PMRT1, a Plasmodium specific parasite plasma membrane transporter is essential for asexual

and sexual blood stage development

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- 4 Jan Stephan Wichers^{1,2,3}, Paolo Mesén-Ramírez², Jing Yu-Strzelczyk⁴, Gwendolin Fuchs^{1,2,3},
- 5 Jan Stäcker², Heidrun von Thien^{1,2}, Arne Alder^{1,2,3}, Isabelle Henshall⁵, Benjamin Liffner⁵,
- 6 Georg Nagel⁴, Christian Löw^{1,6}, Danny Wilson^{5,7}, Tobias Spielmann², Shiqiang Gao^{4#}, Tim-
- 7 Wolf Gilberger^{1,2,3,#}, Anna Bachmann^{1,2,3,&}, Jan Strauss^{1,2,3,6,#,&,†}
- ⁹ Centre for Structural Systems Biology, 22607 Hamburg, Germany.
- 10 ²Bernhard Nocht Institute for Tropical Medicine, 20359 Hamburg, Germany.
- ³Biology Department, University of Hamburg, 20146, Hamburg, Germany
- 12 ⁴Institute of Physiology, Department of Neurophysiology, Biocenter, University of Wuerzburg,
- 13 97070 Würzburg, Germany
- 14 ⁵Research Centre for Infectious Diseases, School of Biological Sciences, University of
- 15 Adelaide, Adelaide 5005, Australia.
- 16 ⁶European Molecular Biology Laboratory, Hamburg Unit, Hamburg, Germany
- ⁷Burnet Institute, 85 Commercial Road, Melbourne 3004, Victoria, Australia.
- 19 *Corresponding authors: gao.shiqiang@uni-wuerzburg.de, gilberger@bnitm.de,
- 20 jan.strauss@geomar.de
- 21 &Contributed equally
- [†]Present address: GEOMAR Helmholtz Centre for Ocean Research Kiel, 24105 Kiel, Germany
- 24 ORCID:

25	Jan Stephan Wichers:	0000-0002-0599-1742
26	Paolo Mesén-Ramírez:	0000-0001-7842-5867
27	Jing Yu-Strzelczyk:	0000-0002-7576-6831
28	Gwendolin Fuchs:	0000-0001-9294-6984
29	Jan Stäcker:	0000-0002-4738-6639
30	Arne Alder:	0000-0003-4918-4640
31	Isabelle Henshall:	0000-0002-5906-0687
32	Benjamin Liffner:	0000-0002-1573-6139
33	Danny Wilson:	0000-0002-5073-1405
34	Georg Nagel:	0000-0001-8174-8712
35	Christian Löw:	0000-0003-0764-7483
36	Tobias Spielmann:	0000-0002-3968-4601
37	Shiqiang Gao:	0000-0001-6190-9443
38	Tim-Wolf Gilberger:	0000-0002-7965-8272
39	Anna Bachmann:	0000-0001-8397-7308
40	Jan Strauss:	0000-0002-6208-791X

Abstract

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Membrane transport proteins perform crucial roles in cell physiology. The obligate intracellular parasite Plasmodium falciparum, an agent of human malaria, relies on membrane transport proteins for the uptake of nutrients from the host, disposal of metabolic waste, exchange of metabolites between organelles and generation and maintenance of transmembrane electrochemical gradients for its growth and replication within human erythrocytes. Despite their importance for *Plasmodium* cellular physiology, the functional roles of a number of membrane transport proteins remain unclear, which is particularly true for orphan membrane transporters that have no or limited sequence homology to transporter proteins in other evolutionary lineages. Therefore, in the current study, we applied endogenous tagging, targeted gene disruption, conditional knockdown and knockout approaches to investigate the subcellular localization and essentiality of six membrane transporters during intraerythrocytic development of *P. falciparum* parasites. They are localized at different subcellular structures – the food vacuole, the apicoplast, and the parasite plasma membrane – and showed essentiality of four out of the six membrane transporters during asexual development. Additionally, the plasma membrane resident transporter 1 (PMRT1, PF3D7 1135300), a unique Plasmodiumspecific plasma membrane transporter, was shown to be essential for gametocytogenesis. Heterologous expression of wild-type and mutation constructs in Xenopus laevis oocytes indicated ion transport upon membrane hyperpolarization and a functional role of negatively charged amino acids protruding into the parasitophorous vacuole lumen. Overall, we reveal the importance of four orphan transporters to blood stage P. falciparum development and provide the first functional characterization of PfPMRT1, an essential parasite membrane transporter.

Importance (150 words)

Plasmodium falciparum-infected erythrocytes possess multiple compartments with designated membranes. Transporter proteins embedded in these membranes do not only facilitate movement of nutrients, metabolites and other molecules between these compartments, but are common therapeutic targets and can also confer antimalarial drug resistance. Orphan membrane transporter in *P. falciparum* without sequence homology to transporters in other evolutionary lineages and distant to host transporters may constitute attractive targets for novel intervention approaches. Here, we localized six of these putative transporters at different subcellular compartments and probed into their importance during asexual parasite growth using reverse genetic approaches. In total, only two candidates turned out to be dispensable for the parasite, highlighting four candidates as putative targets for therapeutic interventions. This study reveals the importance of several orphan transporters to blood stage *P. falciparum* development and provides the first functional characterization of *Pf*PMRT1, an essential parasite membrane transporter.

Introduction

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Plasmodium spp. malaria parasites inhabit diverse intracellular niches and need to import nutrients and export waste across both, host-cell and parasite membranes. Despite this, there are less than 150 putative membrane transporters encoded in the genome of *Plasmodium* falciparum, the most virulent malaria parasite, making up only 2.5% of all encoded genes (P. falciparum 3D7 v3.2: 5280 genes) (1-5), which is reduced compared to other unicellular organisms of similar genome size. The loss of redundant transporters is a typical feature of many intracellular parasites (6) and, as a result, the proportion of transporters that are indispensable for parasite survival increases (2), some of which have been shown to be critical for the uptake of several anti-Plasmodial compounds and/or to be involved in drug resistance (7-20). Moreover, the parasite's intracellular lifestyle resulted in the evolution of additional specialized transporters without human homologues (1). During its intraerythrocytic development, the parasite relies on the uptake of nutrients, such as amino acids, pantothenate or fatty acids, from its host erythrocyte as well as from the extracellular blood plasma (21–24). As P. falciparum resides in a parasitophorous vacuole (PV) in the host erythrocyte, nutrients acquired from the extracellular milieu must traverse multiple membranes: the erythrocyte plasma membrane (EPM), the parasitophorous vacuole membrane (PVM), the parasite plasma membrane (PPM) and eventually membranes of intracellular organelles, such as those of the apicoplast or mitochondria (21, 25-27). The unique requirements of malaria parasite survival have led to the evolution of a number of orphan transporters, whose localization or function cannot be predicted based on sequence homology to transporters in other organisms (4, 28). Despite the likely importance of uniquely adapted transporters to *P. falciparum* survival, subcellular localization, essentiality, function and substrate specificity for most P. falciparum transporters has not been directly determined (2, 21, 26). The best functional evidence available for many Plasmodium-specific transporters comes from a recent knockout screen of these orphan transporters in the rodent malaria parasite *Plasmodium berghei* (28). However, whether observations for different transporters in the P. berghei model are directly transferrable to P. falciparum have yet to be examined. Therefore, in this study, we explored the localization and essentiality of four predicted orphan transporters that had been partially characterised in P. berghei and included two additional transporters with no experimental characterization available.

Results

A subset of orphan transporters characterized in the *P. berghei* malaria model was selected for further characterization in *P. falciparum*. The four transporters selected were reported to be important at different stages of rodent malaria parasite growth with i) *P. berghei* drug/metabolite transporter 2 (*Pf*DMT2: PF3D7_0716900) found to be essential for asexual blood stage development, ii) *P. berghei* zinc transporter 1 (*Pf*ZIP1: PF3D7_0609100) was

117 essential across transmission stages but not blood stages, where there was only a slight 118 growth defect, iii) P. berghei cation diffusion facilitator family protein (PfCDF: PF3D7_0715900) 119 knockout parasites had a defect during transmission stages but not during asexual stages, and 120 berghei major facilitator superfamily domain-containing protein (PfMFS6: 121 PF3D7 1440800) was found to be essential for parasite transmission from mosquitos to a new 122 host, with a growth defect observed at asexual and gametocyte stages but not during mosquito 123 stage parasite growth (28, 29). In order to confirm expression of these transporters in P. 124 falciparum asexual stages, we screened the list of "Genes coding for transport proteins" 125 included in the Malaria Parasite Metabolic Pathways (MPMP) database (1, 30) for proteins 126 with i) RNA-seg (31, 32) and ii) proteomics evidence (33, 34) for expression in asexual blood 127 stages. We identified two additional putative transporters (PF3D7_0523800, PF3D7_1135300) 128 which had yet to be functionally characterized in the P. berghei screen by Kenthirapalan et al. 129 (28) or any other experimental model. Both were subsequently included in our characterization of P. falciparum orphan transporters, and named as 'food vacuole resident transporter 1' 130 131 (FVRT1: PF3D7 0523800) and as 'plasma membrane resident transporter 1' (PMRT1: 132 PF3D7_1135300) based on their subcellular localization. AlphaFold-based structure 133 predictions (35) and results from structure homology search (36) of all six selected transporters 134 are provided in Figure S1.

Localization of putative *P. falciparum* transporters

- 136 To determine subcellular localization, we tagged the six putative transporters endogenously
- with GFP using the selection-linked integration (SLI) system (37) (Figure 1A). Additionally, a
- 138 glmS ribozyme sequence was included in the 3'UTR, which enabled conditional gene
- knockdown upon addition of glucosamine (38). Correct integration of the plasmid into the
- 140 respective genomic locus was verified by PCR and expression of the GFP-fusion protein was
- confirmed by Western blot for each generated cell line (Figure S2A, B).
- 142 All transgenic cell lines expressed the GFP-fusion protein, demonstrating that these
- transporters are expressed in asexual blood stage parasites (Figure 1B-G, S2A). Expression
- 144 levels were sufficient to allow determination of subcellular localization (Figure 1B-G): (i)
- 145 PF3D7 0523800-GFP localized to the food vacuole, (ii) PfDMT2-GFP and PfMFS6-GFP
- 146 apicoplast localization, and (iii) PfZIP1-GFP and PF3D7_1135300-GFP parasite plasma
- membrane (PPM) localization. However, *Pf*CDF-GFP showed an obscure staining pattern with
- a weak spot within the parasite cytosol in ring and trophozoite state parasites, but multiple foci
- in schizont stages (Figure 1D). To pinpoint this localization, an additional cell line with
- endogenously 3xHA-tagged *Pf*CDF was generated, confirming the focal localization of *Pf*CDF
- in asexual stages (Figure S2C).

- 152 Except for *Pf*CDF, the observed localizations of the other five transporters were confirmed by
- 153 co-localization studies using appropriate episomally expressed marker proteins: P40PX-

mCherry (39, 40) for the food vacuole, ACP-mCherry (41, 42) for apicoplast and Lyn-mCherry

155 (37, 43) for PPM. The focal distribution of PfCDF-GFP was co-localized with a rhoptry (ARO-

mCherry (44, 45)) and a micronemes (AMA1-mCherry (46, 47)) marker, but PfCDF-GFP did

not colocalize with either marker (Figure 1H). Additionally, for *Pf*ZIP and PF3D7_1135300 the

158 PPM localization was further confirmed in free merozoites (Figure S2D, E). Accordingly, as

noted above, we named PF3D7 0523800 as 'food vacuole resident transporter 1' (FVRT1)

and PF3D7 1135300 as 'plasma membrane resident transporter 1' (PMRT1).

Targeted-gene disruption (TGD), conditional knockdown and conditional knockout of

putative transporters

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In order to test whether the putative transporters are essential for *P. falciparum* during its intraerythrocytic cycle, we first tried to functionally inactivate them by targeted gene disruption (TGD) using the SLI system (37) (Figure S3A). TGD cell lines were successfully obtained for *Pf*ZIP1 and *Pf*CDF (Figure S3B, C). For *Pf*ZIP1-TGD, the correct integration of the plasmid

into the genomic locus and absence of wildtype locus was verified by PCR and subsequent

growth experiments revealed no growth defect compared to *P. falciparum* 3D7 wildtype

parasites (Figure S2B), suggesting its redundancy during asexual parasite proliferation. For

170 PfCDF-TGD the correct integration of the plasmid into the genomic locus was also verified, but

wildtype DNA was still detectable and remained even upon prolonged culturing under

172 G418/WR selection and limited dilution cloning (Figure S3C). In contrast, six (PfPMRT1,

173 PfDMT2) or eight (PfFVRT1, PfMFS6) independent attempts to obtain TGD cell lines for the

other four transporters with the respective plasmids failed, indicating that these genes have an

indispensable role in blood stage parasite growth.

176 To probe into the function of the putative transporters where we were unable to generate gene-

knockout parasites, we utilized the glmS ribozyme sequence. The corresponding sequence

was integrated into the 3'UTR of the targeted genes. This enabled the induction of conditional

degradation of respective mRNAs upon addition of glucosamine (38) and the assessment of

180 the phenotypic consequences. Upon addition of 2.5 mM glucosamine to young ring stage

parasites we found a 76.8% (+/- SD 3.7) reduction in GFP fluorescence intensity in *Pf*DMT2-

182 GFP parasites, 72.7% (+/- SD 9.4) reduction in *Pf*MFS6-GFP and a 77.7% (+/- SD 6.1)

reduction in PfPMRT1-GFP in schizonts of the same cycle (Figure 2A-C, S4A-C). No

measurable reduction in fluorescence intensity could be detected for PfFVRT1-GFP or PfCDF-

185 GFP expressing parasite lines (Figure S4D-F). For parasite cell lines with a significant

reduction in the expression of the endogenously tagged protein, proliferation was analyzed in

the absence and presence of 2.5 mM glucosamine (Figure 2D, S4G). While no significant effect

on growth was observed for PfMFS6 and PfPMRT1, a growth reduction of 68.5 % (+/- SD 2.1)

over two cycles was observed upon knockdown of PfDMT2. For PfPMRT1, a minor growth

delay was measurable, which resulted in a reduced parasitemia at day 3 upon knockdown

using both, 2.5 mM or 5 mM glucosamine (Figure 2E), and fewer newly formed ring stage parasites at 84 hours post invasion (hpi) (Figure 2F) were observed.

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To better characterize the minor growth phenotype of PfPMRT1-GFP-glmS parasites that might be due to incomplete knockdown, we generated a conditional PfPMRT1 knockout cell line (condΔPMRT1) using the Dimerizable Cre (DiCre) system (48, 49). Again using the SLI system (37), the endogenous PfPMRT1 was disrupted upstream of the region encoding the Nterminal transmembrane domain, but, at the same time introducing a recodonized second functional copy of PfPMRT1 flanked by loxP sites in the genomic locus. This loxP-flanked allelic copy of PfPMRT1 encodes an additional 3x hemagglutinin (HA) tag, which can be conditionally excised upon addition of a rapamycin analog (rapalog) via the enzymatic activity of an episomally expressed DiCre (Figure 3A). First, correct integration of the plasmid into the genomic locus was verified by PCR (Figure 3B). Second, expression and localization of the recodonized HA-tagged protein at the PPM was verified by colocalization with the merozoite plasma membrane marker MSP1 (50) (Figure 3C). Third, excision of the recodonized gene upon rapalog addition was confirmed on genomic level by PCR (Figure 3D) and on protein level by Western blot analysis at 24 hpi and 48 hpi (Figure 3E). To assess the effect of conditional PfPMRT1 knockout on parasite proliferation, we determined growth of the transgenic parasite cell line with and without rapalog over five days (Figure 3F, S5A). In contrast to the glmS-based knockdown experiment, DiCre-based gene excision (induced by the addition of rapalog to young ring stages of cond∆PMRT1 parasite cell cultures) abolished growth within the first replication cycle (Figure 3F, S5A). The specificity of the observed growth phenotype was verified by gene complementation. To achieve this, we episomally expressed recodonized PfPMRT1 with TY1-epitope tag either under the constitutive nmd3 or the weaker sf3a2 promoter (51) in the condΔPMRT1 cell line (Figure 3D, F, S5B, C). Correct localization of the TY1-tagged PfPMRT1 at the PPM was verified by immunofluorescence assays (IFA) (Figure 3G). Notably both, complementation of the *Pf*PMRT1 knockout cell line (cond Δ PMRT1) with recodonized PfPMRT1 either under control of the constitutive nmd3 or the weaker sf3a2 promoter, restored parasite growth (Figure 3F, S5B, C). The level of growth restoration with low level expression of recodonized *Pf*PMRT1 is in line with the results from glmS-knockdown experiments, which showed that a reduction of about 75% in protein expression resulted only in a minor growth perturbation (Figure 2C, D).

