Phylogenetic profiles of all membrane transport proteins of the malaria parasite highlight new drug targets

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ABSTRACT In order to combat the on-going malaria epidemic, discovery of new drug targets remains vital. Proteins that are essential to survival and specific to malaria parasites are key candidates. To survive within host cells, the parasites need to acquire nutrients and dispose of waste products across multiple membranes. Additionally, like all eukaryotes, they must redistribute ions and organic molecules between their various internal membrane bound compartments. Membrane transport proteins mediate all of these processes and are considered important mediators of drug resistance as well as drug targets in their own right. Recently, using advanced experimental genetic approaches and streamlined life cycle profiling, we generated a large collection of Plasmodium berghei gene deletion mutants and assigned essential gene functions, highlighting potential targets for prophylactic, therapeutic, and transmission-blocking anti-malarial drugs. Here, we present a comprehensive orthology assignment of all Plasmodium falciparum putative membrane transport proteins and provide a detailed overview of the associated essential gene functions obtained through experimental genetics studies in human and murine model parasites. Furthermore, we discuss the phylogeny of selected potential drug targets identified in our functional screen. We extensively discuss the results in the context of the functional assignments obtained using gene targeting available to date.

INTRODUCTION The malaria parasite has adopted a highly complex life cycle involving a continuous switching between vertebrate hosts and anopheline mosquitoes. Within humans, Plasmodium species are obligate intracellular parasites moving through three different life-cycle stages. After an infectious mosquito bite, a single phase of preclinical growth within the host liver cells commences [1]. Next, fast proliferating blood-stage parasites are the cause of malaria-associated pathology and severe disease outcome [2]. Finally, some of the asexual blood-stage parasites are triggered to develop into male or female gametocytes, which are required for sexual reproduction following a mosquito blood meal [3,4].

Despite a gradual decline in annual malaria cases and deaths, the parasite remains one of the largest global killers. In 2015, WHO reported 150-300 million cases and 438,000 deaths [5]. Difficulty in combating the disease is exacerbated by the growing resistance to anti-malarial drugs and the absence of an effective vaccine. The continuous need for new therapeutic anti-malarial drugs is unquestioned. Yet, due to limited availability, there is a much more pressing need for drugs that can prevent infections by acting prophylactically on the liver stage of the parasite, and transmission-blocking compounds, which kill the gametocytes thus helping to prevent the spread of the disease. Chemoprophylaxis is not only important for travellers [6] but also particularly for women in endemic areas in their first or second pregnancy [7]. Thus, there is a critical need for novel drugs that may be used prophylactically, therapeutically, or to block transmission [8].

As defined by the Medicines for Malaria Venture in there 2015 annual report [9], ideal medicines for treatment
and protection would both be suitable for mass drug administration programs and require a single encounter treatment, or better still a single exposure treatment, to help improve compliance. Treatment of infection should be effective against all life-cycle stages of all five malaria species infecting humans, including resistant strains, and resistance against the new drug should be difficult to achieve.

Generally, it is believed that these features may be best obtained by the combination of at least two active compounds, one fast acting for immediate clearance of the infection and a second, slower-acting compound providing long duration of efficacy. Furthermore, an ideal treatment would be gametocytocidal to prevent the spread of infection to mosquitoes, while sporontocidal or liver-stage activity would provide a prophylactic component. To complete the wish list of the ideal anti-malarial drug, it should also be active against the so-called hypnozoites, the dormant stages of certain malaria parasite species (most notably *P. vivax*) that are the cause of relapse infection even many years after the infectious mosquito bite. It is obvious that the development of the ideal anti-malarial drug will not be straightforward and will result in a compromise between the long list of desirable attributes and features. In this light, it is particularly important that ongoing functional studies of malaria parasite biology continue to highlight potential new drug targets that have important roles in the different life-cycle stages of the parasite and are conserved among all malaria parasite species but are absent from or have diverged significantly in humans, such that compounds acting on these important *Plasmodium* proteins may do so effectively as well as selectively.

