The Role of Palmitoylation for Protein Recruitment to the Inner Membrane Complex of the Malaria Parasite*

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Background: Recruitment of peripheral proteins to the inner membrane complex (IMC) of the malaria parasite can be mediated by N-terminal acylation.

Results: Characterization of substrate determinants and identification of an IMC-localized palmitoyl acyltransferase PfDHHC1.

Conclusion: Residues close to palmitoylation sites interfere with specific IMC recruitment. PfDHHC1 represents an apicomplexan-specific PAT.

Significance: Dissection of palmitoylation for protein recruitment to the inner membrane complex in P. falciparum.

To survive and persist within its human host, the malaria parasite Plasmodium falciparum utilizes a battery of lineage-specific innovations to invade and multiply in human erythrocytes. With central roles in invasion and cytokinesis, the inner membrane complex, a Golgi-derived double membrane structure underlying the plasma membrane of the parasite, represents a unique and unifying structure characteristic to all organisms belonging to a large phylogenetic group called Alveolata. More than 30 structurally and phylogenetically distinct proteins are embedded in the IMC, where a portion of these proteins display N-terminal acylation motifs. Although N-terminal myristoylation is catalyzed co-translationally within the cytoplasm of the parasite, palmitoylation takes place at membranes and is mediated by palmitoyl acyltransferases (PATs). Here, we identify a PAT (PfDHHC1) that is exclusively localized to the IMC. Systematic phylogenetic analysis of the alveolate PAT family reveals PfDHHC1 to be a member of a highly conserved, apicomplexan-specific clade of PATs. We show that during schizogony this enzyme has an identical distribution like two dual-acylated, IMC-localized proteins (PfISP1 and PfISP3). We used these proteins to probe into specific sequence requirements for IMC-specific membrane recruitment and their interaction with differentially localized PATs of the parasite.

Malaria tropica, the most severe form of human malaria, is caused by the apicomplexan parasite Plasmodium falciparum and represents one of the major human health problems in endemic countries. Half of the world’s population is at risk, and approximately one million people die annually, the majority being children under the age of 5 years in sub-Saharan Africa (1). From an anthropocentric point of view, the most important part of the parasite’s complex life cycle is the asexual and intraerythrocytic developmental cycle, which causes all symptoms associated with malaria. Invasion of human red blood cells by merozoites initiates this intracellular replication cycle. This process is coordinated and powered by the parasite and relies on a sophisticated protein network with various levels of hierarchy and control points (2). Of critical importance to host cell invasion are rhoptries and micronemes, the secretory organelles at the apical pole of the parasite, as well as the inner membrane complex (IMC), a membranous structure consisting of two layers located on the cell’s sub-surface, underneath the plasma membrane (3). Although the secretory organelles are characteristic of all organisms belonging to the apicomplexan phylum, the IMC is a morphological trait of a large phylogenetic group called Alveolata (4). The Alveolata comprise the Dinoflagellata (typically marine flagellates), Ciliata (like Paramecium spp.), and the parasitic Apicomplexa, including the genera Theileria, Eimeria, Toxoplasma, and Plasmodium. For dinoflagellates and ciliates, largely aquatic groups, the IMC appears to

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have a predominantly structural role in cell architecture. However, for apicoplasts, an additional function is its role in cell motility and cytokinesis (5).

To date, over 25 IMC resident proteins have been identified (3, 5). Phylogenetic analysis has demonstrated that in addition to a common core set of conserved proteins, the IMC includes many lineage-specific proteins, reflecting additional adaptations to specialized requirements of the compartment within the respective phylogenetic group (6). Functional specialization of IMC proteins is exemplified by a group of well-characterized proteins that form the so-called glideosome, the motor complex that drives the locomotion of all motile parasite stages (7–9). In addition to IMC resident proteins, studies have recently revealed an additional IMC-associated protein family termed IMC sub-compartment proteins (ISPs) (10). Originally discovered in P. falciparum (11), ISPs have also been the focus of detailed investigations in Toxoplasma gondii (11–13). Although the precise functions of ISPs are still unknown, knock-out studies in T. gondii and Plasmodium berghei suggest that they play a role in cell polarity and daughter cell formation and are not connected to cell motility (11, 12, 14).