Loss of the PPM-localized PfPMRT1 leads to an arrest of parasite development at

223 trophozoite stage and the formation of vacuoles

To determine, which particular parasite stages are affected by the knockout of PfPMRT1, we

added rapalog to tightly synchronized parasites at different time points (4, 20 and 32 hpi)

226 (Figure 4A) and monitored parasite growth by flow cytometry. Additionally, we quantified

227 growth perturbation by microscopy of Giemsa smears at 4, 20, 24, 32, 40, 48, 72 and 96 hpi

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(Figure 4B, S6A, B). When adding rapalog at 4 hpi, parasite development progressed through ring and early trophozoite stages up to 24 hpi with no visible abnormality. Afterwards, parasites with deformed and enlarged vacuoles started to appear and further development occurred to be stalled. At 32 hpi, almost all parasites had developed to late trophozoites/early schizonts in the control, whereas these stages were completely absent in *Pf*PMRT1-deficient parasites. Over 50% of the parasites were pycnotic or possessed large vacuoles, the remaining parasites stayed arrested at the trophozoite stage. Quantification of the percentage of parasites with vacuoles between 20 hpi and 32 hpi revealed 94.8% (+/- SD 4.0) vacuole-positive parasites (Figure 4C). The activation of gene excision at later time points by adding rapalog at 20 hpi or 32 hpi resulted in no or minor growth perturbation in the first cycle with successful re-invasion, but again led to parasites arresting at the trophozoite stage in the second cycle with an accumulation of vacuoles (Figure 4A, S6A, B). In order to get further insights into the morphological changes in PfPMRT1-deficient parasites with its vacuolization, we incubated these parasites with dihydroethidium (DHE) to visualize the parasite cytosol (40). We observed an absence of staining in the vacuoles, suggesting they are not filled with parasite cytosol (Figure 4D). Next, we transfected the condΔPMRT1 cell line with a plasmid encoding the PPM marker Lyn-mCherry (37) and observed Lyn-mCherrypositive vacuoles upon knockout of PfPMRT1 starting to become visible at 24 hpi, indicating that the vacuoles originate from the PPM (Figure 4E). In line with this, vacuole membranes were also stainable with BODIPY TR C5 ceramide in condΔPMRT1 parasites at 32 hpi (Figure 4F).

Depletion of PfPMRT1 results in an early arrest of gametocyte development

RNA-seq data suggest *Pf*PMRT1 is also expressed during other developmental stages, such as gametocytes (52, 53). Therefore, we assessed expression of *Pf*PMRT1-GFP during gametocytogenesis by re-engineering *Pf*PMRT1-GFP-glmS in the inducible gametocyte producer (iGP) '3D7-iGP (55)' parasite line, which allows the robust induction of sexual commitment by conditional expression of gametocyte development 1 protein (GDV1) upon addition of shield-1 (55) (Figure S7A). We show that *Pf*PMRT1 is indeed expressed during all stages of gametocytogenesis and again localizes to the PPM, colocalizing with the PPM-marker Lyn-mCherry (37) (Figure 5A, B). Conditional knockdown of *Pf*PMRT1 via the glmS-ribozyme system (Figure S7B) resulted in a reduction in *Pf*PMRT1-GFP fluorescence intensity of 79.4% (+/- SD 9.2%) at 7 days post induction (dpi) or 75.5% (+/- SD 23.2%) at 10 dpi, without an effect on gametocyte development (Figure S7C–F). In order to exclude that a role of *Pf*PMRT1 in gametocytogenesis is covered by only a partial knockdown resulting in low levels of expressed protein and to determine if *Pf*PMRT1 is essential for gametocytogenesis, we episomally expressed GDV1-GFP-DD in the condΔPMRT1 parasite line, enabling conditional induction of sexual commitment upon addition of shield 1 in these parasites (56). Conditional

knockout of *Pf*PMRT1 in these transgenic parasites at day three post gametocyte induction resulted in pycnotic parasites from day 5 onwards, while excision of *Pf*PMRT1 at day 5 post induction had no effect on gametocyte development (Figure 5C, D). Excision of the recodonized gene upon rapalog addition was confirmed at a genomic level by PCR for both conditions (Figure 5E). Quantification of parasite stages at day 10 post induction of GDV1 expression revealed 77.9% (+/- SD 7.7%) gametocytes and 22.1% (+/- SD 7.7%) pycnotic parasites in the control, while 100% of parasites were already pycnotic in the cultures, with induced knockout by addition of rapalog at day 3 post gametocyte induction by GDV1 expression (Figure 5F). This data indicates that *Pf*PMRT1 is important for early gametocyte development.

PMRT1 is unique to the genus *Plasmodium* and interspecies complementation assays

showed partial functional conservation

PfPMRT1 shows a lack of sequence similarities with known or putative transporters and/or conserved domains shared with known transporter families (2, 5). Our phylogenetic analysis revealed that homologs of PfPMRT1 are present across Plasmodium species with amino acid sequence identities of about 90% in the subgenus Laverania, but about 50% outside Laverania (Figure 6A). However, prediction of the protein structure using AlphaFold (35) indicates two bundles of four transmembrane helices with reasonable similarity of the C-terminal bundle with the photosynthetic reaction center Maquette-3 protein (57) (RMSD of 3.12) (Figure 6B, Figure S1B). In order to test for functional conservation, we expressed the PfPMRT1 homologs of P. vivax (PVP01_0936100) and P. knowlesi (PKNH_0933400) episomally as C-terminal Ty-1 fusion proteins under the nmd3 promoter in the condΔPMRT1 parasites. Both protein variants localized correctly at the PPM, as shown by IFA (Figure 6C), and were able to partially restore growth after two cycles to 64.8% (+/- SD 9.8%) and 65.1% (+/- SD 7.4%) compared to the control (Figure 6D, S8). Excision of the recodonized endogenous Pfpmrt1 gene upon rapalog addition was confirmed at a genomic level by PCR (Figure 6E). These data indicate that PMRT1 is functionally conserved within the genus Plasmodium.

Functional characterization of PfPMRT1 in Xenopus oocytes revealed

293 hyperpolarization-activated ion transport activity

PfPMRT1 encodes a 410 amino acid protein with eight predicted (58) transmembrane domains (TM) (Figure S1). The N- and C-terminal parts of PfPMRT1 are both predicted (59) to be facing the cytosolic side of the parasite. Surface electrostatics indicate a clear polarity of PfPMRT1 with negative charges facing the parasitophorous vacuole (PV) lumen and positive charges inside the parasite cytosol (Figure 7A). The loops protruding into the PV lumen of PfPMRT1 are generally larger than the cytosolic loops and possess several negatively charged amino acids, especially the first loop between TM1 and TM2 with 10 negatively charged amino acids.

In contrast, the loop between TM4 and TM5 on the parasite cytosol (PC) side contains six positively charged amino acids. These two stretches of charged amino acids were hypothesized to be important for protein function and were replaced with neutral amino acids to test this hypothesis (Figure 7B). Full-length, WT PfPMRT1 as well as the two mutants (MutPV, with sequence EDDIDEYNIKGNEEE modified to QNAIAQYNIKGNAAA; MutPC, with sequence KIEKHFRKKF modified to AIQAAFAMAF), each as C-terminal eYFP fusion proteins, were expressed in Xenopus laevis oocytes in order to characterize its function (Figure 7D). The PfPMRT1-WT expressing oocytes showed significant inward currents when clamped at -160 mV, indicating a hyperpolarization-activated ion transport (Figure 7C). The current of MutPV-expressing oocytes was significantly reduced compared to PfPMRT1-WT ($W = 10, n_1$ = 10, n_2 = 7, P = 0.0147, two tailed Wilcoxon rank sum test), while in MutPC-expressing oocytes the observed current was only slightly smaller than in WT controls (Figure 7D, E), suggesting that the negatively charged amino acids of the first loop protruding into the PV lumen are important for the function of PfPMRT1. To translate these results to the parasite, we expressed PfPMRT1-WT and the two sequence mutants as C-terminal Ty-1 fusion complementation constructs under the *nmd3* promoter in condΔ*Pf*PMRT1 parasites. These experiments showed, that PfPMRT1-MutPC was able to completely restore parasite growth, while PfPMRT1-MutPV showed only 59.2% (SD +/- 9.5) growth compared to the control after two developmental cycles (Figure 7F, G, S9 A-C). These data support the functional role of the negatively charged amino acids of the first loop protruding into the parasitophorous vacuole lumen for transport activity during the intraerythrocytic developmental cycle of the parasite.

Discussion

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To date, the predicted 'transportome' of *P. falciparum* consists of 117 putative transport systems (encoded by 144 genes) classified as channels (n=19), carriers (n=69), and pumps (n=29) (2). Functions of the vast majority of transporter genes were inferred from sequence homology to model organisms, however, given their lack of homology, 39 gene products could not be associated with any functional or subcellular localization and were categorized as orphan transporters accordingly (4). Here, we concentrated on four candidates from the orphan group, which were already partially characterized in *P. berghei*, and included two additional so far uncharacterized putative transporter proteins.

far uncharacterized putative transporter proteins.

We localized *Pf*FVRT1-GFP – annotated on PlasmoDB (60) as putative divalent metal transporter – at the food vacuole of the parasite, which is in line with a previously predicted food vacuole association (1) and its reported homology (1, 61) to the conserved eukaryotic endosomal/lysosomal natural resistance-associated macrophage protein (NRAMP) transporter (62) in our structure similarity search. Repeated attempts to generate a TGD cell line failed, indicating an important role of this transporter during asexual blood stage

337 development which is in agreement with data from a P. falciparum genome wide essentiality 338 screen (63). 339 In concordance with recently published data identifying PbDMT2 and PbMFS6 as leaderless 340 apicoplast transporters (29), we localized GFP-fusion proteins of PfDMT2 and PfMFS6 at the 341 apicoplast. Successful knockdown of PfDMT2 resulted in a growth defect in the second cycle 342 after induction, resembling the described delayed death phenotype of other apicoplast genes 343 that were functionally inactivated (29, 64-66). It suggests an essential role of PfDMT2 in 344 apicoplast physiology, as observed by Sayers et al. (29) for the rodent malaria P. berghei. This 345 is further supported by our failed attempts to disrupt this gene using the SLI system. 346 We also failed to disrupt the PfMFS6 locus, which is in agreement with the gene knockout 347 studies in P. berghei that led to a markedly decreased multiplication rate (28, 29, 67). Nevertheless, glmS-based knock-down, although comparable to PfDMT2-GFP knockdown 348 349 (72.7% versus 76.8% reduction in GFP fluorescence, respectively) had no effect on parasite 350 proliferation in our study. This might indicate that these reduced levels of PfMFS6, in contrast 351 to reduced levels of PfDMT2, are sufficient for normal asexual replication in vitro. 352 Another candidate, PfCDF, annotated as putative cation diffusion facilitator family protein, 353 showed multiple cytosolic foci within the parasite with no co-localization with apical organelle 354 markers. The homologue in Toxoplasma gondii, TqZnT (TqGT1 251630) shows a similar cellular distribution (68). It has recently been shown to transport Zn²⁺, to localize to vesicles at 355 356 the plant-like vacuole in extracellular tachyzoites and to be present at dispersed vesicles throughout the cytoplasm of intracellular tachyzoites (68). Our data suggests that PfCDF is not 357 358 essential for parasite in vitro blood stage development even though we were not able to 359 generate a clonal wild-type free TGD cell line. Its redundancy is further supported by published 360 data, showing that CDF proteins are non-essential for in vivo blood stage development in P. 361 yoelli (69) and P. berghei (28, 67) and by its high (1.0) mutagenesis index score in a P. 362 falciparum genome-wide mutagenesis screen (63). Finally, two putative transporters, PfZIP1 and PfPMRT1, localized to the PPM. We show that 363 364 PfZIP1 is non-essential for P. falciparum in vitro blood stage development, in line with a high 365 (0.7) mutagenesis index score in a P. falciparum genome-wide mutagenesis screen (63). 366 However, this is in contrast to the reported strong fitness loss in P. berghei (67) knockout 367 mutants and failed knockout attempts in P. yoelli and P. berghei in vivo mouse models (29, 69). These observations may reflect differences between *Plasmodium* species or differing 368 369 requirements for *in vitro* and *in vivo* growth conditions. 370 PfPMRT1 is annotated as a conserved Plasmodium membrane protein with unknown function. 371 It has been described as a protein showing structural characteristics of a transporter, without 372 sharing sequence similarities with known or putative transporters and/or conserved domains 373 of known transporter families (2, 5). Our phylogenetic analysis confirmed PMRT1 as unique

for Plasmodium species with high sequence conservation only within the Laverania subgenus (70). In line with data from genome-wide mutagenesis screens (63, 67) and reported failed knockout attempts in P. yoelli (69), we found that PfPMRT1 is essential for parasite growth, as its functional inactivation resulted in growth arrest at the trophozoite stage accompanied by the accumulation of PPM-derived vacuoles within the parasite. In contrast, conditional knockdown resulted only in a growth delay, indicating that minor residual PfPMRT1 protein levels appear to be sufficient to promote parasite growth. This finding was validated by episomal expression of an allelic copy under the control of the weak sf3a2 promoter (51) in the PfPMRT1 knockout parasites. Additionally, we found that PfPMRT1 is essential for early gametocytogenesis. Interestingly, the induction of the knockout at stage II–III had no effect on gametocytogenesis. This might be due to sufficient amounts of PfPMRT1 already present at the PPM, but could also indicate that the function of the transporter is not required for later stage gametocyte maturation. The PfPMRT1-WT-expressing Xenopus oocytes showed a hyperpolarization-activated ion transport activity, which is reduced in the variant featuring mutations in the first loop that extends into the parasitophorous vacuole. But it remains to be determined whether the hyperpolarization-activated ion transporting activity is the main physiological function of PfPMRT1. Additional functions such as transporting of other molecules, or cooperative interaction with other parasite proteins might play a role under physiological conditions. However, the mutation analysis suggested that the negatively charged amino acids of the first loop protruding into the PV lumen are important for the function of *Pf*PMRT1. Interestingly, AlphaFold predicts smaller helical structures within this negatively charged loop, which possibly function as flexible gates and/or could be involved substrate binding. For future work, further functional and pharmacological characterization of this transporter will provide insights into its biological role in different stages of the parasites life cycle, as transcriptomic data indicates – along with expression in blood stages (31, 32) – PfPMRT1 is expressed in oocysts of P. falciparum (54, 71) and P. berghei (72).

Material and methods

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Cloning of plasmid constructs for parasite transfection

- 404 For endogenous tagging using the SLI system (37) a 889 bp (for *Pf*PMRT1; PF3D7_1135300),
- 405 905 bp (*Pf*FVRT1; PF3D7 0523800), 827bp (*Pf*ZIP1; PF3D7 0609100), 873 bp (*Pf*DMT2;
- 406 PF3D7_0716900), 877 bp (*Pf*MFS6; PF3D7_1440800), 785 bp (*Pf*CDF; PF3D7_0715900)
- 407 long homology region (HR) was amplified using 3D7 gDNA and cloned into pSLI-GFP-glmS
- 408 (73) (derived from pSLI-GFP (37)) using the Notl/Mlul restriction site. In order to generate

- 409 PfPMRT1-2xFKBP-GFP a 1000 bp long HR was amplified using 3D7 gDNA and cloned into
- 410 pSLI-2xFKBP-GFP (37).
- 411 For SLI-based targeted gene disruption (SLI-TGD) (37) a 501 bp (PfPMRT1), 378 bp
- 412 (PfFVRT1), 511 bp (PfZIP1), 399 bp (PfDMT2), 396 bp (PfMFS6), 741 bp (PfCDF) long
- 413 homology region was amplified using 3D7 gDNA and cloned into the pSLI-TGD plasmid (37)
- 414 using Notl and Mlul restriction sites.
- For conditional deletion of *Pf*PMRT1, the first 492 bp of the *Pf*PMRT1 gene were PCR amplified
- 416 to append a first loxP site and a recodonized T2A skip peptide. The recodonized full-length
- 417 coding region of PfPMRT1 was synthesized (GenScript, Piscataway, NJ, USA) and PCR
- 418 amplified with primers to add a second loxP site after the gene to obtain a second fragment.
- Both fragments were cloned into pSLI-3xHA (51), using Notl/SpeI and AvrII/XmaI sites. This
- 420 resulted in plasmid pSLI-PfPMRT1-loxP and the resulting transgenic cell line after successful
- 421 genomic modification was transfected with pSkip-Flox (37) using 2 μg/ml Blasticidin S to obtain
- 422 a line expressing the DiCre fragments (cond∆PMRT1).
- 423 For complementation constructs, the recodonized PfPMRT1 gene was PCR amplified using
- 424 primers to append the TY1 sequence and cloned via Xhol and AvrII or KpnI into pEXP1comp
- 425 (51) containing yDHODH as a resistance marker and different promoters (nmd3
- 426 (PF3D7_0729300), sf3a2 (PF3D7_0619900)) driving expression of the expression cassette.
- 427 This resulted in plasmids c-nmdrPfPMRT1-ty1 and c-sf3a2PfPMRT1-ty1.
- 428 PfPMRT1 homologues of P. vivax (PVP01_0936100) (74) and P. knowlesi (PKNH_0933400)
- 429 (75) were amplified from parasite qDNA and cloned into p^{nmd3}EXP1comp (51) via the Xhol/AvrII
- 430 restriction site. For co-localization experiments the plasmids pLyn-FRB-mCherry (37), P40PX-
- mCherry (40), pARL-crtACP-mCherry (42), pARL-ama1ARO-mCherry (44) and pARL-ama1AMA1-
- 432 mCherry (47) were used. For conditional gametocyte induction yDHODH was amplified by
- 433 PCR from pARL-ama1AMA1-mCherry-yDHODH (47) and cloned into GDV1-GFP-DD-
- 434 hDHFR(56)(56) using the Xhol/Xhol restriction site. For expression in Xenopus oocysts
- 435 PfPMRT1 was amplified from c-nmdrPfPMRT1-ty1 and DNA fragments were ligated and
- inserted into the oocyte expression vector pGEM-HE (76), containing a C-terminal eYFP using
- 437 the BamHI/Xhol restriction site. As a control of plasma membrane-targeted channel protein,
- 438 the cyclic nucleotide-gated potassium channel SthK (77) was also cloned into the pGEM-
- 439 HE vector with C-terminal eYFP using the BamHI/XhoI restriction site.
- The pGEM-HE plasmids containing SthK, PfPMRT1 or the mutant were linearized by Nhel
- 441 digestion before the RNA generation. Mutations were made by QuikChange Site-Directed
- 442 Mutagenesis. The complementary RNA (cRNA) was synthesized by in vitro transcription using
- the AmpliCap-MaxT7 High Yield Message Maker Kit (Epicentre).
- 444 Sequences of the MutPC and MutPV variants were amplified from pGEM plasmid constructs
- and cloned into c-nmd3PfPMRT1-ty1 using the Xhol/AvrII restriction site.