*Plasmodium* membrane transport proteins (MTP), such as the chloroquine resistance transporter (CRT) and the ATP-binding cassette (ABC) transporter family, including the multidrug resistance proteins (MDR) and the multidrug resistance-associated proteins (MRP), are well known for their roles in anti-malarial drug resistance [10,11]. MTPs are also generally considered potential drug targets in their own right [12,13]. Spiroindolones and dihydroisouquinolones are new classes of potent anti-malarial drugs, currently under clinical testing, that have both been shown to act via the *Plasmodium falciparum* cation ATPase, ATP4, causing severe disturbance of Na⁺ homeostasis in the parasite [14-17].

To survive and thrive, malaria parasites utilize a range of transport processes to import nutrients, export waste, and redistribute ions and small organic molecules between different sites and organelles [12,18]. MTPs of different classes facilitate these processes. Following the functional and phylogenetic classification of MTPs from the Transporter Classification Database [http://www.tcdb.org] [19], *Plasmodium* MTPs can be classified as: α-type channels and β-barrel porins (TCDB Class 1.A/B); P-P-bond hydrolis-driven transporters, here referred to as pumps (TCDB Class 3.A); porters, including uniporters, symporters, and antiporters (TCDB Class 2.A); and unclassified, putative MTPs.

Experimental genetics is a powerful means to further explore critical functions of *Plasmodium* MTPs for parasite survival throughout its complex life cycle [20,21]. Due to more efficient experimental and computational methods and access to the entire *in vivo* life cycle, the majority of such studies have been performed using murine malaria model species, notably *Plasmodium berghei*, and have focused on single MTPs (Table S1). Two of our most recent studies have more than doubled the number of targeted genes, and have generated loss-of-function mutants in both *P. falciparum* and *P. berghei* [22,23]. A systematic study of the MDR family demonstrated that four of seven members fulfill essential functions during blood-stage development, highlighting these as potential drug targets [22]. While targeting 35 orphan MTPs, Kenthirapalan et al. produced the largest collection of *P. berghei* knock-out parasites available to date [23]. The 29 available mutant lines provide a powerful resource for further studies of malaria parasite transport processes. In addition to highlighting six genes essential for blood-stage survival including five pumps, they also revealed potential prophylactic (MFS6) and transmission-blocking (ZIP1) drug targets [23].

Here, we performed comprehensive orthology profiling by reciprocal Blast of all identified 139 *Plasmodium* MTPs (consisting of the list published by Martin et al. [24] expanded with newly identified candidates) against a selection of 41 species from the entire breadth of the eukaryotic kingdom. To further validate the potential of the eight newly identified drug targets, we have explored their phylogenetic relationships in detail. These results are discussed extensively in the context of available insights from functional genetics studies of malaria parasite MTPs.

**RESULTS AND DISCUSSION**

**Comprehensive orthology assignments of *Plasmodium* MTPs**

To identify the levels of conservation of the 139 *P. falciparum* MTPs (Table S1) within the eukaryotic domain, we first performed an extensive orthology assignment using Blast against a set of 41 species (Table S2). We selected a subset of species to represent the enormous diversity in the eukaryotic domain by including sequences from all kingdoms, with an emphasis on protozoan species, commonly used model species, and parasite species of great medical or veterinary importance. These include the clinically second most important human malaria parasite *Plasmodium vivax* and the murine malaria model parasite *P. berghei. Plasmodium* species belong to a large monophyletic group of largely obligate intracellular parasites, the *Apicomplexa*. We included six additional apicomplexan parasites in our analysis: two piroplasms (*Babesia bovis* and *Theileria annulata*) that cause cattle fever and like *Plasmodium* species belong to the *Aconoidasida*, and four *Conoidasida*, including parasites of medical (*Toxoplasma gondii* and *Cryptosporidium parvum*) and veterinary (*Neospora caninum* and *Eimeria tenella*) importance.