PFISP are relatively small (144 and 148 amino acids for PFISP1/PF10_0107 and PFISP3/PF14_0578, respectively) and display N-terminal myristoylation and palmitoyl motif. It was previously shown that, in contrast to the glideosome-associated protein 45 (GAP45), the N-terminal domain of the ISPs is sufficient for IMC membrane recruitment (10, 15). Although myristoylation of proteins is catalyzed by a cytosolic N-myristoyltransferase (PF14_0127; see Ref. 16), palmitoylation is catalyzed by palmitoyl acyltransferases (PATs) and is associated with distinct membrane systems (17–19). The function of palmitoylation is not restricted to membrane attachment, but it also regulates protein-protein interaction and therefore multiple cellular processes (20). To date, over 400 P. falciparum proteins have been shown to be palmitoylated (21), including proteins essential for asexual development and erythrocyte invasion. PATs are polytopic membrane proteins, possessing four or more transmembrane domains (19, 22). Organization complements of PATs vary from over 20 members in metazoan species to only seven in the budding yeast Saccharomyces cerevisiae (23). Tetrahymena, Toxoplasma, and Plasmodium possess 38, 18, and 12 predicted PATs, respectively. PATs reside in different tissues and sub-cellular localizations and are thought to direct the recruitment of proteins to specific membranes: for example, Vac8 to vacuole membrane in yeast (24), RAS2 in mammalian cell lines to the plasma membrane (25), and TgARO in T. gondii to the rhoptry membrane (26, 27). A localization map of PATs in T. gondii and P. berghei was recently established, with two PATs found to localize to the IMC (26, 28). Here, we expand these studies to P. falciparum. We deliver a detailed phylogenetic analysis and dissect the specific sequence requirements that are essential for dual-acylated proteins to be localized to the IMC.

EXPERIMENTAL PROCEDURES

Cell Culture and Transfection of P. falciparum—P. falciparum (3D7) was cultured in human O+ erythrocytes according to standard procedures using complete Roswell Park Memorial Institute medium (29). Gametocytes were produced using a modified version of the established protocol (6, 30). For transfection, ring-stage parasites (10%) were electroporated with 100 μg of plasmid DNA resuspended in cytomix as described previously (31). Transfectants were selected using 10 μM WR99210 (for pARL-based vectors) or 30 μM blasticidin (for pBcam-based vectors), respectively. For single crossover integration, the following transfection parasites were alternately grown with and without WR99210 pressure (4 weeks for each interval off-drug) to promote integration of the transfection vectors in the endogenous loci. Double transfectant parasites expressing PfISP3-mCherry and PfDHHC1-GFP were generated by transfecting 100 μg of pB-PfISP3-mCherry into the stable transgenic cell line expressing 3D7-PfDHHC1-GFP. Double transfectant parasites expressing PfISP3-GFP and GAP45-mCherry were generated by transfecting 100 μg of pB-GAP45mCherry into the transgenic cell line expressing PfISP3-GFP. Selection was carried out using 30 μM blasticidin.

Nucleic Acids and Constructs—3D7 parasites expressing full-length PfDHHC1 and PfDHHC2 as a GFP fusion protein were generated by PCR amplification of the respective genes. PCR amplifications were performed using Phusion DNA polymerase (New England Biolabs). PCR amplicons were digested with Kpn1 and AvrII, cloned into the derivative of pARL-1a-GFP. Expression of the GFP fusion construct is controlled by the late stage promoter of ama1 (32). To circumvent the internal Kpn1 site in the genes coding PfDHHC3 and PfDHHC9, both open reading frames were cloned into the BamHI/Sall of a derivative of pBcamR (33). This vector mediates the overexpression of mCherry fusion proteins under the control of the ama1 promoter (6). The GFP replacement construct for tagging endogenous PfDHHC1 was designed using pARL-1a-GFP vector. 1 kb of the PfDHHC1 gene was cloned into the NotI/AvrII site of the pARL-1a-GFP releasing the crt promoter that controls the expression of the fusion protein in the original vector (34). All constructs were sequenced for accuracy. Oligonucleotides used for these constructs are summarized in Table 1.

Full-length and truncated versions of PfISP3, PfISP1, and GAP45 were previously cloned and localized in P. falciparum as overexpressing GFP- or mCherry fusion proteins (6, 10, 15). To dissect sequence requirements for membrane attachment nucleic acid substitutions introduced into these genes were generated by oligonucleotide directed mutagenesis. DNA fragments were digested with Kpn1 and AvrII, cloned into a derivative of pARL-based vectors. For single crossover integration, the respective oligonucleotides used for these constructs are summarized in Table 1.

Western Blot Analysis—Synchronized P. falciparum cultures were saponin-lysed using 0.03% saponin (Sigma). Parasite pellets were resuspended in adequate amount of PBS and 5% SDS loading dye. Proteins were separated on 10% SDS-PAGE and transferred to nitrocellulose membranes (Schleicher & Schuell). Monoclonal mouse anti-GFP (Roche Applied Science) or monoclonal rat anti-mCherry (Chromotek) was diluted 1:1000 in 10% (w/v) skim milk. The secondary antibody was horseradish peroxidase (HRP)-conjugated goat anti-rabbit immunoglobulin G (IgG) (1:3000, Jackson ImmunoResearch) and HRP-conjugated goat anti-mouse IgG (1:3000, Jackson ImmunoResearch).
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**Table 1**

| Oligonucleotides used in this study |
|------------------------------------|
| Restriction sites are underlined. |