Oligonucleotides and plasmids used in this study are listed in Table S1A and S1B.

P. falciparum culture and transfection

- Blood stages of *P. falciparum* 3D7 were cultured in human erythrocytes (O+). Cultures were
- 449 maintained at 37°C in an atmosphere of 1% O₂, 5% CO₂ and 94% N₂ using RPMI complete
- 450 medium containing 0.5% Albumax according to standard protocols (78). To maintain
- 451 synchronized parasites, cultures were treated with 5% sorbitol (79).
- Induction of gametocytogenesis was done as previously described (55, 56). Briefly, GDV1-
- 453 GFP-DD expression was achieved by addition of 4 µM shield-1 to the culture medium and
- 454 gametocyte cultures were treated with 50 mM N-acetyl-D-glucosamine (GlcNAc) for five days
- starting 72 hours post shield-1 addition to eliminate asexual parasites(80). Alternatively,
- 456 asexual ring stage cultures with >10% parasitemia were synchronized with Sorbitol (79)
- cultured for 24 hours and treated with 50 mM N-acetyl-D-glucosamine (GlcNAc) (80) for five
- 458 days.

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- 459 For transfection, Percoll-purified (81) late-schizont-stage parasites were transfected with 50
- 460 µg of plasmid DNA using Amaxa Nucleofector 2b (Lonza, Switzerland) as previously
- 461 described(82). Transfectants were selected either using 4 nM WR99210 (Jacobus
- 462 Pharmaceuticals), 2 μg/ml Blasticidin S (Life Technologies, USA), or 0.9 μM DSM1 (83) (BEI
- Resources; https://www.beiresources.org). In order to select for parasites carrying the genomic
- 464 modification using the SLI system (37), G418 (Sigma-Aldrich, St. Louis, MO) at a final
- 465 concentration of 400 μg/ml was added to 5% parasitemia culture. The selection process and
- 466 testing for integration were performed as previously described (37).
- 467 For SLI-TGD, a total of six (*Pf*PMRT1, *Pf*DMT2, *Pf*ZIP1, *Pf*CDF) or eight (*Pf*FVRT1, *Pf*MFS6)
- independent 5 ml cultures containing the episomal plasmid were selected under G418 for at
- 469 least eight weeks.

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Imaging and immunofluorescence analysis (IFA)

- 471 Images of Xenopus oocysts were taken 3 days after injection using a Carl Zeiss LSM 5
- Pascal Laser Scanning Confocal Microscope, with a 40x/0.8 water immersion objective. The
- eYFP fluorescence was excited at 514 nm of the Argon laser and obtained at 530 600 nm.
- 474 Fluorescence images of infected erythrocytes were observed and captured using a Zeiss
- 475 Axioskop 2plus microscope with a Hamamatsu Digital camera (Model C4742-95), a Leica D6B
- 476 fluorescence microscope equipped with a Leica DFC9000 GT camera and a Leica Plan
- 477 Apochromat 100x/1.4 oil objective or an Olympus FV3000 with a x100 MPLAPON oil objective
- 478 (NA 1.4).
- 479 Microscopy of unfixed IEs was performed as previously described (84). Briefly, parasites were
- 480 incubated in RPMI1640 culture medium with Hoechst-33342 (Invitrogen) for 15 minutes at
- 481 37°C prior to imaging. 7 μl of IEs were added on a glass slide and covered with a cover slip.

BODIPY TR C5 ceramide (Invitrogen) staining was performed by adding the dye to 32 hours 482 post invasion parasites in a final concentration of 2.5 µM in RPMI as previously described (84). 483 484 For DHE staining of the parasite cytosol (40), 80 µl of resuspended parasite culture were 485 incubated with DHE at a final concentration of 4.5 µg/ml in the dark for 15 minutes prior to 486 imaging. 487 IFAs were performed as described previously (85). Briefly, IEs were smeared on slides and 488 air-dried. Cells were fixed in 100% ice cold methanol for 3 minutes at -20°C. Afterwards, cells 489 were blocked with 5% milk powder for 30 minutes. Next primary antibodies were diluted in 490 PBS/3% milk powder and incubated for 2 hours, followed by three washing steps in PBS. 491 Secondary antibodies were applied for 2 hours in PBS/3% milk powder containing 1 µg/ml 492 Hoechst-33342 (Invitrogen) or DAPI (Roche) for nuclei staining, followed by 3 washes with 493 PBS. One drop of mounting medium (Mowiol 4-88 (Calbiochem)) was added and the slide 494 sealed with a coverslip for imaging. To assess the localisation of the endogenously HA-tagged PfPMRT1 IFAs were performed in 495 496 suspension with Compound 2-stalled schizonts (86) to distinguish protein located at the PPM 497 from that located at the PVM as previously done (51, 87). For this, trophozoite stages were 498 treated with Compound 2 (1 µM) overnight, and arrested schizonts were harvested, washed in 499 PBS, and fixed with 4% paraformaldehyde/0.0075% glutaraldehyde in PBS. Cells were 500 permeabilized with 0.5% Triton X-100 in PBS, blocked with 3% BSA in PBS, and incubated 501 overnight with primary antibodies diluted in 3% BSA in PBS. Cells were washed 3 times with 502 PBS and incubated for 1 hour with Alexa 488 nm or Alexa 594 nm conjugated secondary 503 antibodies specific for human and rat IgG (Invitrogen) diluted 1:2,000 in 3% BSA in PBS and 504 containing 1 µg/ml DAPI. Cells were directly imaged after washing 5 times with PBS 505 Antisera used: 1:200 mouse anti-GFP clones 7.1 and 13.1 (Roche), 1:500 rat anti-HA clone 506 3F10 (Roche), 1:1000 human anti-MSP1 (88), 1:10000 mouse anti-TY1 (ThermoFischer 507 Scientific Cat.No: MA5-23513). Contrast and intensities were linear adjusted if necessary and 508 cropped images were assembled as panels using Fiji (89), Corel Photo-Paint X6 and Adobe 509 Photoshop CC 2021. 510 **Immunoblots** 511 Immunoblots were performed using saponin lysed infected erythrocytes. Parasite proteins 512 were separated on a 10% SDS-PAGE gel using standard procedures and transferred to a nitrocellulose membrane (Amersham™Protran™ 0.45 µm NC, GE Healthcare) using a 513 514 transblot device (Bio-Rad) according to manufacturer's instructions. 515 Rabbit anti-aldolase (90) and anti-sbp1 (90) antibodies were diluted 1:2,000, mouse anti-GFP 516 clones 7.1 and 13.1 (Roche) antibody was diluted 1:500 or 1:1,000 and rat anti-HA clone 3F10

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(Roche) antibody was diluted 1:1,000.

The chemiluminescent signal of the HRP-coupled secondary antibodies (Dianova) was 519 visualized using a Chemi Doc XRS imaging system (Bio-Rad) and processed with Image Lab 520 Software 5.2 (Bio-Rad). To perform loading controls and ensure equal loading of parasite material anti-aldolase antibodies were used. The corresponding immunoblots were incubated two times in stripping buffer (0.2 M glycine, 50 mM DTT, 0.05% Tween 20) at 55°C for 1 hour and washed 3 times with TBS for 10 minutes.

Growth Assay

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A flow cytometry-based assay adapted from previously published assays (40, 91) was performed. For this, parasite cultures were resuspended and 20 µl samples were transferred to an Eppendorf tube. 80 µl RPMI containing Hoechst-33342 and dihydroethidium (DHE) was added to obtain final concentrations of 5 µg/ml and 4.5 µg/ml, respectively. Samples were incubated for 20 minutes (protected from UV light) at room temperature, and parasitemia was determined using an LSRII flow cytometer by counting 100,000 events using the FACSDiva software (BD Biosciences) or using an ACEA NovoCyte flow cytometer.

Stage distribution assay

In order to obtain tightly synchronized parasite cultures, percoll purified schizonts(81) were cultured for four hours together with fresh erythrocytes, followed by sorbitol synchronization and resulting in a four-hour age window of parasites. Next, the culture was divided in four dishes and rapalog was added at a final concentration of 250 nM immediately to one dish and at 20 hours post invasion (hpi) and 32 hpi to the respective dishes. Giemsa smears and samples for flow cytometry were collected at the indicated timepoints. The parasitemia was determined using a flow cytometry assay and the stages were determined microscopically counting at least 50 infected erythrocytes per sample and timepoint.

Gametocyte stage distribution assay

542 Giemsa-stained blood smears 10 days post induction of GDV1 expression were obtained and 543 at least 10 fields of view were recorded using a 63x objective per treatment and time point. 544 Erythrocyte numbers were then determined using the automated Parasitemia software (http://www.gburri.org/parasitemia/) while the number of gametocytes, pycnotic and asexual 545 546 parasites was determined manually in >1800 erythrocytes per sample. This assay was done 547 blinded.

GImS-based knockdown

GlmS based knockdown assay was adapted from previously published assays (38, 73). To induce knockdown 2.5 or 5 mM glucosamine was added to highly synchronous early rings stage parasites. As a control, the same amount of glucosamine was also added to 3D7 wildtype

- parasites. For all analyses, the growth medium was changed daily, and fresh glucosamine
- were added every day.
- 554 Knockdown was quantified by fluorescence live cell microscopy at day 1 and 3 of the growth
- assay. Parasites with similar size were imaged, and fluorescence was captured with the same
- 556 acquisition settings to obtain comparable measurements of the fluorescence intensity.
- 557 Fluorescence intensity (integrated density) was measured with Fiji(89), and background was
- subtracted in each image. The data were analyzed with Graph Pad Prism version 8.
- 559 GlmS based knockdown experiments in gametocytes were performed as described previously
- 560 (92). Briefly, synchronized ring stage cultures were induced by the addition of shield-1. At day
- 3 post induction the culture was spilt into two dishes and one dish was cultured in the presence
- of 2.5 mM glucosamine for the remaining ten days. Knockdown was quantified by fluorescence
- live cell microscopy at day 7 and 10 post induction, as described above and gametocyte
- 564 parasitemia was determined at day 10 post induction using the automated Parasitemia
- software (http://www.gburri.org/parasitemia/).

DiCre mediated conditional knockout

- 567 The parasites containing the integrated pSLI-PfPMRT1-loxP construct were transfected with
- 568 pSkip-Flox (37) using 2 μg/ml Blasticidin S to obtain a line expressing the DiCre fragments. To
- induce excision, the tightly synchronized parasites (detailed description see growth assay)
- 570 were split into 2 dishes and rapalog was added to one dish (Clontech, Mountain View, CA) to
- a final concentration of 250 nM. The untreated dish served as control culture. Excision was
- verified at genomic level after 24 and 48 hours of cultivation by PCR and on protein level by
- 573 Western blot using anti-HA antibodies.

Phylogenetic analysis

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- 575 A blastp search of the PMRT1 sequence (PlasmoDB (60): PF3D7 1135300; UniProt: Q8II12)
- 576 was performed against the nr database (9 May 2021) using Geneious Prime 2021.2.2
- 577 (https://www.geneious.com) and an E-value of 10e-0 (BLOSUM62 substitution matrix). Blast
- 578 hits were filtered for sequences from taxa represented in the currently favored haemosporidian
- parasite phylogeny (93). The phylogeny derived from an amino acid alignment using Bayesian
- 580 framework with a partitioned supermatrix and a relaxed molecular clock
- 581 (18_amino_acid_partitioned_BEAST_relaxed_clock_no_outgroup.tre; (93)) was visualized
- with associated data using the R package ggtree v3.3.0.900 (94, 95). A multiple protein
- 583 sequence alignment of PMRT1 and homologous sequences was performed using MAFFT
- v7.490 (96) using the G-INS-I algorithm to obtain a highly accurate alignment. Protein statistics
- 585 were calculated using Geneious Prime 2021.2.2 (https://www.geneious.com) and EMBOSS
- 586 pepstats v6.6.0.0 (97).

Prediction of protein structures

AlphaFold structure predictions (35) were retrieved from https://alphafold.ebi.ac.uk and the PDB used for DALI protein structure homology search (36). PyMOL Molecular Graphics System, Version 2.5.2 Schrödinger was used for visualization of all structures, generation of figures and the calculation of the root mean square deviation (RMSD) between the predicted crystal structure of PfPMRT1 and the Maquette-3 protein (PDB: 5vit (57)) by cealign. The Adaptive Poisson-Boltzmann Solver (APBS) within PyMOL was used to predict the surface 594 electrostatics of PfPMRT1.

Electrophysiology experiments

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- Stage V and VI oocytes were surgically removed from female Xenopus laevis by immersion in water containing 1 g/L Tricain and isolated from theca and follicle layers by digestion with 0.14 mg ml⁻¹ collagenase I. Oocytes were injected with 20 ng cRNA and were incubated at 16°C for 3 days in ND96 solution (96 mM NaCl, 2 mM KCl, 1 mM CaCl₂, 1 mM MgCl₂, 10 mM HEPES, pH7.4). Electrophysiological measurements with Xenopus oocytes were performed at room temperature (20-23°C) with a two-electrode voltage clamp amplifier (TURBO TEC-03X, npi electronic GmbH, Tamm, Germany). The bath solutions for electrophysiological recording are indicated in each figure legend. Electrode capillaries (Φ=1.5 99 mm, Wall thickness 0.178 mm, Hilgenberg) were filled with 3 M KCl, with tip openings of 0.4–1 MΩ. USB-6221 DAQ device (National Instruments) and WinWCP (v5.5.3, Strathclyde University, UK) are used for data acquisition. Origin2020 Pro and the R software environment (98) were used for data analysis. Data was visualized in R using ggplot2 (99) and patchwork (100).
- 609 Parasite icons were generated using BioRender (biorender.com) and statistical analysis was 610 performed using GraphPad Prism version 8 (GraphPad Software, USA).

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- Writing original manuscript: JSW, AB, JSTR, DW
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- All authors read and approved the manuscript.
- 643 Figures

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- 644 Figure 1: Subcellular localization of six putative P. falciparum transporters during
- 645 asexual blood stage development.
- 646 (A) Schematic representation of endogenous tagging strategy using the selection-linked
- 647 integration system (SLI). pink, human dihydrofolate dehydrogenase (hDHFR); grey, homology
- region (HR); green, green fluorescence protein (GFP) tag; dark grey, T2A skip peptide; blue,
- 649 neomycin resistance cassette; orange, glmS cassette. Stars indicate stop codons, and arrows
- depict primers (P1 to P4) used for the integration check PCR. (B-G) Localization of (B)
- 651 PfFVRT1-GFP-glmS, (C) PfZIP1-GFP-glmS, (D) PfCDF-GFP-glmS, (E) PfDMT2-GFP-glmS,
- 652 (F) PfMFS6-GFP-glmS and (G) PfPMRT1-GFP-glmS by live-cell microscopy in ring,
- trophozoite and schizont stage parasites. Nuclei were stained with Hoechst-33342. (H) Co-
- 654 localization of the GFP-tagged putative transporters with marker proteins P40PX-mCherry
- 655 (food vacuole), ACP-mCherry (apicoplast), Lyn-mCherry (parasite plasma membrane), ARO-
- 656 mCherry (rhoptry) and AMA1-mCherry (microneme) as indicated. Nuclei were stained with
- 657 Hoechst- 33342. Scale bar, 2 μm.

Figure 2: Conditional knockdown of putative transporter indicate importance of *Pf*DMT2 and *Pf*PMRT1 for parasites fitness.

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(A-C) Live cell microscopy and quantification of knockdown by measuring mean fluorescence intensity (MFI) density and size (area) of (A) PfDMT2-GFP-qlmS (B) PfMFS6-GFP-qlmS and (C) PfPMRT1-GFP-glmS parasites 40 hours after treatment without (control) or with 2.5 mM glucosamine. Scale bar, 2 µm. Statistics are displayed as mean +/- SD of three (A-B) or four (C) independent experiments and individual data points are color-coded by experiments according to Superplots guidelines (101). P-values displayed were determined with two-tailed unpaired t-test. (D) Growth of parasites treated without (control) or with 2.5 mM glucosamine determined by flow cytometry is shown as relative parasitemia values after two cycles. Shown are means +/- SD of three (PfPMRT1-GFP-glmS, PfDMT2-GFP-glmS, PfMFS6-GFP-glmS) and five (3D7 wild type parasites) independent growth experiments. P-values displayed were determined with unpaired t test with Welch correction and Benjamin Hochberg for multiple testing correction. Individual growth curves are shown in Figure S4G. (E) Growth of PfPMRT1glmS and 3D7 parasites after treatment with 2.5 mM (left panel) or 5 mM glucosamine (right panel) compared to untreated control parasites over five consecutive days. (F) Mean +/- SD distribution of ring and schizont stage parasites in PfPMRT1-glmS and 3D7 cell lines treated without (control), with 2.5 mM or 5 mM glucosamine at 80 hours post addition of glucosamine of three independent experiments.

Figure 3: *Pf*PMRT1 is essential for asexual blood stage development.