Apicomplexan parasites are characterized by a complex at the apical end that is used to penetrate and enter a wide variety of host cells. However, this is not the only peculiar subcellular structure of these eukaryotic single cell para-
sites. Being eukaryotes, one would naturally expect most of the commonly shared organelles to be present, but while most do retain a single and thoroughly reduced mitochondrion, some have only vague remnants of the cell’s power house in the form of mitosomes [25]. An additional organelle of endosymbiotic heritage present in the majority of apicomplexan parasites (though it has been lost in Cryptosporidium species) is the former-photosynthetic apicoplast [26]. This plastid is of red algal origin, a trait shared across a variety of highly divergent species, including other alveolates, such as Chromerida and dinoflagellates, but not ciliates. For our analysis, we included four alveolates, including two ciliates and one of the closest relatives of the Apicomplexa, Vitrella brassicaformis.

The true evolutionary relationships between highly diverse protozoa lineages are unresolved and still a matter of dispute but many interesting parallels can be observed. Thus Cryptomonads, Haptophyta, and a significant proportion of Stramenopiles, all harbour plastids of red algal origin, although at present it is not clear whether these plastids originate from single or multiple secondary endosymbiotic events [27-29]. In our analysis, we included one cryptomonad, Guillardia theta, as well as four stramenopiles including the oomycyte Phytophthora infestans. The latter is a plant parasite causing potato blight, which does not harbour a plastid, but it does share another interesting feature with malaria parasites. Both oomycetes and malaria parasites extensively remodel the host they infect by exporting a large repertoire of effector proteins using comparable strategies [30]. Further protist representative sequences were taken from two Rhizaria, six Excavata, including three kinetoplastid parasites, and two Amoebozoa.

From the plant kingdom, we took a green and a red alga and the most commonly used model plant Arabidopsis thaliana. Ophistokonts were represented by two fungi (baker’s yeast and the pathogenic Cryptococcus neoformans) and by nine animals, including the parasitic flatworm Schistosoma haematobium, the malaria mosquito Anopheles gambiae, a number of widely used model organisms, and of course mouse and human.

Potential prophylactic drug targets

In our recent functional screen of orphan MTPs, we identified a critical role for a major facilitator superfamilly member, MFS6, during liver-stage development, in addition to important functions in the blood [23]. Though we identified a putative orthologue in V. brassicaformis (Figure 1), MFS6 is largely Plasmodium-specific and appears absent from humans. Its important functions during both liver- and blood-stage development justify further exploration of the protein as a target for compounds with combined prophylactic and therapeutic activity. P. berghei MRP2-deficient parasites demonstrate a complete arrest in the liver, a phenotype that was also observed in P. falciparum counterpart MRPs [31]. Like MFS6, MRP2 appears to be largely Plasmodium-specific (Figure 1) providing another promising drug target. However, for these two MTPs both their function and their respective substrates remain unresolved.

Evidence is mounting that transport processes of metal ions, in particular of heavy metals such as iron, copper, and zinc, could provide efficient targets for chemoprophylactic treatments. Although the exact mechanisms are unclear, P. berghei liver-stage development is influenced by host iron homeostasis [32,33]. Recently, a vascular iron transporter was described that plays an important (although not critical) role during liver infection in vivo and in vitro [34]. Parasites lacking an alternative zinc–iron permease (ZIPCO) are severely affected in liver-stage development in vitro and show a delay in prepatency of two days [35]. A similar delay in prepatency was observed following needle injection of ctr2+ sporozoites despite developing normally in culture [23]. Interestingly, these parasites were most severely affected during natural transmission by infectious mosquito bites or following subcutaneous injection of the sporozoites. Parasites deficient in the copper channel 1 gene (CTR1) completely failed to transmit, but this was at least in part attributable to a much reduced and delayed sporozoite production, and the infectivity of these sporozoites remains to be determined in more detail to establish whether CTR1 like CTR2 plays an important role in the liver [23]. Despite the fact that the putative heavy metal transporting MTPs with a demonstrated important role during the establishment of new infection are not strictly essential, they may still prove interesting targets for prophylactic interventions, since none of the MTPs discussed above appear to have a clear reciprocal orthologue in humans and show only partial sequence matches (Figure 1).

Potential transmission-blocking drug targets

The importance of heavy metal homeostasis appears not to be restricted to mosquito-to-mouse/human transition. Also sexual blood-stage parasites, parasite fertility, and effective colonization of the mosquito midgut appear to be strongly dependent on the correct distribution of these cations.

