 pg column (GE Healthcare) equilibrated with 25 mM Tris–HCl (pH 7.5), 250 mM NaCl and 1 mM MgCl2 buffer. Fractions containing HAD4 were pooled, concentrated with a centrifugal filter, flash-frozen and stored at −80°C.

**HAD4 enzymatic assays**

All assays were performed in clear 96-well half-area plates using a FLUOstar Omega microplate reader (BMG Labtech) at 37°C. Reaction rates were determined using GraphPad Prism software. All reaction rates represent the mean and standard error of at least three experiments, each with technical replicates. Substrates were purchased from Sigma-Aldrich, except for GTP, dGTP, dATP, dCTP, dTTP, dUTP (Roche), dTMP (BioBasic Canada), DOXP (Echelon Biosciences) and fructose 1-phosphate (Santa Cruz Biotechnology). Other substrates tested that had activity < 0.1 µmol/min/mg were as follows: cytosine triphosphate; adenosine triphosphate; uridine diphosphate; deoxyuridine triphosphate; deoxyadenosine triphosphate; deoxythymidine triphosphate; inosine triphosphate; deoxy-adenosine triphosphate; D/L-glyceraldehyde-3-phosphate; sorbitol-6-phosphate; sodium pyrophosphate; deoxyxylulose-5-phosphate; galactose-1-phosphate, sucrose-6-phosphate; 2-deoxy-ribose-5-phosphate; glucose-6-phosphate; glyceral-2-phosphate; mannos-6-phosphate; thiamine monophosphate; fructose-6-phosphate; thiamine pyrophosphate; fructose-6-phosphate; thiamine pyrophosphate; fructose-6-phosphate; thiamine pyrophosphate; fructose-6-phosphate; thiamine pyrophosphate.
sedoheptulose-7-phosphate; racemic glycerol-1-phosphate; 2-phosphoglyceric acid; 2,3-diphosphoglycerate; myo-inositol-2-phosphate; mannose-1-phosphate; trehalose-6-phosphate; glucose-1-phosphate; phosphoenol pyruvate; and dihydroxyacetone phosphate.

Phosphatase activity and phosphate inhibition were measured using the substrate para-nitrophenyl phosphate (pNPP) (New England Biolabs). Reactions were performed in 50 µl volumes with 1 mM pNPP, 50 mM Tris–HCl (pH 7.5), 5 mM MgCl₂ and inorganic phosphate (0 mM–42 mM). Reactions contained 1 µg purified recombinant enzyme. Para-nitrophenyl production was quantified by absorbance at 405 nm.

Enzyme activity against phosphorylated substrates was measured using the EnzChek Phosphate Assay Kit (Invitrogen) according to supplier instructions. Each 50 µl assay contained 200 ng recombinant purified enzyme and 1 mM substrate. The reaction was quantified by absorbance at 360 nm, and reactions were linear with respect to time and enzyme concentration.

Enzyme activity against flavin mononucleotide (FMN) was measured using the BIOMOL Green kit (Enzo Life Sciences) to account for the overlap in absorbance between FMN and the EnzChek kit. Each 50 µl assay contained 200 ng recombinant purified enzyme, 25 mM Tris–HCl (pH 7.5), 250 mM NaCl, 1 mM MgCl₂ and 1 mM substrate. A time course was taken to obtain kinetic data, and the reaction was quantified by absorbance at 620 nm.

Substrate abbreviations are as follows: dGMP, deoxyguanosine monophosphate; dIMP, deoxyinosine monophosphate; dCMP, deoxycytosine monophosphate; GMP, guanosine monophosphate; IMP, inosine monophosphate; XMP, xanthosine monophosphate; AMP, adenosine monophosphate; 8oxodGMP, 8-oxo-deoxyguanosine monophosphate; dAMP, deoxyadenosine monophosphate; GDP, guanosine diphosphate; UMP, uridine monophosphate; CMP, cytosine monophosphate; dTMP, deoxothymidine monophosphate; dGDP, deoxyguanosine diphosphate; GTP, guanosine triphosphate; dADP, deoxyadenosine diphosphate; ADP, adenosine diphosphate; TTP, thymidine triphosphate; dGTP, deoxyguanosine triphosphate; PLP, pyridoxal-5-phosphate; RSP, ribose-5-phosphate; FBP, fructose-1,6-bisphosphate; E4P, erythrose-4-phosphate; and 3PGA, 3-phosphoglyceric acid.

Data availability

Mass spectrometry data are available at MetaboLights (https://www.ebi.ac.uk/metabolights/) MTBLS1815 (http://www.ebi.ac.uk/metabolights/MTBLS1815).

Expanded View for this article is available online.

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Author contributions

Project design: SAC, SAR, MJM. Mass spectrometry: SAC, DJC. Gene knock-out and protein localisation studies: SAC, MVT. Mutant phenotype analysis: SAC, PF, AOJ, EM, MK. Project supervision and management: SAC, MJM, LT, SAR. Data analysis and manuscript preparation: SAC, MJM. Funding Acquisition: MJM, LT, SAR, AOJ.

Conflict of interest

The authors declare that they have no conflict of interest.

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