(A) Simplified schematic of DiCre-based conditional PfPMRT1 knockout using selection-linked integration (SLI). Pink, human dihydrofolate dehydrogenase (hDHFR); grey, homology region (HR); green, T2A skip peptide; light blue, recodonized *Pf*PMRT1; dark blue, 3xHA tag, yellow, neomycin phosphotransferase resistance cassette; orange, loxp sequence. Scissors indicate DiCre mediated excision sites upon addition of rapalog. Stars indicate stop codons, and arrows depict primers (P1 to P5) used for the integration check PCR and excision PCR. (B) Diagnostic PCR of unmodified wildtype and transgenic cond∆PMRT1 knock-in (KI) cell line to check for genomic integration using Primer P1-P4 as indicated in (A). (C) Immunofluorescence assay (IFA) of cond∆PMRT1 late stage schizont parasites showing localization of PfPMRT1-3xHA at the parasite plasma membrane (PPM) co-localizing with the merozoite surface protein 1 (MSP1). (D) Diagnostic PCR to verify the excision at genomic level at 24 hpi / 20 hours post rapalog addition for cond∆PMRT1 and at 48 hpi for cond∆PMRT1, c-nmd3PfPMRT1-ty1 and csf3a2PfPMRT1-ty1 parasites using Primer P1-P5 as indicated in (A). Black arrow head, original locus; red arrow head, excised locus. (E) Western blot using α-HA to verify knockout of PfPMRT1 on protein level 4, 24 and 48 hours post invasion. Expected molecular weight of PfPMRT1-3xHA: 53.3 kDa. Antibodies detecting Aldolase and SBP1 were used as loading

controls. **(F)** Growth curves of condΔPMRT1, c-nmd3PfPMRT1-ty1 and c-sf3a2PfPMRT1-ty1 parasites +/- rapalog monitored over five days by flow cytometry. One representative growth curve is depicted (replicates in Figure S5). Summary is shown as relative parasitemia values, which were obtained by dividing the parasitemia of rapalog treated cultures by the parasitemia of the corresponding untreated ones. Shown are means +/- SD of three (condΔPMRT1, c-nmd3PfPMRT1-ty1) or four (c-sf3a2PfPMRT1-ty1) independent growth experiments. **(G)** IFA of condΔPMRT1 complemented with C-terminal TY1-tagged PfPMRT1 constructs expressed either under the constitutive nmd3 or the weak sf3a2 promoter to verify PPM localization. Scale bar, 2 μm.

Figure 4: Knockout of *Pf*PMRT1 results in accumulation of PPM-derived vacuoles and growth arrest at the trophozoite stage.

(A) Parasite stage distribution in Giemsa smears displayed as heatmap showing percentage of parasite stages for tightly synchronized (+/- 2 h) 3D7 control and condΔPMRT1 (rapalog treated at 4 hpi, 20 hpi or 32 hpi) parasite cultures over two consecutive cycles. A second replicate is shown in Figure S6A (B) Giemsa smears of control and at 4 hpi rapalog treated condΔPMRT1 parasites over two cycles. Scale bar, 5 μm. (C) Live cell microscopy of 4 hour window synchronized 3D7 control and condΔPMRT1 parasites +/- rapalog stained with dihydroethidium (DHE) at 20–32 hpi. (D) Quantification of parasites displaying vacuoles (green) for 4 hour window synchronized 3D7 control and rapalog treated condΔPMRT1 parasites. Shown are percentages of normal parasites versus parasites displaying vacuoles as means +/- SD of three independent experiments. (E) Live cell microscopy of 8 hour window synchronized 3D7 control and rapalog treated condΔPMRT1 parasites, episomally expressing the PPM marker Lyn-mCherry at 24–40 hpi. (F) Live cell microscopy of 3D7 control and condΔPMRT1 parasites +/- rapalog stained with BODIPY TR C5 ceramide at 32 hpi. Scale bar, 2 μm.

Figure 5: PfPMRT1 is essential for early gametocyte development.

(A) Live cell microscopy of 3D7-iGP-*Pf*PMRT1-GFP parasites across the complete gametocyte development. White arrow heads indicate remaining GDV1-GFP signal observed in close proximity to the Hoechst signal, as previously reported (56, 92, 102, 103). (B) Live cell microscopy of *Pf*PMRT1-GFP parasites expressing the PPM marker Lyn-mCherry. Nuclei were stained with Hoechst-33342. Scale bar, 2 μm. (C) Experimental setup of gametocyte induction upon GDV1-GFP-DD expression (+shield-1) and conditional *Pf*PMRT1 knockout (+rapalog) and elimination of asexual blood stage parasites (+GlcNac). (D) Gametocyte development over 12 days of condΔPMRT1/GDV1-GFP-DD parasites without (control) or with

rapalog addition at day 3 (3 dpi) or day 5 (5 dpi) after induction of sexual commitment by conditional expression of GDV1-GFP upon addition of shield-1. Scale bar, 5 μ m. (**E**) Diagnostic PCR to verify the excision on genomic level at 5 dpi and 12 dpi. Black arrow head, original locus; red arrow head, excised locus. (**F**) Representative Giemsa smears and quantification of parasite stage distribution at day 10 post induction for parasites treated without (control) or with rapalog at day 3 post induction. For each condition parasitemia and parasite stages distribution in ($n_{control}$ = 3370, 2304, 2759 and $n_{rapalog}$ = 3010, 1830, 2387) erythrocytes were determined and are displayed as percentage (gametocyte parasitemia for control (1.3%, 0.4%, 1.1%) and for rapalog treated (0%, 0%, 0%). Nuclei were stained with Hoechst-33342. Scale bar, 10 μ m.

Figure 6: PMRT1 is a genus-specific transporter with conserved function.

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(A) Phylogenetic tree of haemosporidian parasites (modified from (93)) containing PMRT1 homologous sequences associated with data on pairwise amino acid sequence identity to PfPMRT1. The phylogeny is derived from Bayesian Inference using BEAST using a fully partitioned amino acid dataset and lognormal relaxed molecular clock (93). Silhouettes depict representatives of the vertebrate hosts for each lineage and white filled bars indicate pairwise identities of PMRT1 homologs used for subsequent complementation assays. (B) Structural alignment of predicted PfPMRT1 structure with Maguette-3 protein (PDB: 5vit) (57). Both structures have a root mean square deviation (RMSD) over the aligned α-carbon position of 3.12 over 184 residues calculated in PyMol. (C) IFA of c- nmd3Pk-ty1 and c- nmd3Pv-ty1 parasites to verify correct localization of the expressed complementation fusion proteins at the parasite plasma membrane. Nuclei were stained with Hoechst-33342. Scale bar, 2 µm. (D) Growth of cond∆PMRT1 parasites complemented with PfPMRT1 homologs from P. vivax (PVP01 0936100) and P. knowlesi (PKNH 0933400). Shown are relative parasitemia values, which were obtained by dividing the parasitemia of rapalog treated cultures by the parasitemia of the corresponding untreated controls together with means +/- SD from three c-nmd3Pf-ty1 (\(\delta\) c-nmd3PfPMRT1-tv1 Figure 3D. S5B) and six (c-nmd3Pk-tv1, c-nmd3Pv-tv1) independent growth experiments. One sample t-test (E) Diagnostic PCR to verify the excision of PfPMRT1 on genomic level at 48 hpi for c-nmd3Pf-ty1, c-nmd3Pk-ty1 and c-nmd3Pv-ty1 parasites. Black arrow head, original locus; red arrow head, excised locus.

Figure 7: In vitro characterization of PfPMRT1 in Xenopus oocytes.

- 765 (A) Surface electrostatics of the predicted PfPMRT1 structure generated by APBS within
- 766 PyMol. **(B)** Scheme of *Pf*PMRT1 membrane topology prediction and the two mutation sites:
- Outer loop mutant (MutPV) between TM 1 2 with negative charged amino acids (indicated in

red) and inner loop mutant (MutPC) between TM 4 - 5 with positive charged amino acids (indicated in blue) were changed to neutral amino acids. The cartoon was generated by Protter(104) and PyMol. (C) Confocal images of a non-injected oocyte, PfPMRT1, MutPV and MutPC expressing oocytes. Scale bar, 100 µm. (D) Superimposed traces of control (injected with a cyclic nucleotide-gated potassium channel SthK(77)), PfPMRT1, MutPV and MutPC expressing oocytes with the protocol shown in the upper left. The grey traces are original recordings from 7 to 10 oocytes, averaged traces are shown in black. Bath solution contains (in mM): 120 NMG CI, 1 MgCl₂, 2 CaCl₂, 10 HEPES, pH7.6. (E) Current-voltage (I-V) curves of data points from (D) taken at the end of each holding potential. Both, MutPV and the SthKcontrol are significantly different from wild type PfPMRT1 with P values of 0.017 and 0.014, respectively (Mean \pm SD, n = 7 - 10). (F) Diagnostic PCR to verify the excision on genomic level at 48 hours post rapalog addition in cond∆PMRT1, c-nmd3MutPC-ty1 and c-nmd3MutPV-ty1 parasites +/- rapalog. Black arrow head, original locus; red arrow head, excised locus. (G) Parasite growth shown as relative parasitemia values, which were obtained by dividing the parasitemia of rapalog treated cultures by the parasitemia of the corresponding untreated controls. Shown are means +/- SD of four (c-sf3a2ty1), three (c-nmd3ty1, c-nmd3MutPC-ty1), or six (c-nmd-MutPV-ty1) independent growth experiments. One sample t-test was used to test for statistical differences between complemented cell lines and plus and minus rapalog treatment.

Figure S1: Structure predictions and structure homology search of candidate proteins

- 788 **(A)** AlphaFold structure predictions of the six selected orphan transporters visualized in PyMol.
- 789 **(B)** Results from protein structure comparison server Dali using the AlphaFold-generated PDB
- 790 files of the selected transporters as input structure. Shown are the top five non-redundant hits
- with Z score (significance estimate), msd (difference between the root-mean-square-deviation
- 792 (rmsd) value associated with a protein structure pair and the rmsd value that would have been
- observed in the case that the two structures had the same crystallographic resolution), lali
- 794 (number of aligned positions), nres (number of residues in the matched structure) and %id (the
- 795 percentage sequence identity in the match).

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Figure S2: Validation of generated transgenic cell lines by PCR and Western blot.

- 798 (A) Confirmatory PCR of unmodified wildtype (WT) and transgenic knock-in (KI) cell lines
- 799 (PF3D7 0523800-GFP-glmS (*Pf*FVRT1), PF3D7 0609100-GFP-glmS (*Pf*ZIP1),
- 800 PF3D7 0715900-GFP-glmS (*Pf*CDF), PF3D7 0716900-GFP-glmS (*Pf*DMT2),
- 801 PF3D7_1440800-GFP-glmS (PfMFS6) and PF3D7_1135300-GFP-glmS (PfPMRT1)) to check
- 802 for genomic integration at the 3'- and 5'-end of the locus. Position of the primer used are
- 803 indicated with numbered arrows in Figure 1A. (B) Western Blot analysis of wildtype (3D7) and

knock-in (KI) cell lines using mouse anti-GFP to detect the tagged full-length protein (upper panel) and rabbit anti-aldolase to control for equal loading (lower panel). Protein size is indicated in kDa. Expected molecular weight for GFP fusion proteins: *Pf*FVRT1 (107.5 kDa), *Pf*ZIP1 (69.0 kDa), *Pf*DMT2 (66.4 kDa), *Pf*MFS6 (98.8 kDa), *Pf*PMRT1 (77.5 kDa), *Pf*CDF (91.6 kDa) (C) Localization of *Pf*CDF-3xHA by IFA in ring, trophozoite and schizont parasites. Nuclei were stained with Hoechst. Diagnostic PCR of unmodified wildtype (WT) and transgenic knock-in (KI) cell line. (D) Localization of *Pf*PMRT1_2xFKBP-GFP across the IDC. Nuclei were stained with DAPI. Scale bar, 2 μm. Diagnostic PCR of unmodified wildtype (WT) and transgenic knock-in (KI) cell line. (E) Localization of *Pf*ZIP1-GFP in merozoites. Nuclei were stained with DAPI. Scale bar, 2 μm.

Figure S3: Targeted gene disruption (TGD) of PfZIP1 and PfCDF.

A) Schematic representation of TGD strategy using the selection-linked integration system (SLI). pink, human dihydrofolate dehydrogenase (hDHFR); grey, homology region (HR); green, green fluorescence protein (GFP) tag; dark grey, T2A skip peptide; blue, neomycin resistance cassette. Stars indicate stop codons, and arrows depict primers (P1 to P4) used for the integration check PCR. (B) Localization of *Pt*ZIP1-TGD-GFP in ring, trophozoite and schizont parasites. Nuclei were stained with Hoechst-33342. Scale bar, 2 μm. Confirmatory PCR of unmodified wildtype (WT) and transgenic targeted gene disruption (TGD) cell line. Growth curves of PfZIP1-TGD vs. 3D7 parasites monitored over five days by FACS. Three independent growth experiments were performed and a summary is shown as percentage of growth compared to 3D7 parasites. (C) Localization of *Pt*CDF-TGD in ring, trophozoite and schizont parasites. Nuclei were stained with DAPI. Confirmatory PCR of unmodified wildtype (WT) and transgenic targeted gene disruption (TGD) cell line. Scale bar, 1 μm.

Figure S4: Conditional knockdown via glmS system.

- Live cell microscopy of (A) PfFVRT1-GFP-glmS, (B) PfCDF-GFP-glmS, (C) PfZIP1-GFP-glmS
- 831 (D) PfDMT2-GFP-glmS, (E) PfMFS6-GFP-glmS and (F) PfPMRT1-GFP-glmS parasites 40
- 832 hours after treatment without (control) or with 2.5 mM Glucosamine. Nuclei were stained with
- 833 Hoechst-33342. Scale bar, 2 μm. (G) Individual growth curves of the growth assays shown in
- Figure 2D.

Figure S5: Conditional knockout of PfPMRT1 via DiCre-based system

- 837 Replicates of growth curves of condΔPMRT1, c-nmd3PfPMRT1-ty1 and c-sf3a2PfPMRT1-ty1
- 838 parasites +/- rapalog monitored over five days by FACS shown in Figure 3.

Figure S6: Conditional knockout of PfPMRT1

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- (A) Parasite stage distribution in Giemsa smears displayed as heatmap showing percentage of stages for control, 4 hpi, 20 hpi or 32 hpi rapalog treated 4 hour window synchronized condΔPMRT1parasite cultures over one cycle. (B) Giemsa smears of control and 4 hpi, 20 hpi or 32 hpi rapalog treated parasites at 4, 16, 20, 24, 32, 40 and 48 hpi. Scale bar, 5 μm.
- Figure S7: Conditional knockdown of *Pf*PMRT1 has no effect during gametocyte development.
 - (A) Confirmatory PCR of unmodified wildtype (WT) and transgenic 3D7-iGP-PfPMRT1-GFPglmS to check for genomic integration at the 3'- and 5'-end of the locus. Position of the primer used are indicated with numbered arrows in Figure 1A. (B) Schematic representation of the experimental setup. (C) Live cell microscopy of 3D7-iGP-PfPMRT1-GFP stage I - V gametocytes. Scale bar, 2 µm. (D) Giemsa smears of stage I – V gametocytes cultured either without (control) or with 2.5 mM glucosamine. Scale bar, 5 µm. (E) Quantification of knockdown by measuring mean fluorescence intensity (MFI) density and size (area) of parasites at day 7 and day 12 post induction of gametocytogenesis cultured either without (control) or with 2.5 mM glucosamine. Scale bar, 2 µm. Statistics are displayed as mean +/- SD of four independent experiments and individual data points are displayed as scatterplot color-coded by experiments according to Superplots guidelines(101)(101). P-values displayed were determined with two-tailed unpaired t-test. (F) For each condition gametocytemia at day 10 post gametocyte induction was determined by counting between 1256-2653 (mean 2147) cells per condition in Giemsa-stained thin blood smears. Displayed are means +/- SD of independent growth experiments with the number of experiments (n) indicated. P-values displayed were determined with two-tailed unpaired t-test.
- Figure S8: Individual growth curves of c- ^{nmd3}Pk-ty1 (A) and c- ^{nmd3}Pv-ty1 (B) parasites +/rapalog monitored over two IDCs by FACS shown in Figure 6.
- Figure S9: Individual growth curves of c-^{nmd3}MutIn-ty1 (A) and c-^{nmd3}MutOut-ty1 (B) parasites
 +/- rapalog monitored over two IDCs by FACS shown in Figure 7. (C) IFA of c-^{nmd3}MutPC-ty1
 and c-^{nmd3}MutPV-ty1 parasites to verify correct localization at the parasite plasma membrane.
 Nuclei were stained with Hoechst-33342. Scale bar, 2 µm.