Where the two copper channels may act in copper transport in the liver stages, a copper-transporting P-type ATPase (CuTP) was shown to be central to fertility of both male and female gametes [36]. Activation of cutp5 male microgametes, a process known as exflagellation, is reduced to ~10% of wild-type and this reduction could be phenocopied using an intracellular copper chelator. Cross-fertilization studies and an even more pronounced reduction in oocyst numbers indicate additional important roles in the female gametes. ZIP1, a paralogue of the liver-stage-specific ZIPCO, is crucial for mosquito colonization [23]. Exflagellation in zip1– parasites is reduced to naught, which is directly attributable to a nearly complete absence of male gametocyte formation. Since ZIP1-deficient parasites also have a slightly reduced blood-stage multiplication rate, ZIP1 is potentially a very attractive transmission-blocking drug target. Indeed, if a compound would be able to target ZIPCO and ZIP1, this would be a triple-acting drug that could be used prophylactically, therapeutically, and to block spread of the disease. One disadvantage could be
FIGURE 1: Orthologies of membrane transport proteins of the malaria parasite. Extensive amino acid based reciprocal homology searches were performed to establish orthologues of the all Plasmodium channels/pores (TCDB Class 1.A/B [19]; pink), pumps (3.A; purple), porters (2.A; cyan), and unclassified, putative MTPs (yellow) in representative species of the entire breadth of the eukaryotic domain. Dot sizes indicate the fraction of the sequence length over which homology was detected; red dots indicate reciprocal orthologues. When a protein has not been given a name, these are indicated by their P. falciparum geneID number. MTPs for which the encoding genes were deleted successfully are indicated in blue, unsuccessful gene deletions suggesting essential functions during blood-stage development are highlighted in red. ®, species harbouring plastids of red algal origin; *, putative heavy metal transporting MTPs) Please see the continuation of the table on the next page.
that humans harbour fourteen zinc-iron permeases [37], although only one of these bears any sequence similarity to the *Plasmodium* copies. Nevertheless, caution is needed to ensure that the compound is sufficiently parasite-specific. Phylogenetic profiling of the identified ZIP sequences supports a relatively recent gene duplication at the root of the Aconoidasida (Figure 2), and the relatively long distance from the vertebrate ZIPs suggest that specific targeting of the *Plasmodium* ZIPs might be feasible.

Other MTPs with roles during mouse/human-to-mosquito host transition have been identified, such as the Ca$^{2+}$/H$^+$ exchanger (CAX) [38], a cation diffusion factor or putative zinc transporter (CDF) [23], and the pantothenate transporter [23,39]. Unfortunately, not a single MTP has been identified that is critical for both male and female gametocyte formation, although it remains to be tested if zip1− female gametocytes remain fertile or have lost the capacity to reproduce.

**Potential therapeutic drug targets**

Two types of channels have been discussed as potential therapeutic drug targets. Two potassium channels (K1 and K2) were refractory to gene deletion in *P. falciparum* [40]. However, absence of supporting evidence that this was not merely due to the technical difficulties of targeting *P. falciparum* genes and the fact that PbK1 could be readily deleted [41] suggest that these channels may not be strictly necessary for blood-stage survival. The suitability of aquaglyceroporin (AQP) has also been subject of controversy. Initial studies demonstrated that blood-stage growth of *aqp* parasites was strongly affected [42]. Independently, we were only able to replicate a minor defect in Swiss-Webster mice [43]. Using a sensitive flow cytometry-based method in NMRI mice, we saw no difference in growth rates of WT and *aqp* parasites either growing in direct competition or in individual mice [44]. The relevant difference between NMRI and Swiss-Webster mice that may lead to this observation is unclear.

Interestingly, a vast majority of the currently identified resistance markers, e.g. the ABC family [11], as well as the single validated druggable MTP, ATP4 [16,17], are primary active transporters that require ATP to fuel their activity. As the parasite invests energy in their functioning, it is perhaps not surprising that many pumps play crucial roles at some stages during the parasite’s life cycle. Indeed, of all nineteen targeted pumps, eleven were shown to be refractory to gene deletion (Table S1). In addition to the four MDRs (MDR1, MDR4, MDR6, and MDR7) [22], these in-
clude the putative cation transporting ATPase, ATP4, (S. Kenthirapalan, K. Matuschewski, T.W.A. Kooij, unpublished data), ATP6 [45] and the V-type proton ATPase subunit G [46], ABC3, and four predicted aminophospholipid transporters [23].