Table S1: Oligonucleotides and plasmids used in this study

References

- 876 1. Martin RE, Henry RI, Abbey JL, Clements JD, Kirk K. 2005. The "permeome" of the
- malaria parasite: an overview of the membrane transport proteins of Plasmodium
- falciparum. Genome Biol 6:R26.
- 879 2. Martin RE. 2020. The transportome of the malaria parasite. Biol Rev Camb Philos Soc
- 880 95:305–332.
- 881 3. Gardner MJ, Hall N, Fung E, White O, Berriman M, Hyman RW, Carlton JM, Pain A,
- Nelson KE, Bowman S, Paulsen IT, James K, Eisen JA, Rutherford K, Salzberg SL,
- Craig A, Kyes S, Chan M-S, Nene V, Shallom SJ, Suh B, Peterson J, Angiuoli S, Pertea
- M, Allen J, Selengut J, Haft D, Mather MW, Vaidya AB, Martin DMA, Fairlamb AH,
- Fraunholz MJ, Roos DS, Ralph SA, McFadden GI, Cummings LM, Subramanian GM,
- Mungall C, Venter JC, Carucci DJ, Hoffman SL, Newbold C, Davis RW, Fraser CM,
- Barrell B. 2002. Genome sequence of the human malaria parasite Plasmodium
- 888 falciparum. Nature 419:498–511.
- 889 4. Martin RE, Ginsburg H, Kirk K. 2009. Membrane transport proteins of the malaria
- parasite. Mol Microbiol. John Wiley & Sons, Ltd (10.1111).
- 891 5. Weiner J, Kooij TWA. 2016. Phylogenetic profiles of all membrane transport proteins of
- the malaria parasite highlight new drug targets. Microb Cell 3:511–521.
- 893 6. Dean P, Major P, Nakjang S, Hirt RP, Martin Embley T. 2014. Transport proteins of
- parasitic protists and their role in nutrient salvage. Front Plant Sci. Frontiers Research
- 895 Foundation.
- 896 7. Foote SJ, Thompson JK, Cowman AF, Kemp DJ. 1989. Amplification of the multidrug
- resistance gene in some chloroquine-resistant isolates of P. falciparum. Cell 57:921-
- 898 930.
- 899 8. Fidock DA, Nomura T, Talley AK, Cooper RA, Dzekunov SM, Ferdig MT, Ursos LMB,
- 900 Bir Singh Sidhu A, Naudé B, Deitsch KW, Su XZ, Wootton JC, Roepe PD, Wellems TE.
- 901 2000. Mutations in the P. falciparum digestive vacuole transmembrane protein PfCRT
- and evidence for their role in chloroquine resistance. Mol Cell 6:861–871.
- 903 9. Veiga MI, Dhingra SK, Henrich PP, Straimer J, Gnädig N, Uhlemann AC, Martin RE,
- 904 Lehane AM, Fidock DA. 2016. Globally prevalent PfMDR1 mutations modulate
- Plasmodium falciparum susceptibility to artemisinin-based combination therapies. Nat
- 906 Commun 7.
- 907 10. Rijpma SR, Van Der Velden M, Bilos A, Jansen RS, Mahakena S, Russel FGM,
- Sauerwein RW, Van De Wetering K, Koenderink JB. 2016. MRP1 mediates folate
- transport and antifolate sensitivity in Plasmodium falciparum. FEBS Lett 590:482–492.
- 910 11. Mesén-Ramírez P, Bergmann B, Elhabiri M, Zhu L, Thien H von, Castro-Peña C,
- Gilberger T-W, Davioud-Charvet E, Bozdech Z, Bachmann A, Spielmann T. 2021. The

- parasitophorous vacuole nutrient pore is critical for drug access in malaria parasites and modulates the fitness cost of artemisinin resistance. Cell Host Microbe 0:283.
- 914 12. Cowman AF, Galatis D, Thompson JK. 1994. Selection for mefloquine resistance in
- 915 Plasmodium falciparum is linked to amplification of the pfmdr1 gene and cross-
- 916 resistance to halofantrine and quinine. Proc Natl Acad Sci U S A 91:1143–1147.
- 917 13. Mok S, Liong KY, Lim EH, Huang X, Zhu L, Preiser PR, Bozdech Z. 2014. Structural
- polymorphism in the promoter of pfmrp2 confers Plasmodium falciparum tolerance to
- 919 quinoline drugs. Mol Microbiol 91:918–934.
- 920 14. Lim MYX, LaMonte G, Lee MCS, Reimer C, Tan BH, Corey V, Tjahjadi BF, Chua A,
- Nachon M, Wintjens R, Gedeck P, Malleret B, Renia L, Bonamy GMC, Ho PCL, Yeung
- 922 BKS, Chow ED, Lim L, Fidock DA, Diagana TT, Winzeler EA, Bifani P. 2016. UDP-
- galactose and acetyl-CoA transporters as Plasmodium multidrug resistance genes. Nat
- 924 Microbiol 1.
- 925 15. Richards SN, Nash MN, Baker ES, Webster MW, Lehane AM, Shafik SH, Martin RE.
- 926 2016. Molecular Mechanisms for Drug Hypersensitivity Induced by the Malaria
- 927 Parasite's Chloroquine Resistance Transporter. PLoS Pathog 12.
- 928 16. Cowell AN, Istvan ES, Lukens AK, Gomez-Lorenzo MG, Vanaerschot M, Sakata-Kato
- T, Flannery EL, Magistrado P, Owen E, Abraham M, La Monte G, Painter HJ, Williams
- 930 RM, Franco V, Linares M, Arriaga I, Bopp S, Corey VC, Gnädig NF, Coburn-Flynn O,
- Reimer C, Gupta P, Murithi JM, Moura PA, Fuchs O, Sasaki E, Kim SW, Teng CH,
- 932 Wang LT, Akidil A, Adjalley S, Willis PA, Siegel D, Tanaseichuk O, Zhong Y, Zhou Y,
- Llinás M, Ottilie S, Gamo FJ, Lee MCS, Goldberg DE, Fidock DA, Wirth DF, Winzeler
- EA. 2018. Mapping the malaria parasite druggable genome by using in vitro evolution
- 935 and chemogenomics. Science (80-) 359:191–199.
- 936 17. Rocamora F, Gupta P, Istvan ES, Luth MR, Carpenter EF, Kümpornsin K, Sasaki E,
- Calla J, Mittal N, Carolino K, Owen E, Llinás M, Ottilie S, Goldberg DE, Lee MCS,
- 938 Winzeler EA. 2021. PfMFR3: A Multidrug-Resistant Modulator in Plasmodium
- 939 falciparum. ACS Infect Dis 7:811–825.
- 940 18. Kirk K. 2004. Channels and transporters as drug targets in the Plasmodium-infected
- 941 erythrocyte. Acta Trop 89:285–298.
- 942 19. Koenderink JB, Kavishe RA, Rijpma SR, Russel FGM. 2010. The ABCs of multidrug
- 943 resistance in malaria. Trends Parasitol. Trends Parasitol.
- 944 20. Murithi JM, Deni I, Pasaje CFA, Okombo J, Bridgford JL, Gnädig NF, Edwards RL, Yeo
- 945 T, Mok S, Burkhard AY, Coburn-Flynn O, Istvan ES, Sakata-Kato T, Gomez-Lorenzo
- 946 MG, Cowell AN, Wicht KJ, Le Manach C, Kalantarov GF, Dey S, Duffey M, Laleu B,
- Lukens AK, Ottilie S, Vanaerschot M, Trakht IN, Gamo F-J, Wirth DF, Goldberg DE,
- 948 Odom John AR, Chibale K, Winzeler EA, Niles JC, Fidock DA. 2021. The Plasmodium

- 949 falciparum ABC transporter ABCl3 confers parasite strain-dependent pleiotropic
- 950 antimalarial drug resistance. Cell Chem Biol 0.
- 951 21. Counihan NA, Modak JK, de Koning-Ward TF. 2021. How Malaria Parasites Acquire
- 952 Nutrients From Their Host. Front Cell Dev Biol 9:582.
- 953 22. Saliba KJ, Horner HA, Kirk K. 1998. Transport and metabolism of the essential vitamin
- pantothenic acid in human erythrocytes infected with the malaria parasite Plasmodium
- 955 falciparum. J Biol Chem 273:10190–10195.
- 956 23. Gulati S, Ekland EH, Ruggles K V., Chan RB, Jayabalasingham B, Zhou B, Mantel PY,
- Lee MCS, Spottiswoode N, Coburn-Flynn O, Hjelmqvist D, Worgall TS, Marti M, Di
- Paolo G, Fidock DA. 2015. Profiling the Essential Nature of Lipid Metabolism in Asexual
- 959 Blood and Gametocyte Stages of Plasmodium falciparum. Cell Host Microbe 18:371–
- 960 381.
- 961 24. Mamoun C Ben, Prigge ST, Vial H. 2010. Targeting the lipid metabolic pathways for the
- treatment of malaria. Drug Dev Res. John Wiley & Sons, Ltd.
- 963 25. Garten M, Beck JR. 2021. Structured to conquer: transport across the Plasmodium
- parasitophorous vacuole. Curr Opin Microbiol. Elsevier Ltd.
- 965 26. Kloehn J, Lacour CE, Soldati-Favre D. 2021. The metabolic pathways and transporters
- of the plastid organelle in Apicomplexa. Curr Opin Microbiol 63:250–258.
- 967 27. Beck JR, Ho CM. 2021. Transport mechanisms at the malaria parasite-host cell
- 968 interface. PLoS Pathog. Public Library of Science.
- 969 28. Kenthirapalan S, Waters AP, Matuschewski K, Kooij TWA. 2016. Functional profiles of
- orphan membrane transporters in the life cycle of the malaria parasite. Nat Commun
- 971 7:10519.
- 972 29. Sayers CP, Mollard V, Buchanan HD, McFadden GI, Goodman CD. 2018. A genetic
- 973 screen in rodent malaria parasites identifies five new apicoplast putative membrane
- 974 transporters, one of which is essential in human malaria parasites. Cell Microbiol
- 975 20:e12789.
- 976 30. Ginsburg H. 2006. Progress in in silico functional genomics: the malaria Metabolic
- 977 Pathways database. Trends Parasitol 22:238–240.
- 978 31. Otto TD, Wilinski D, Assefa S, Keane TM, Sarry LR, Böhme U, Lemieux J, Barrell B,
- Pain A, Berriman M, Newbold C, Llinás M. 2010. New insights into the blood-stage
- 980 transcriptome of Plasmodium falciparum using RNA-Seq. Mol Microbiol 76:12–24.
- 981 32. Wichers JS, Scholz JAM, Strauss J, Witt S, Lill A, Ehnold LI, Neupert N, Liffner B,
- 982 Lühken R, Petter M, Lorenzen S, Wilson DW, Löw C, Lavazec C, Bruchhaus I, Tannich
- 983 E, Gilberger TW, Bachmann A. 2019. Dissecting the gene expression, localization,
- membrane topology, and function of the plasmodium falciparum STEVOR protein
- 985 family. MBio 10:e01500-19.

- 986 33. Treeck M, Sanders JL, Elias JE, Boothroyd JC. 2011. The phosphoproteomes of plasmodium falciparum and toxoplasma gondii reveal unusual adaptations within and
- beyond the parasites' boundaries. Cell Host Microbe 10:410–419.
- 989 34. Pease BN, Huttlin EL, Jedrychowski MP, Talevich E, Harmon J, Dillman T, Kannan N,
- Doerig C, Chakrabarti R, Gygi SP, Chakrabarti D. 2013. Global analysis of protein
- 991 expression and phosphorylation of three stages of plasmodium falciparum
- intraerythrocytic development. J Proteome Res 12:4028–4045.
- 993 35. Jumper J, Evans R, Pritzel A, Green T, Figurnov M, Ronneberger O, Tunyasuvunakool
- K, Bates R, Žídek A, Potapenko A, Bridgland A, Meyer C, Kohl SAA, Ballard AJ, Cowie
- A, Romera-Paredes B, Nikolov S, Jain R, Adler J, Back T, Petersen S, Reiman D,
- Clancy E, Zielinski M, Steinegger M, Pacholska M, Berghammer T, Bodenstein S, Silver
- D, Vinyals O, Senior AW, Kavukcuoglu K, Kohli P, Hassabis D. 2021. Highly accurate
- protein structure prediction with AlphaFold. Nature 596:583–589.
- 999 36. Holm L. 2020. DALI and the persistence of protein shape. Protein Sci 29:128–140.
- 1000 37. Birnbaum J, Flemming S, Reichard N, Soares AB, Mesén-Ramírez P, Jonscher E,
- Bergmann B, Spielmann T. 2017. A genetic system to study Plasmodium falciparum
- protein function. Nat Methods 14:450–456.
- 1003 38. Prommana P, Uthaipibull C, Wongsombat C, Kamchonwongpaisan S, Yuthavong Y,
- Knuepfer E, Holder AA, Shaw PJ. 2013. Inducible Knockdown of Plasmodium Gene
- Expression Using the glmS Ribozyme. PLoS One 8:e73783.
- 1006 39. Tawk L, Chicanne G, Dubremetz J-F, Richard V, Payrastre B, Vial HJ, Roy C, Wengelnik
- 1007 K. 2010. Phosphatidylinositol 3-Phosphate, an Essential Lipid in Plasmodium, Localizes
- to the Food Vacuole Membrane and the Apicoplast. Eukaryot Cell 9:1519–1530.
- 1009 40. Jonscher E, Flemming S, Schmitt M, Sabitzki R, Reichard N, Birnbaum J, Bergmann B,
- Höhn K, Spielmann T. 2019. PfVPS45 Is Required for Host Cell Cytosol Uptake by
- Malaria Blood Stage Parasites. Cell Host Microbe 25:166-173.e5.
- 1012 41. Waller RF, Reed MB, Cowman AF, McFadden GI. 2000. Protein trafficking to the plastid
- of Plasmodium falciparum is via the secretory pathway. EMBO J 19:1794–1802.
- 1014 42. Birnbaum J, Scharf S, Schmidt S, Jonscher E, Hoeijmakers WAM, Flemming S,
- Toenhake CG, Schmitt M, Sabitzki R, Bergmann B, Fröhlke U, Mesén-Ramírez P,
- Blancke Soares A, Herrmann H, Bártfai R, Spielmann T. 2020. A Kelch13-defined
- 1017 endocytosis pathway mediates artemisinin resistance in malaria parasites. Science (80-
- 1018) 367:51–59.
- 1019 43. Inoue T, Heo W Do, Grimley JS, Wandless TJ, Meyer T. 2005. An inducible translocation
- strategy to rapidly activate and inhibit small GTPase signaling pathways. Nat Methods
- 1021 2:415–418.
- 1022 44. Cabrera A. Herrmann S. Warszta D. Santos JM. John Peter AT. Kono M. Debrouver S.

- Jacobs T, Spielmann T, Ungermann C, Soldati-Favre D, Gilberger TW. 2012. Dissection
- of Minimal Sequence Requirements for Rhoptry Membrane Targeting in the Malaria
- 1025 Parasite. Traffic 13:1335–1350.
- 1026 45. Geiger M, Brown C, Wichers JS, Strauss J, Lill A, Thuenauer R, Liffner B, Wilcke L,
- Lemcke S, Heincke D, Pazicky S, Bachmann A, Löw C, Wilson DW, Filarsky M, Burda
- 1028 P-C, Zhang K, Junop M, Gilberger TW. 2020. Structural Insights Into PfARO and
- 1029 Characterization of its Interaction With PfAIP. J Mol Biol 432:878–896.
- 1030 46. Peterson MG, Marshall VM, Smythe JA, Crewther PE, Lew A, Silva A, Anders RF, Kemp
- DJ. 1989. Integral membrane protein located in the apical complex of Plasmodium
- 1032 falciparum. Mol Cell Biol 9:3151–3154.
- 1033 47. Wichers JS, Wunderlich J, Heincke D, Pazicky S, Strauss J, Schmitt M, Kimmel J,
- Wilcke L, Scharf S, von Thien H, Burda P, Spielmann T, Löw C, Filarsky M, Bachmann
- 1035 A, Gilberger TW. 2021. Identification of novel inner membrane complex and apical
- annuli proteins of the malaria parasite Plasmodium falciparum. Cell Microbiol
- 1037 23:e13341.
- 1038 48. Jullien N, Goddard I, Selmi-Ruby S, Fina J-L, Cremer H, Herman J-P. 2007. Conditional
- 1039 Transgenesis Using Dimerizable Cre (DiCre). PLoS One 2:e1355.
- 1040 49. Andenmatten N, Egarter S, Jackson AJ, Jullien N, Herman JP, Meissner M. 2013.
- 1041 Conditional genome engineering in Toxoplasma gondii uncovers alternative invasion
- 1042 mechanisms. Nat Methods 10:125–127.
- 1043 50. Holder AA, Lockyer MJ, Odink KG, Sandhu JS, Riveros-Moreno V, Nicholls SC, Hillman
- 1044 Y, Davey LS, Tizard MLV, Schwarz RT, Freeman RR. 1985. Primary structure of the
- precursor to the three major surface antigens of Plasmodium falciparum merozoites.
- 1046 Nature 317:270–273.
- 1047 51. Mesén-Ramírez P, Bergmann B, Tran TT, Garten M, Stäcker J, Naranjo-Prado I, Höhn
- 1048 K, Zimmerberg J, Spielmann T. 2019. EXP1 is critical for nutrient uptake across the
- parasitophorous vacuole membrane of malaria parasites. PLoS Biol 17:e3000473.
- 1050 52. López-Barragán MJ, Lemieux J, Quiñones M, Williamson KC, Molina-Cruz A, Cui K,
- Barillas-Mury C, Zhao K, Su X zhuan. 2011. Directional gene expression and antisense
- transcripts in sexual and asexual stages of Plasmodium falciparum. BMC Genomics
- 1053 12:587.
- 1054 53. Lasonder E, Rijpma SR, Van Schaijk BCL, Hoeijmakers WAM, Kensche PR, Gresnigt
- 1055 MS, Italiaander A, Vos MW, Woestenenk R, Bousema T, Mair GR, Khan SM, Janse CJ,
- 1056 Bártfai R, Sauerwein RW. 2016. Integrated transcriptomic and proteomic analyses of P.
- Falciparum gametocytes: Molecular insight into sex-specific processes and translational
- repression. Nucleic Acids Res 44:6087–6101.
- 1059 54. Gómez-Díaz E. Yerbanga RS, Lefèvre T, Cohuet A, Rowley MJ, Ouedraogo JB, Corces

- 1060 VG. 2017. Epigenetic regulation of Plasmodium falciparum clonally variant gene expression during development in Anopheles gambiae. Sci Rep 7.
- 1062 55. Boltryk SD, Passecker A, Alder A, Carrington E, van de Vegte-Bolmer M, van Gemert
- 1063 G-J, van der Starre A, Beck H-P, Sauerwein RW, Kooij TWA, Brancucci NMB,
- 1064 Proellochs NI, Gilberger T-W, Voss TS. 2021. CRISPR/Cas9-engineered inducible
- gametocyte producer lines as a valuable tool for Plasmodium falciparum malaria
- transmission research. Nat Commun 12:4806.
- 1067 56. Filarsky M, Fraschka SA, Niederwieser I, Brancucci NMB, Carrington E, Carrió E, Moes
- S, Jenoe P, Bártfai R, Voss TS. 2018. GDV1 induces sexual commitment of malaria
- parasites by antagonizing HP1-dependent gene silencing. Science (80-) 359:1259-
- 1070 1263.
- 1071 57. Ennist NM, Stayrook SE, Dutton PL, Moser CC. 2017. 5VJT: De Novo Photosynthetic
- 1072 Reaction Center Protein Equipped with Heme B and Zn(II) cations. doi:
- 1073 <u>10.2210/pdb5vjt/pdb</u>
- 1074 58. Sonnhammer EL, von Heijne G, Krogh A. 1998. A hidden Markov model for predicting
- transmembrane helices in protein sequences. Proceedings Int Conf Intell Syst Mol Biol
- 1076 6:175–82.
- 1077 59. Käll L, Krogh A, Sonnhammer ELL. 2004. A combined transmembrane topology and
- signal peptide prediction method. J Mol Biol 338:1027–1036.
- 1079 60. Aurrecoechea C, Brestelli J, Brunk BP, Dommer J, Fischer S, Gajria B, Gao X, Gingle
- 1080 A, Grant G, Harb OS, Heiges M, Innamorato F, Iodice J, Kissinger JC, Kraemer E, Li W,
- 1081 Miller JA, Nayak V, Pennington C, Pinney DF, Roos DS, Ross C, Stoeckert CJ,
- Treatman C, Wang H. 2009. PlasmoDB: a functional genomic database for malaria
- parasites. Nucleic Acids Res 37:D539–D543.
- 1084 61. Wunderlich J, Rohrbach P, Dalton JP. 2012. The malaria digestive vacuole. Front Biosci
- 1085 (Schol Ed) 4:1424–48.
- 1086 62. Tabuchi M, Yoshimori T, Yamaguchi K, Yoshida T, Kishi F. 2000. Human
- NRAMP2/DMT1, which mediates iron transport across endosomal membranes, is
- localized to late endosomes and lysosomes in HEp-2 cells. J Biol Chem 275:22220-
- 1089 22228.
- 1090 63. Zhang M, Wang C, Otto TD, Oberstaller J, Liao X, Adapa SR, Udenze K, Bronner IF,
- 1091 Casandra D, Mayho M, Brown J, Li S, Swanson J, Rayner JC, Jiang RHY, Adams JH.
- 1092 2018. Uncovering the essential genes of the human malaria parasite Plasmodium
- falciparum by saturation mutagenesis. Science 360:eaap7847.
- 1094 64. Kennedy K, Cobbold SA, Hanssen E, Birnbaum J, Spillman NJ, McHugh E, Brown H,
- Tilley L, Spielmann T, McConville MJ, Ralph SA. 2019. Delayed death in the malaria
- 1096 parasite Plasmodium falciparum is caused by disruption of prenylation-dependent

- intracellular trafficking. PLOS Biol 17:e3000376.
- 1098 65. Pasaje CFA, Cheung V, Kennedy K, Lim EE, Baell JB, Griffin MDW, Ralph SA. 2016.
- Selective inhibition of apicoplast tryptophanyl-tRNA synthetase causes delayed death
- in Plasmodium falciparum. Sci Rep 6:1–13.
- 1101 66. Yeh E, DeRisi JL. 2011. Chemical rescue of malaria parasites lacking an apicoplast
- defines organelle function in blood-stage plasmodium falciparum. PLoS Biol 9.
- 1103 67. Bushell E, Gomes AR, Sanderson T, Anar B, Girling G, Herd C, Metcalf T, Modrzynska
- 1104 K, Schwach F, Martin RE, Mather MW, McFadden GI, Parts L, Rutledge GG, Vaidya
- AB, Wengelnik K, Rayner JC, Billker O. 2017. Functional Profiling of a Plasmodium
- Genome Reveals an Abundance of Essential Genes. Cell 170:260-272.e8.
- 1107 68. Chasen NM, Stasic AJ, Asady B, Coppens I, Moreno SNJ. 2019. The Vacuolar Zinc
- 1108 Transporter TgZnT Protects Toxoplasma gondii from Zinc Toxicity. mSphere 4:e00086-
- 1109 19.
- 1110 69. Jiang Y, Wei J, Cui H, Liu C, Zhi Y, Jiang Z, Li Z, Li S, Yang Z, Wang X, Qian P, Zhang
- 1111 C, Zhong C, Su X, Yuan J. 2020. An intracellular membrane protein GEP1 regulates
- xanthurenic acid induced gametogenesis of malaria parasites. Nat Commun 11:1764.
- 1113 70. Liu W, Sundararaman SA, Loy DE, Learn GH, Li Y, Plenderleith LJ, Ndjango JBN,
- Speede S, Atencia R, Cox D, Shaw GM, Ayouba A, Peeters M, Rayner JC, Hahn BH,
- Sharp PM. 2016. Multigenomic delineation of Plasmodium species of the Laverania
- subgenus infecting wild-living chimpanzees and gorillas. Genome Biol Evol 8:1929–
- 1117 1939.
- 1118 71. Zanghì G, Vembar SS, Baumgarten S, Ding S, Guizetti J, Bryant JM, Mattei D, Jensen
- ATR, Rénia L, Goh YS, Sauerwein R, Hermsen CC, Franetich J-F, Bordessoulles M,
- Silvie O, Soulard V, Scatton O, Chen P, Mecheri S, Mazier D, Scherf A. 2018. A Specific
- 1121 PfEMP1 Is Expressed in P. falciparum Sporozoites and Plays a Role in Hepatocyte
- 1122 Infection. Cell Rep 22:2951–2963.
- 1123 72. Howick VM, Russell AJC, Andrews T, Heaton H, Reid AJ, Natarajan K, Butungi H,
- Metcalf T, Verzier LH, Rayner JC, Berriman M, Herren JK, Billker O, Hemberg M,
- 1125 Talman AM, Lawniczak MKN. 2019. The malaria cell atlas: Single parasite
- transcriptomes across the complete Plasmodium life cycle. Science (80-) 365.
- 1127 73. Burda P-C, Crosskey T, Lauk K, Zurborg A, Söhnchen C, Liffner B, Wilcke L, Strauss J,
- 1128 Jeffries CM, Svergun DI, Wilson DW, Wilmanns M, Gilberger T-W, Pietsch E, Strauss
- J, Jeffries CM, Svergun DI, Wilson DW, Wilmanns M, Gilberger T-W. 2020. Structure-
- 1130 Based Identification and Functional Characterization of a Lipocalin in the Malaria
- Parasite Plasmodium falciparum. Cell Rep 31:107817.
- 1132 74. Auburn S, Böhme U, Steinbiss S, Trimarsanto H, Hostetler J, Sanders M, Gao Q, Nosten
- 1133 F, Newbold CI, Berriman M, Price RN, Otto TD. 2016. A new Plasmodium vivax

- reference sequence with improved assembly of the subtelomeres reveals an abundance
- of pir genes. Wellcome Open Res 1:4.
- 1136 75. Pain A, Böhme U, Berry AE, Mungall K, Finn RD, Jackson AP, Mourier T, Mistry J,
- Pasini EM, Aslett MA, Balasubrammaniam S, Borgwardt K, Brooks K, Carret C, Carver
- TJ, Cherevach I, Chillingworth T, Clark TG, Galinski MR, Hall N, Harper D, Harris D,
- Hauser H, Ivens A, Janssen CS, Keane T, Larke N, Lapp S, Marti M, Moule S, Meyer
- 1140 IM, Ormond D, Peters N, Sanders M, Sanders S, Sargeant TJ, Simmonds M, Smith F,
- 1141 Squares R, Thurston S, Tivey AR, Walker D, White B, Zuiderwijk E, Churcher C, Quail
- 1142 MA, Cowman AF, Turner CMR, Rajandream MA, Kocken CHM, Thomas AW, Newbold
- 1143 CI, Barrell BG, Berriman M. 2008. The genome of the simian and human malaria
- parasite Plasmodium knowlesi. Nature 455:799–803.
- 1145 76. Liman ER, Tytgat J, Hess P. 1992. Subunit stoichiometry of a mammalian K+ channel
- determined by construction of multimeric cDNAs. Neuron 9:861–871.
- 1147 77. Beck S, Yu-Strzelczyk J, Pauls D, Constantin OM, Gee CE, Ehmann N, Kittel RJ, Nagel
- 1148 G, Gao S. 2018. Synthetic Light-Activated Ion Channels for Optogenetic Activation and
- 1149 Inhibition. Front Neurosci 12:643.
- 1150 78. Trager W, Jensen JB. 1997. Continuous culture of Plasmodium falciparum: its impact
- on malaria research. Int J Parasitol 27:989–1006.
- 1152 79. Lambros C, Vanderberg JP. 1979. Synchronization of Plasmodium falciparum
- 1153 Erythrocytic Stages in Culture. J Parasitol 65:418.
- 1154 80. Ponnudurai T, Lensen AHW, Meis JFGM, Meuwissen JHE. 1986. Synchronization of
- Plasmodium falciparum gametocytes using an automated suspension culture system.
- 1156 Parasitology 93:263–274.
- 1157 81. Rivadeneira E, Wasserman M, Espinal C. 1983. Separation and Concentration of
- 1158 Schizonts of Plasmodium falciparum by Percoll Gradients. J Protozool 30:367–370.
- 1159 82. Moon RW, Hall J, Rangkuti F, Ho YS, Almond N, Mitchell GH, Pain A, Holder AA,
- Blackman MJ. 2013. Adaptation of the genetically tractable malaria pathogen
- 1161 Plasmodium knowlesi to continuous culture in human erythrocytes. Proc Natl Acad Sci
- 1162 110:531–536.
- 1163 83. Ganesan SM, Morrisey JM, Ke H, Painter HJ, Laroiya K, Phillips MA, Rathod PK, Mather
- MW, Vaidya AB. 2011. Yeast dihydroorotate dehydrogenase as a new selectable
- marker for Plasmodium falciparum transfection. Mol Biochem Parasitol 177:29–34.
- 1166 84. Grüring C, Spielmann T. 2012. Imaging of live malaria blood stage parasites. Methods
- 1167 Enzymol 506:81–92.
- 1168 85. Bachmann A, Scholz JAM, Janßen M, Klinkert M-Q, Tannich E, Bruchhaus I, Petter M.
- 2015. A comparative study of the localization and membrane topology of members of
- the RIFIN. STEVOR and PfMC-2TM protein families in Plasmodium falciparum-infected

- 1171 erythrocytes. Malar J 14:274.
- 1172 86. Collins CR, Hackett F, Strath M, Penzo M, Withers-Martinez C, Baker DA, Blackman
- MJ. 2013. Malaria Parasite cGMP-dependent Protein Kinase Regulates Blood Stage
- 1174 Merozoite Secretory Organelle Discharge and Egress. PLoS Pathog 9:e1003344.
- 1175 87. Tonkin CJ, Van Dooren GG, Spurck TP, Struck NS, Good RT, Handman E, Cowman
- AF, McFadden GI. 2004. Localization of organellar proteins in Plasmodium falciparum
- using a novel set of transfection vectors and a new immunofluorescence fixation
- method. Mol Biochem Parasitol 137:13–21.
- 1179 88. Blackman MJ, Whittle H, Holder AA. 1991. Processing of the Plasmodium falciparum
- major merozoite surface protein-1: identification of a 33-kilodalton secondary
- processing product which is shed prior to erythrocyte invasion. Mol Biochem Parasitol
- 1182 49:35–44.
- 1183 89. Schindelin J, Arganda-Carreras I, Frise E, Kaynig V, Longair M, Pietzsch T, Preibisch
- S, Rueden C, Saalfeld S, Schmid B, Tinevez J-Y, White DJ, Hartenstein V, Eliceiri K,
- Tomancak P, Cardona A. 2012. Fiji: an open-source platform for biological-image
- 1186 analysis. Nat Methods 9:676–682.
- 1187 90. Mesén-Ramírez P, Reinsch F, Blancke Soares A, Bergmann B, Ullrich AK, Tenzer S,
- Spielmann T. 2016. Stable Translocation Intermediates Jam Global Protein Export in
- 1189 Plasmodium falciparum Parasites and Link the PTEX Component EXP2 with
- 1190 Translocation Activity. PLoS Pathog 12:e1005618.
- 1191 91. Malleret B, Claser C, Ong ASM, Suwanarusk R, Sriprawat K, Howland SW, Russell B,
- Nosten F, Rénia L. 2011. A rapid and robust tri-color flow cytometry assay for monitoring
- malaria parasite development. Sci Rep 1:118.
- 1194 92. Wichers JS, van Gelder C, Fuchs G, Ruge JM, Pietsch E, Ferreira JL, Safavi S, von
- Thien H, Burda P-C, Mesén-Ramirez P, Spielmann T, Strauss J, Gilberger T-W,
- Bachmann A. 2021. Characterization of Apicomplexan Amino Acid Transporters
- 1197 (ApiATs) in the Malaria Parasite Plasmodium falciparum. mSphere
- 1198 https://doi.org/10.1128/mSphere.00743-21.
- 1199 93. Galen SC, Borner J, Martinsen ES, Schaer J, Austin CC, West CJ, Perkins SL. 2018.
- The polyphyly of Plasmodium: Comprehensive phylogenetic analyses of the malaria
- parasites (Order Haemosporida) reveal widespread taxonomic conflict. R Soc Open Sci
- 1202 5.
- 1203 94. Yu G, Smith DK, Zhu H, Guan Y, Lam TT-Y. 2017. ggtree: an r package for visualization
- and annotation of phylogenetic trees with their covariates and other associated data.
- 1205 Methods Ecol Evol 8:28–36.
- 1206 95. Yu G. 2020. Using ggtree to Visualize Data on Tree-Like Structures. Curr Protoc
- 1207 Bioinforma 69:e96.

- 1208 96. Katoh K, Standley DM. 2013. MAFFT Multiple Sequence Alignment Software Version
 1209 7: Improvements in Performance and Usability. Mol Biol Evol 30:772–780.
- 1210 97. Madeira F, Park YM, Lee J, Buso N, Gur T, Madhusoodanan N, Basutkar P, Tivey ARN,
 1211 Potter SC, Finn RD, Lopez R. 2019. The EMBL-EBI search and sequence analysis tools
- 1212 APIs in 2019. Nucleic Acids Res 47:W636–W641.
- 1213 98. R Core Team. 2017. R: A language and environment for statistical computing. R Found 1214 Stat Comput Vienna, Austria.
- 1215 99. Wickham H. 2016. Ggplot2: elegant graphics for data analysis. Springer.
- 1216 100. Lin Pedersen T. 2020. patchwork, The Composer of Plots.
- 1217 101. Lord SJ, Velle KB, Mullins RD, Fritz-Laylin LK. 2020. SuperPlots: Communicating reproducibility and variability in cell biology. J Cell Biol 219.
- 1219 102. Tibúrcio M, Hitz E, Niederwieser I, Kelly G, Davies H, Doerig C, Billker O, Voss TS, 1220 Treeck M. 2021. A 39-Amino-Acid C-Terminal Truncation of GDV1 Disrupts Sexual 1221 Commitment in Plasmodium falciparum. mSphere 6.
- 1222 103. Eksi S, Morahan BJ, Haile Y, Furuya T, Jiang H, Ali O, Xu H, Kiattibutr K, Suri A, Czesny 1223 B, Adeyemo A, Myers TG, Sattabongkot J, Su X zhuan, Williamson KC. 2012. 1224 Plasmodium falciparum Gametocyte Development 1 (Pfgdv1) and Gametocytogenesis 1225 Early Gene Identification and Commitment to Sexual Development. PLoS Pathog 8.
- 1226 104. Omasits U, Ahrens CH, Müller S, Wollscheid B. 2014. Protter: Interactive protein feature 1227 visualization and integration with experimental proteomic data. Bioinformatics 30:884– 1228 886.
- 1230 **Table S1:** Oligonucleotides (A) and plasmids (B) used in this study.