Our orthology profiling indicates that the ABC transporters, with the exception of MDR1 (and to lesser extent MDR2), are poorly conserved across the eukaryotic kingdom, further highlighting their potential as anti-malarial drug targets (Figure 1). ABC3 appears to be a unique, *Plasmodium*-specific ABC family member and is characterized by the presence of two transmembrane domains consisting of multiple transmembrane helices interspersed by a single nucleotide-binding domain. An extensive phylogenetic profiling of all ABC transporters identified in 16 eukaryotic and 55 prokaryotic genomes assigned *Pf*ABC3 to a poorly supported clade together with sequences from two archaea from different kingdoms and sequences from five bacteria from four different phyla [47]. Thus, the orthology and phylogeny of ABC3, along with demonstrated essential function during blood-stage development, strengthen its potential as a therapeutic drug target.

Another class of primary active transporters that was highlighted for its druggable potential consists of putative aminophospholipid-transporting P₄-type ATPases, from hereon referred to as flippases [23]. In addition to ATP2, ATP7, and ATP8, these also include two putatively bifunctional proteins that also harbour guanylyl cyclase (GC) activity [48]. GCα, like the other flippases, was essential for blood-stage development, whereas GCβ plays a critical role in colonization of the mosquito midgut [23,49,50]. While parts of the GCs are conserved, no bifunctional

![Phylogenetic tree of putative flippases. Four of five Plasmodium berghei putative aminophospholipid-transporting P₄-type ATPases were shown to be essential for blood-stage development in vivo. This is a maximum likelihood tree based on the complete sequences of the identified orthologues along with models of all the domain architectures. Nodes with support values <50 are left unresolved. This unrooted tree shows low resolution at the base, making it difficult to interpret their phylogenetic history despite the fact that the proteins share their structures. ATP8 forms a well-supported clade with orthologues from a variety of species, while the apicomplexan branches with poor orthology assignments (GCα, GCβ, and ATP7) resolve well supporting the candidacy of GCα and ATP7 as therapeutic drug targets.](image-url)
orthologues were found outside the apicomplexan clade (Figure 1). The phylogenetic tree of the flippases confirmed this notion (Figure 3). Orthology and phylogenetic profiles further indicate that ATP2 and ATP8 are rather well conserved including human orthologues, whereas ATP7 is largely apicomplexan-specific (Figure 1). Combined, these data suggest that ATP7 and GCa may form attractive targets for novel anti-malarial compounds. Of note, for one P. falciparum gene encoding a putative flippase (PF3D7_1468600), no orthologue exists in P. berghei and a possible essential role has not yet been established.

Of the largest group of MTPs, the porters (TCDB Class 2.A) [19], 28 genes have been targeted in P. berghei, only three times without success (Table S1). In addition to the well-studied CRT [51], these include a hexose transporter HT [52], which is well conserved including in humans (Figure 1), and the putative drug metabolite transporter DMT2 [23]. The latter presents a particularly interesting case considering the orthology discovery. Orthologues were identified in all apicomplexan parasites, with the exception of Cryptosporidium, in other alveolates, but not ciliates, and in the cryptomonad, G. theta, and most stramenopiles. Most species have a single DMT2 but T. gondii, N. caninum, and V. brassicafcomis have multiple copies. While the exact phylogenetic relationships of the alveolates, cryptomonads, and stramenopiles is still a topic of debate, many of these chromalveolates, as they are commonly referred to, harbour a plastid of red algal origin. Ciliates and Cryptosporidium species do not have such a plastid and thus it is tempting to speculate that DMT2 localizes to this endosymbiotic organelle. Indeed, the red alga Cyanidioschizon merolae appears to harbour a sequence with a very weak similarity that was initially just below the cut-off E-value applied to establish significant homologies. However, the presence of a rather well conserved copy in P. infestans, that lacks a relic plastid, and presence of weak similarities in three other unrelated eukaryotes (Giardia lamblia, Dictyostelium discoideum, and Chlamydomonas reinhardtii) suggest that the evolutionary history of this gene may be more complicated. When building a maximum likelihood tree for all identified homologues, including the few hits from unrelated species, the tree does not resolve well (data not shown). However, when only including sequences from the chromalveolates and the red alga, the tree is well-supported showing phylum-specific clades with Cyanidioschizon merolae as the outmost group of the tree (data not shown). Attempts to include other sequences as outgroup, e.g. Plasmodium DMT1 sequences, were unsuccessful due to a lack in sequence similarity and the consequently poorly resolving sequence alignments. The most consistent results were obtained when using a single DMT2 homologue of one of the three unrelated species as an outgroup (Figure 4).