1231 **A**

inter nal #	Target or Primer name	Sequence	Purpose
284	PF3D7_071 6900 rv	GGGacgcgtTAACATCAATTTTGCTTTTTTGGG	
285	PF3D7_071 6900 fw	GGGgcggccgctaaGGGTGTGGTAATTCAAGTACG	
280	PF3D7_052 3800 rv	GGGacgcgtATTTCGTTGAATATAATTTTTTTAATTG	
281	PF3D7_052 3800 fw	GGGgcggccgctaaGTTTTCTTGTTGGTATTTTAGCTG	
282	PF3D7_060 9100 rv	GGGacgcgtATGATTATGACCATGATCATGATC	
283	PF3D7_060 9100 fw	GGGgcggccgctaaGGATAGCAGGTGTTACGGTTTCTTTATC	cloning GFP-glmS
288	PF3D7_144 0800 rv	GGGacgcgtATTAGTAATAGAATTTTTCATCTTG	
289	PF3D7_144 0800 fw	GGGgcggccgctaaCATTTGCTTCAAATTTGATGAG	
290	Pf3D7_1135 300 rv	GGGacgcgtAGAAGTTTTTGGGGCATATTTCTTTG	
338	Pf3D7_0715 900 (11)	GGGgcggccgctaaCTGGAAATAAAATAGATGGAACGTCTTG	
339	Pf3D7_0715 900 (11)	GGGacgcgtAGTATCCCCTTTCAATGTGGAAC	

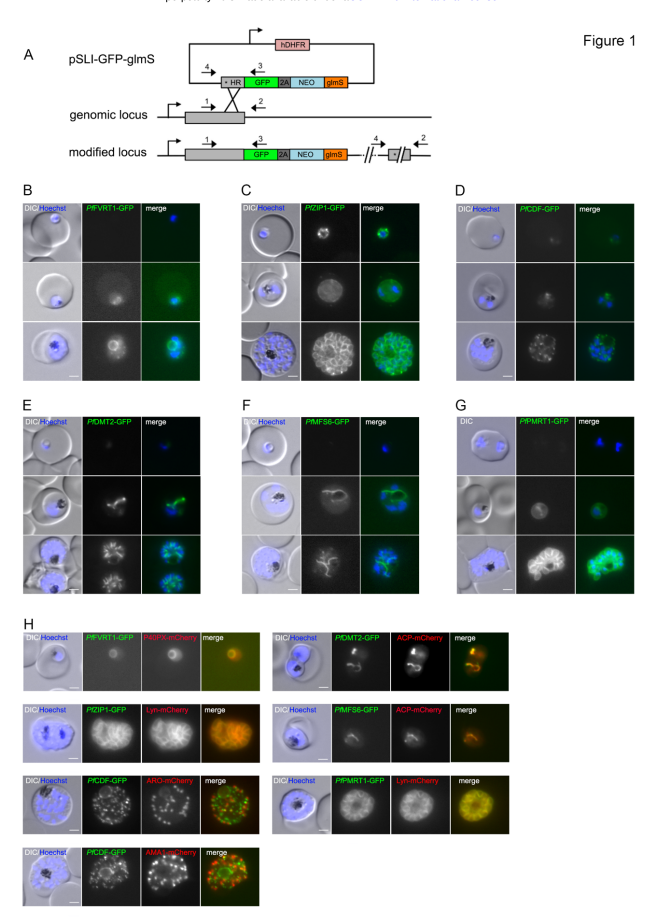
	Pf3D7_1135		
291	300 fw	GGGgcggccgctaaGTTATATAAAAGGATGATTGG	
315	PF3D7_052 3800 (1) TGD fw	GCGGCCGCTAAAGGAGCACTAAAGGCCAAAGGAAGT	
316	PF3D7_052 3800 (1) TGD rv	acgcgtAATAATGTCATCCTTTTCATTAATATT	
317	PF3D7_060 9100 (2) TGD fw	GCGGCCGCTAAGATTTGTTATTTGCAAAAATAATTTGTATT	
318	PF3D7_060 9100 (2) TGD rv	acgcgtATGTctataaaaaaaataaatcacatac	
319	PF3D7_071 6900 (3) TGD fw	GCGGCCGCTAAAAAAATGAAATTATATTTTGTACGACATTC	
320	PF3D7_071 6900 (3) TGD rv	acgcgtGGCAATACCAAAGGTAATTAATAAAATTCC	cloning TGD
323	PF3D7_144 0800 (5) TGD fw	GCGGCCGCTAAACAATGTTTAAGGATGATGAAAATAATTTT	Cidning 1GD
324	PF3D7_144 0800 (5) TGD rv	acgcgtTGTATTAACAACAACTATAAATACAGAACT	
	Pf3D7_0715 900 SLI-Tdg F	GGTGCGGCCGCGTGGATAAGATGTCGCGTTTG	
	Pf3D7_0715 900 SLI-Tgd R	GGTACGCGTGTCCTACATTTTTATCATCTTCCTC	
	PF3D7_113 5300 – fw Notl TGD	CTCGgcggccgctaaAAAAGTATGATCTCTGGAATATCCAG	
	PF3D7_113 5300 – rv Mlul TGD	TCCTacgcgtATTTATAAAGAGATCTGTTTTATTATC	
	PF3D7_113 5300 fw NotI loxP	CTCGGCGGCCCCGGTAAATCTCTGGAATATCCAGCAAAATTG	
	PF3D7_113 5300 rv Spel loxP:	TCCTACTAGTATTTATAAAGAGATCTGTTTTATTATC	cloning 1135300
	PF3D7_113 5300 fw AvrII loxP:	CTCGCCTAGGATGAAGTCAATGATAAGCGGTATTAG	loxp
	PF3D7_113 5300 rv Xmal loxP:	TCCTCCCGGGGCTTGTCTTAGGTGCGTACTTTTTAC	
294	PF3D7_060 9100 int_fw	GGTAGTAACAACCTTTGGTTGTTTTATTCC	
295	PF3D7_060 9100 int_rv	tatgtggaaggttaattaaatggacaaggg	
296	PF3D7_071 6900 int_fw	GAAATTATATTTTGTACGACATTCATACTA	
298	PF3D7_144 0800 int_fw	GCACAGAACATTTAAGAAGCAATGATTTTA	7
299	PF3D7_144 0800 int_rv	caacttgactagccaaaatgttggttctgg	1
300	PF3D7_113 5300 int_fw	GATCTCTGGAATATCCAGCAAAATTGTTG	1
302	PF3D7_071 6900 int_rv	atatttatttatttgaaccgataagctag	integration check PCR
305	PF3D7_113 5300 int_rv	gccatatatatatacatattaaatataaagac	5551(7.51)
308	PF3D7_052 3800 int_fw	GGCCAAAGGAAGTTGGCTAACGGGGTGGTAG	7
309	PF3D7_052 3800 int_rv	gtgttcattcatttaccttttgaatgg	
369	TGD (PF3D7_060 9100) int	atatatgttttaaagcttcaaaatg	
370	check fw TGD (PF3D7_060	gaggagaagcaaaaacgaaagttaac	

			•
	9100) int check rv		
	TGD		1
	(Pf3D7 071		
392	5900) int	CCCACCGAAATGAACTCTTCGTTGC	
	check fw		
	TGD		
	(Pf3D7_071		
393	5900) int	CTTTACCTTTAGAGGAGGAATTATTAG	
	check rv		
	SLI TGD it		
394	fw Adelaide	gtggaattgtgagcggataac	
	Pf3D7 1135		
327	300 compl	ctcgagATGAAGTCAATGATAAGCGG	
	Pf3D7 1135		
328	300 compl	cctaggGCTTGTCTTAGGTGCGTACT	
	PKNH_0933		cloning
553	400 fw 553	GGGctcgagATGAAGGGAACGTACGTAG	complement
	PKNH_0933		ation
554	400 rv 554	GGGcctaggCACCGCCTTCGAGGCGTAC	constrcuts
	PVP01 093		
555	6100 fw 555	GGGctcgagATGAAGGGAACGTACGTCG	
	PVP01 093		
556	6100 rv 556	GGGcctaggCACCGCCTTCGAGGCGTAC	
	Pf3D7_1135		
p37	300 fw 5'	GATTTTGATATATGATTATAGGATAG	
μσ.	(TKo)		excision
	Neo 40 rv	CGAATAGCCTCTCCACCCAAG	PCR primer
	Pf3D7_1135		cloning
	300 fw Notl	CTCGgcggccgctaaTTATTATAAATCATATAATAAAATAAATG	1135300-
	Pf3D7_1135		2xFKBP-
	300 rv Avrll	TCCTcctaggAGAAGTTTTTGGGGCATATTTCTTTG	GFP
	Pf3D7_0715		cloning
443	900 (11)	GggtaccAGTATCCCCTTTCAATGTGGAAC	0715900-
773	Kpnl	oggiace Ao TATOOOOTT TO AATO TOO AAO	3xHA
	PF3D		OAL II C
	MutOut F	GGAAATGCGGCCGCGTTACTTAAGGACGACAAAATATTCA	
	PF3D	TAAGTAACGCGGCCGCATTTCCCTTTATATTGTATTGTGCTATTGCGTTCTGTG	1
	MutOut R	AAAGACTAACGTACTTGTTTTG	cloning of
	PF3D MutIn		MutIn and
	qcF	tCAAGCAGCctttGCTATGGCAttccttattttctacataaagatggga	MutOut
	PF3D MutIn		1
	gcR	CATAGCaaagGCTGCTTGaatcGCgaagtaaagaacaacgaactcta	
	Pf3D7 1135		
	300 BamH I	cgggatccaccatgaagtcaatgataagcggtat	cloning
	fw	ogggaroodoodigaagroadigalaagoggiar	1135300-
	Pf3D7_1135		YFP
	300 Xhol rv	CCGctcgagaccgcttgtcttaggtgcgt	
	2007		l

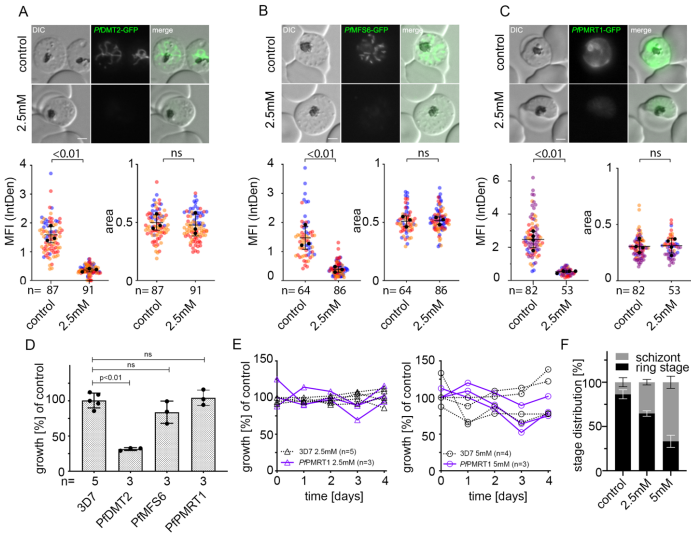
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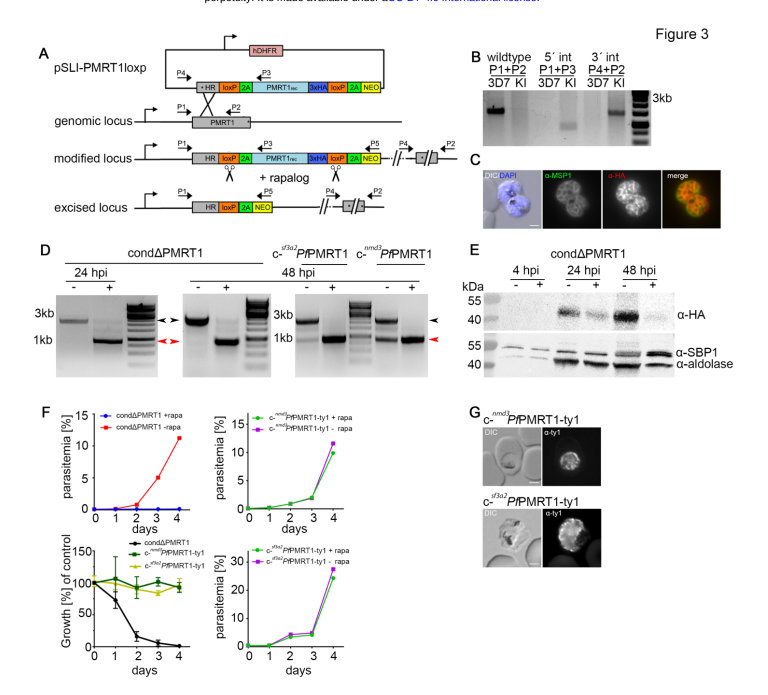
pSLI- <i>Pf</i> FVRT1-GFP-glmS	this study
pSLI- <i>Pf</i> ZIP1-GFP-glmS	this study
pSLI- <i>Pf</i> CDF-GFP-glmS	this study
pSLI- <i>Pf</i> DMT2-GFP-glmS	this study
pSLI- <i>Pf</i> MFS6-GFP-glmS	this study
pSLI- <i>Pf</i> PMRT1-GFP-glmS	this study
pSLI- <i>Pf</i> FVRT1-GFP-TGD	this study
pSLI- <i>Pf</i> ZIP1-GFP-TGD	this study
pSLI- <i>Pf</i> CDF-GFP-TGD	this study
pSLI- <i>Pf</i> DMT2-GFP-TGD	this study
pSLI- <i>Pf</i> MFS6-GFP-TGD	this study

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Jonscher et al. 2018 (40)
Birnbaum et al. 2017 (37)
Birnbaum et al. 2020 (42)
Cabrera et al. 2012 (44)
Wichers et al. 2021 (47)
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Birnbaum et al. 2017 (37)
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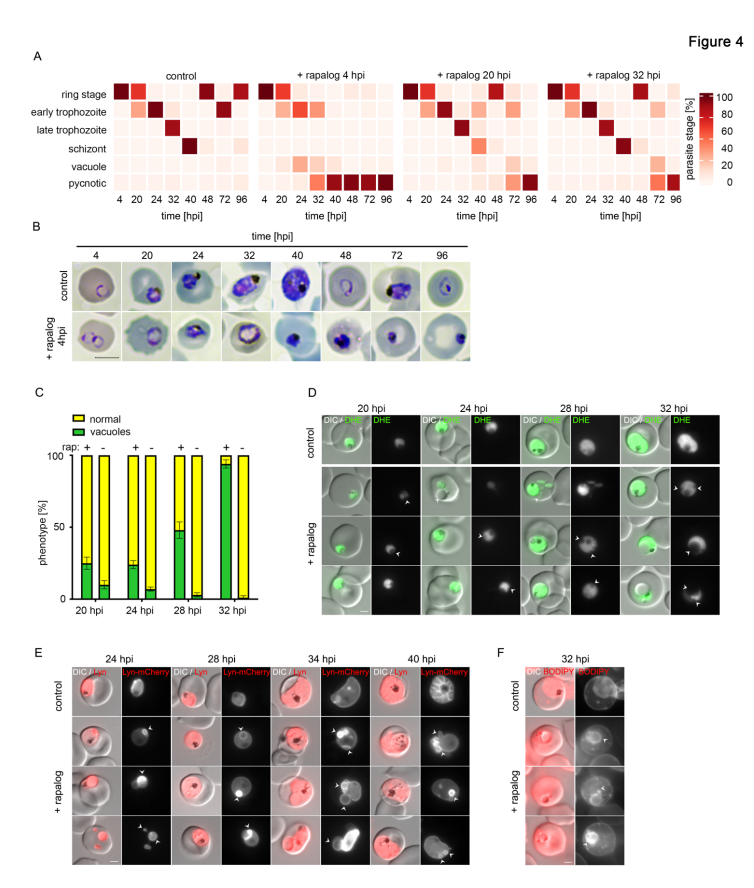


Figure 5

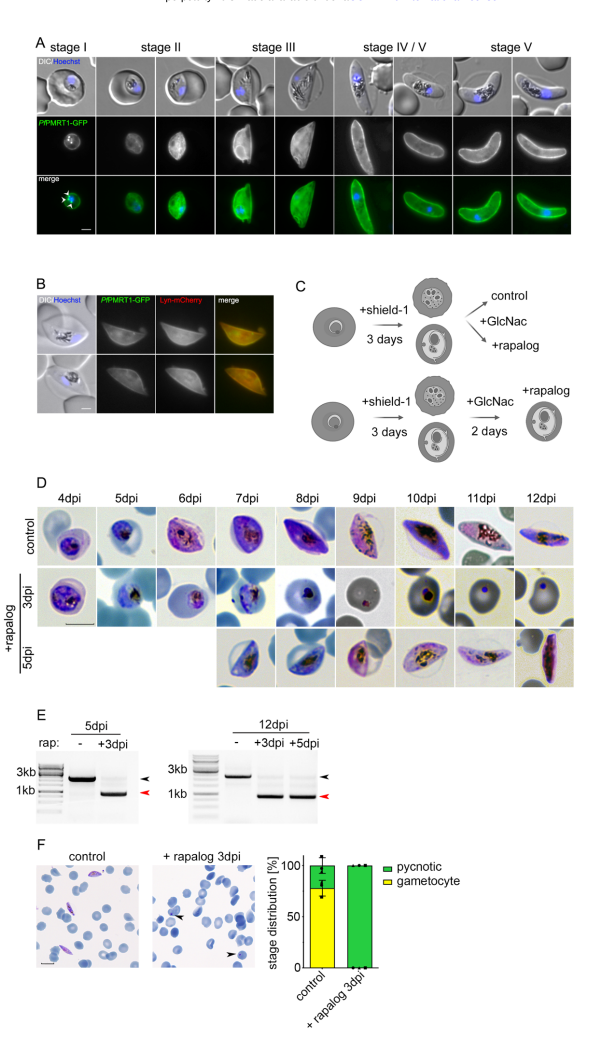
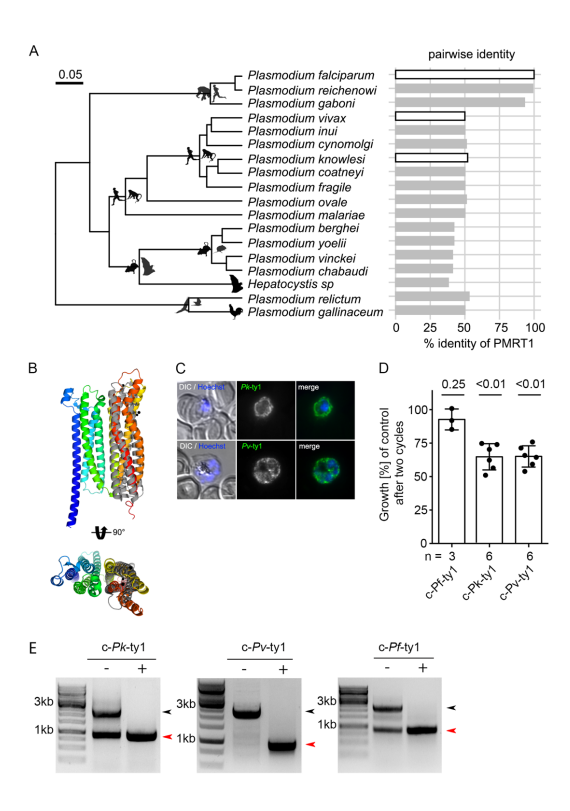