Despite the significant uncertainties and remaining unresolved questions about the origin of DMT2, these data could well suggest that this essential MTP is localized in the parasite’s apicoplast. Our initial localization studies were hampered by very low expression levels of the protein and hence difficult to interpret, but appeared to indicate an intraparasitic staining including a specific, small structure that may well be the apicoplast [23]. In blood-stage parasites, the single critical role of the apicoplast was shown to be the production of isopentenyl pyrophosphate (IPP) [53] and it is tempting to speculate that DMT2 is the dedicated IPP transport protein. However, DMT2 may also be involved in critical processes for the maintenance of the organelle and the IPP biosynthesis pathway, e.g. through the import of sulphur or iron into the organelle for essential iron-sulphur cluster biosynthesis [54].

FIGURE 4: Phylogenetic tree of drug metabolite transporter 2 (DMT2). Plasmodium berghei DMT2 was shown to be essential for blood-stage development in vivo. This is a maximum likelihood tree of all homologues found in chromalveolate species and the red alga Cyanidioschizon merolae using the Giardia lamblia homologue as an outgroup. Nodes with support values <50 are left unresolved. Apicomplexan DMT2 sequences, including the different paralogous sequences, form a distinct and well-supported clade, while the red algal distant homologue and the majority of other chromalveolates sequences sit unresolved at the base of this clade.
Conclusions
In conclusion, our extensive orthology assignment and phylogenetic profiling, in combination with published experimental genetics studies, particularly in the murine malaria model parasite *P. berghei*, support the candidacy of a number of prophylactic, therapeutic, and transmission-blocking drug targets. Further studies into the biochemical and structural properties of these MTPs are required and deserve prioritization.

MATERIALS AND METHODS
Identification of putative orthologues
We first selected candidate protein sequences by homology search using Blast (blastp, version 2.2.29+) [55], against a set of pre-selected proteomes (Table S2) applying a threshold for being considered significant of E=5e-5. Next, for each candidate, we performed a reciprocal Blast search against the *Plasmodium falciparum* genome. From March 19th, 2016, hits on the same sequence were analysed and the overall coverage of the identified homology was calculated as the fraction of the query length. Candidates which returned the same protein as the original query (i.e., reciprocal blast hits) were kept as putative orthologues, while remaining hits were retained as unspecified homologues. We further inspected the putative homologues using the RADS algorithm [56] and the Needleman-Wunsch global alignment algorithm.

Phylogenetic reconstruction
Groups of putative orthologues for selected proteins were aligned using the programme Clustal Omega v. 1.2.1 [57] with default parameters. The alignments were then manually inspected and regions of low coverage and poor conservation were masked. Maximum likelihood phylogeny reconstruction was performed with the TreePuzzle [58] with 10,000 iterations and a mixed model of rate homogeneity.

Domain analysis
Domains of protein sequences were detected with hmmscan from the HMMer package v. 3.1b1 (http://hmmer.org) and the HMM profile collection from the PfamA database (October 2015). Significant overlaps were solved by E-value precedence. Domain architectures were analysed in combination with phylogenetic trees using the DoMosaic programme [59].

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SUPPLEMENTAL MATERIAL
All supplemental data for this article are available online at www.microbialcell.com.

CONFLICT OF INTEREST
The authors declare no existing conflicts of interest.

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