Figure 6



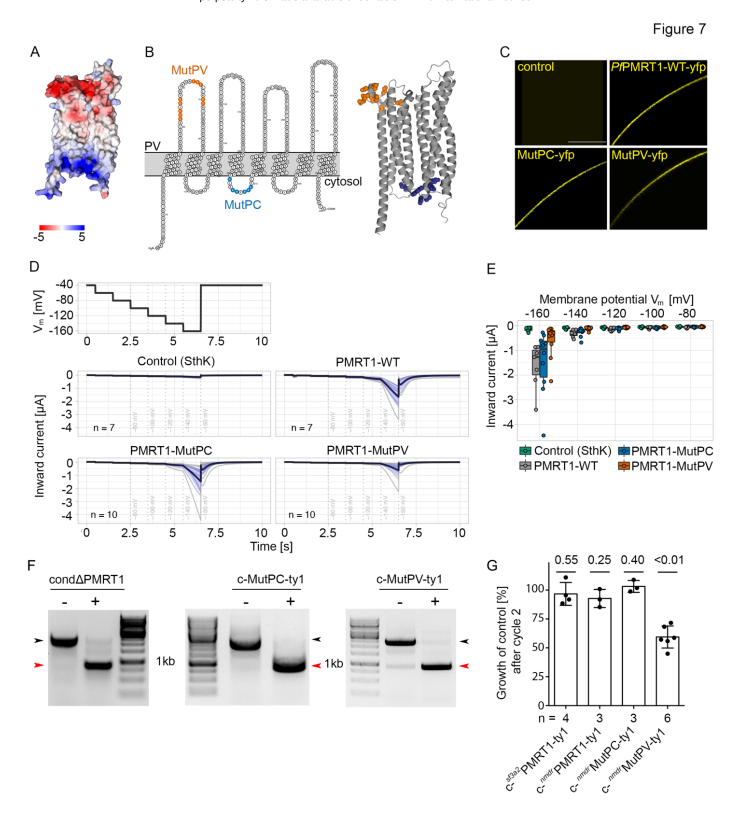
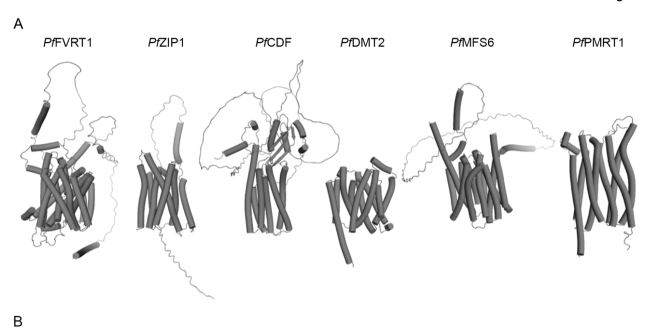


Figure S1



	hit no	. PDB-chain	Z	msd	lali	nres	%id	PDB description on www.rcsb.org
	. 1	6c3i-A	45.9	2.0	379	386	21	Crystal structure of the Deinococcus radiodurans Nramp/MntH divalent transition metal transporter G45R mutant in an inward occluded state
D/EVDT4	, 2	5m94-A	39.9	2.6	377	401	24	Crystal structure of Staphylococcus capitis divalent metal ion transporter (DMT) in complex with nanobody
ž	3	6tl2-A	38.9	2.8	440	494	21	Crystal structure of Eremococcus coleocola manganese transporter in complex with an aromatic bis-isothiourea substituted compound
20	4	6s3k-A	25.1	3.8	357	573	8	KimA from Bacillus subtilis in inward-facing, occluded state (APC Family Permease)
	5	6cse-C	24.7	3.4	344	428	13	Crystal structure of sodium/alanine symporter AgcS with L-alanine bound
	1	6pgi-A	16.1	3.6	206	230	15	Asymmetric functions of a binuclear metal cluster within the transport pathway of the ZIP transition metal transporters
2	2	7c76-B	9.5	3.5	150	483	5	Cryo-EM structure of human TLR3 in complex with UNC93B1
PfZIP1	3	6m2l-A	9.0	3.6	154	447	6	Crystal structure of Plasmodium falciparum hexose transporter PfHT1 bound with C3361
q	4	4gbz-A	8.9	3.6	159	475	7	The structure of the MFS (major facilitator superfamily) proton:xylose symporter XylE bound to D-glucose
	5	6ob6-B	8.8	3.6	155	357	12	Human equilibrative nucleoside transporter-1, S-(4-nitrobenzyl)-6-thioinosine bound, merohedrally twinned
	1	6xpd-B	27.1	2.0	300	303	34	Cryo-EM structure of human ZnT8 double mutant - D110N and D224N, determined in outward-facing conformation
4	2	3h90-C	18.4	2.8	268		15	
PFCDF	3	6vd9-A		1.4	71	73	32	
	- 4	6vd8-B		1.3	76	78	37	Metal-bound C-terminal domain of CzcD transporter from Pseudomonas aeruginosa
	5	3w62-A		2.0	73	79	14	MamM-CTD E289A
	. 1	6ukj-A		3.4	291	350		
È	2	5ogk-E		3.6	285			Crystal structure of a nucleotide sugar transporter with bound nucleotide sugar
D#DMT2	3	6xbo-A						X-ray crystal structure of the mouse CMP-Sialic acid transporter in complex with 5-methyl CMP
à	. 4	5i20-D		3.1			9	Crystal structure of protein (drug metabolite transporter YddG)
	5	5y78-A		4.0	288			Crystal structure of the triose-phosphate/phosphate translocator in complex with inorganic phosphate
PfPMRT1 PfMFS6	, 1	6t1z-A		3.0		393		LmrP from L. lactis, in an outward-open conformation, bound to Hoechst 33342
	2	6s4m-A		2.3		426	17	Crystal structure of the human organic anion transporter MFSD10 (TETRAN)
	3	3wdo-A		3.1	364	453		Structure of E. coli YajR transporter
		3o7p-A		3.0		410	11	Crystal structure of the E.coli Fucose:proton symporter, FucP (N162A)
	5	7ckr-A		2.7		382	7	Cryo-EM structure of the human MCT1/Basigin-2 complex in the presence of anti-cancer drug candidate BAY-8002 in the outward-open conformation
	. 1	6z0c-A		3.3			4	Structure of in silico modelled artificial Maquette-3 protein
	2	5vjt-A		3.3			3	De Novo Photosynthetic Reaction Center Protein Equipped with Heme B and Zn(II) cations
2	3	7p5c-B		3.2			4	Cryo-EM structure of human TTYH3 in Ca2+ and GDN (Protein Tweety Homolog 3)
Ā	: 4	6njm-E		2.0	116		0	Architecture and subunit arrangement of native AMPA receptors (Glutamate-gated AMPA receptor)
	5	6y90-A	12.0	3.5	140	172	8	Structure of full-length CD20 in complex with Rituximab Fab (B-Lymphocyte Antigen CD20)

Dali Z score: significance estimate

msd: difference between the msd value associated with a protein structure pair and the msd value that would have been observed in the case that the two structures had the same crystallographic resolution lali: number of aligned positions

nres: number of residues in the matched structure

 $\%\mbox{id}$: the percentage sequence identity in the match

rmsd: root-mean-square-deviation over aligned $\alpha\text{-carbon}$ position

Figure S2

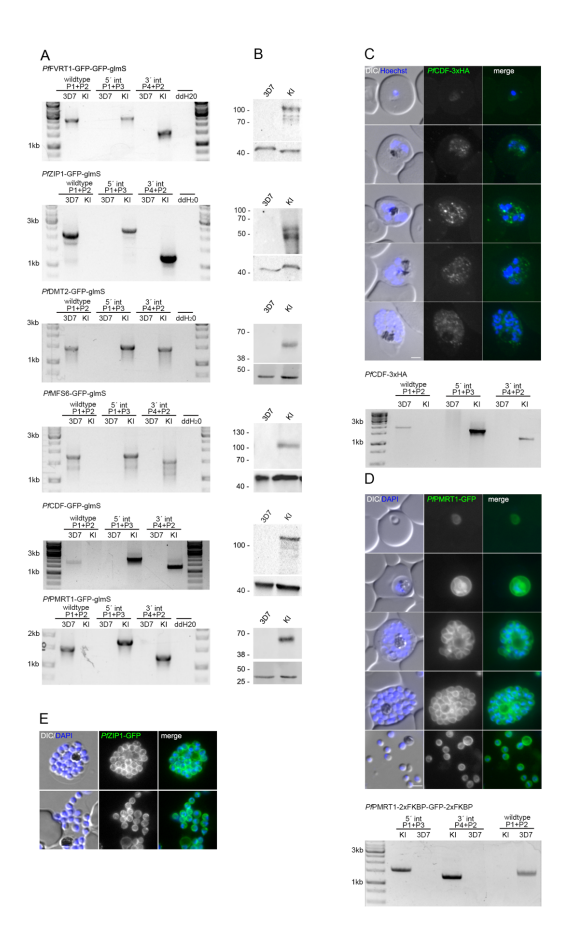
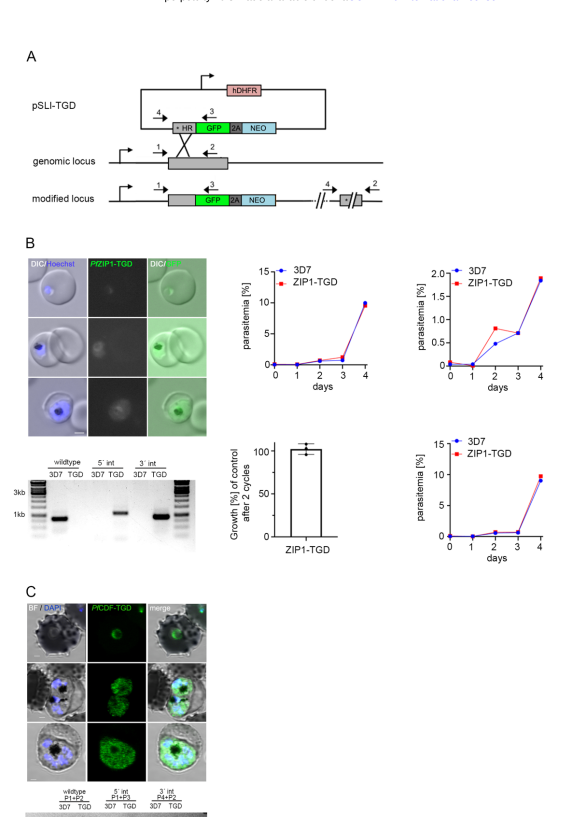


Figure S3



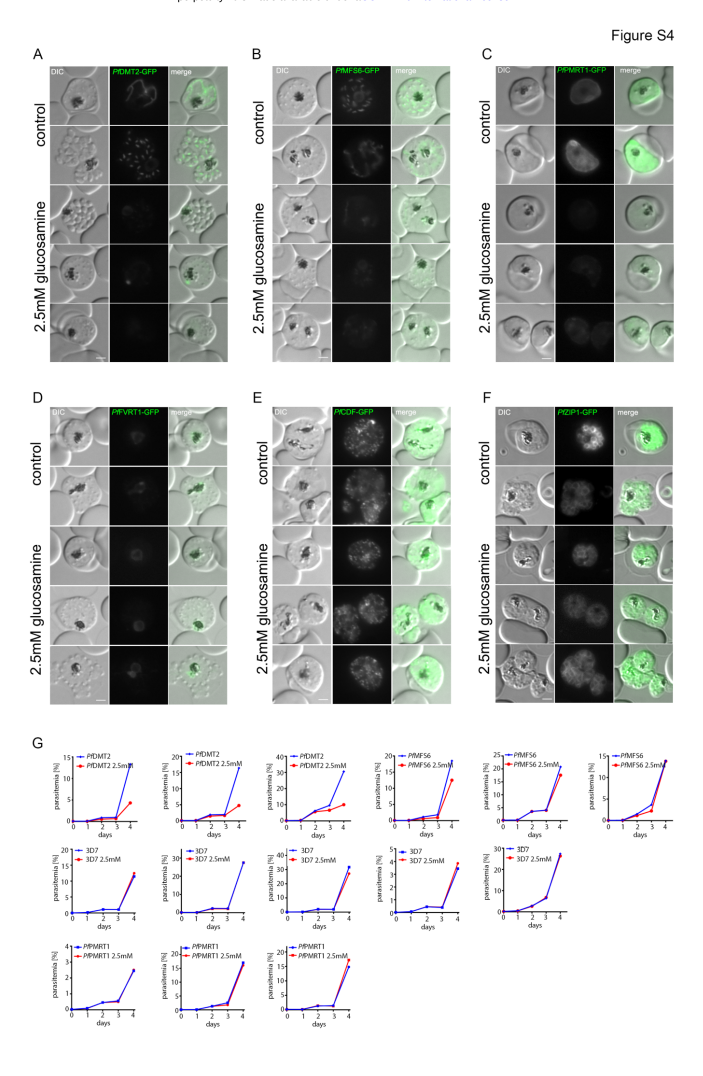
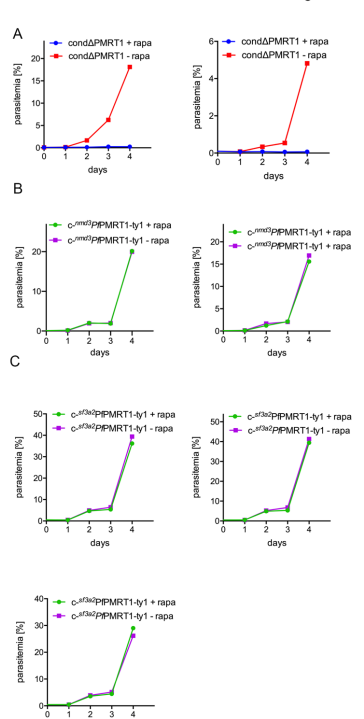
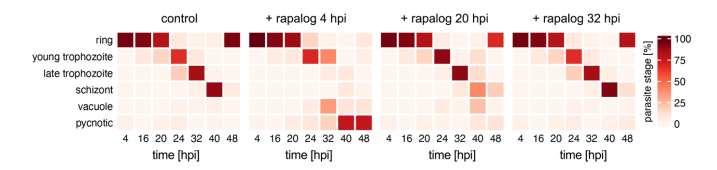


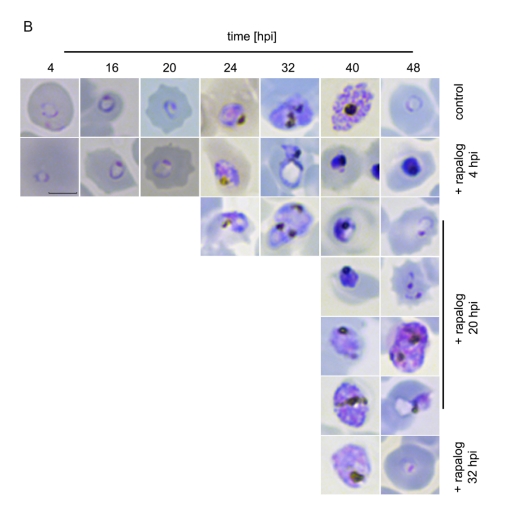
Figure S5



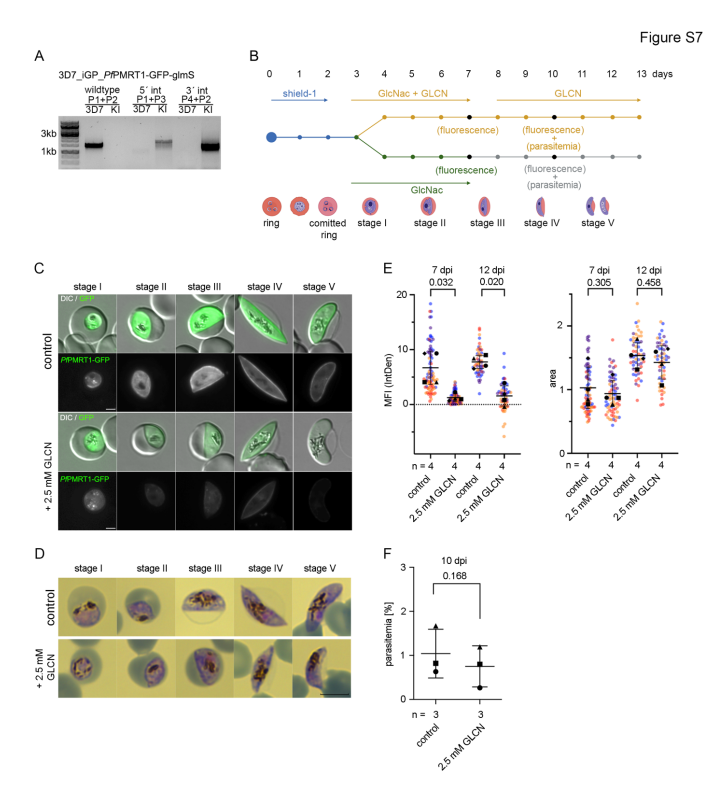
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Figure S6





Α



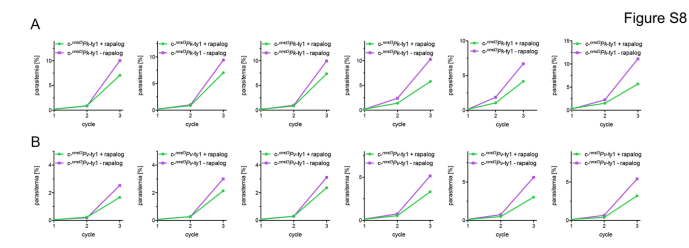
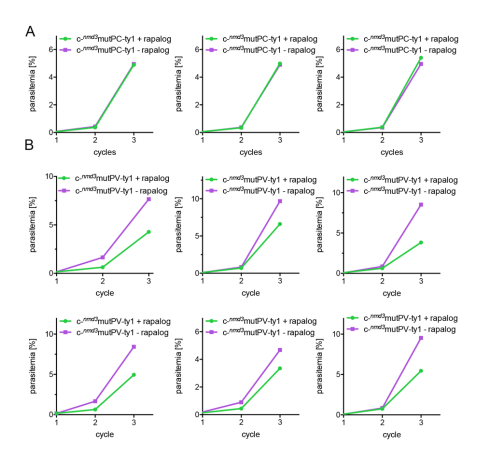
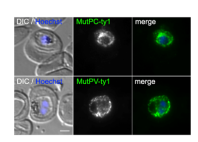


Figure S9





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