## P. falciparum PfsISP3 constructs

| Primer     | Sequence |
|------------|----------|
| PfsISP3 FL fw | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 FL rv | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP3 CmFW | GAGAGGATCCATGGGAATTTAGGTTGAGT |
| PfsISP3 CmFL | GAGAGGATCCATGGGAATTTAGGTTGAGT |
| PfsISP3 G2A FL fw | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 G2A FL rv | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP3 C5A C6A FL fw | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 C5A C6A FL rv | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP3 C5A C5A FW | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 C5A C5A RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP3 C5A FW | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 C5A RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP3 C6A FW | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 C6A RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP3 N9D 10K FW | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 N9D 10K RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP3 N9D FW | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 N9D RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP3 D10G FW | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 D10G RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP3 D10K FW | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 D10K RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP3 D10R FW | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP3 D10R RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |

## P. falciparum PfsISP1 constructs

| Name       | Sequence |
|------------|----------|
| PfsISP1 FL rv | GATTCCTAGCCGATTTTTTTTTTTTTTTATATTTTTC |
| PfsISP1 C7A, C8A FW | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PfsISP1 C7A, C8A RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP1 G2A FW | GATTCCTAGCCGATTTTTTTTTTTTTTTATATTTTTC |
| PfsISP1 G2A RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP1 FL | GATTCCTAGCCGATTTTTTTTTTTTTTTATATTTTTC |
| PfsISP1 RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PfsISP1 D11K FW | GATTCCTAGCCGATTTTTTTTTTTTTTTATATTTTTC |
| PfsISP1 D11K RV | GCCGCTAGTGGCATTCAAACAATTTTTTG |

## P. falciparum PAT constructs

| Name       | Sequence |
|------------|----------|
| PFDHHC1.3 FL | GCCGCGTACCATGGGAATTTAGGTTGAGT |
| PFDHHC1.3 FL | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PFDHHC1 FL | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PFDHHC2 FL | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PFDHHC2 FL | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PFDHHC9 FL | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PFDHHC9 FL | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PFDHHC3 FL | GCCGCTAGTGGCATTCAAACAATTTTTTG |
| PFDHHC3 FL | GCCGCTAGTGGCATTCAAACAATTTTTTG |
ImmunoResearch). The immunoblots were developed by chemiluminescence using Immobilon Western Chemiluminescent DAB substrate (Millipore) and visualized using the ChemiDoc XR5+ System (Bio-Rad).

**Live Microscopy and Immunofluorescence Assays**—Images of unfixed GFP-expressing parasites were captured using a Zeiss Axio Skop 2plus microscope with a Hamamatsu Digital camera (model C4742-95, Zeiss AxioVision) with 1 µg/ml DAPI (Roche Applied Science) for nuclear stain. Immunofluorescence microscopy was performed on parasites fixed with a mixture of 4% formaldehyde and 0.0075% glutaraldehyde. Parasites were permeabilized with 0.1% Triton X-100 and blocked with 3% bovine serum albumin (BSA). First, parasites were incubated for 1 h with anti-GAPM2 (1:500, see Ref. 6) or RALP (1:500, see Ref. 35). Subsequently, cells were washed three times with PBS and incubated for an hour with Alexa-Fluor 594 goat anti-rabbit IgG antibodies (1:2000, Molecular Probes) and 1 µg/ml DAPI (Roche Applied Science). After removal of unbound antibodies by washing three times with PBS, coverslips were mounted with Fluoromount G (Sigma) and kept at 4 °C until evaluation.

**S-Acyl Biotin Switch Assay**—Saponin-lysed parasite pellets derived from 100 ml of synchronized late stage parasites were subjected to biotin switch protocol as described previously (15, 21).

**Solubility Assays**—*P. falciparum* proteins were extracted using 0.03% saponin (Sigma) from a synchronized late stage parasite culture (10 ml) as described previously (15). In short, the extraction was done sequentially; the lysate was resuspended in 100 µl of double distilled H2O, freeze-thawed three times, and centrifuged at 21,000 × g for 5 min. After this hypotonic lysis, the pellet was washed once with PBS and resuspended in 100 µl of freshly prepared 0.1 M Na2CO3 and kept on ice for 30 min to extract peripheral membrane proteins. The pellet was washed once with PBS and extracted for 30 min with 100 µl of 1% Triton X-100 and centrifuged at 21,000 × g for 5 min to obtain the integral membrane protein fraction in the supernatant. The final pellet was washed once with PBS and resuspended on 100 µl of PBS containing the insoluble fraction. Equal amounts of all supernatants were analyzed by immunoblotting with adequate amounts of 5 × SDS loading dye. Proteins were detected using anti-GFP antibodies, and the cytosolic protein GAPDH was used as a control.

**Phylogenetic Analysis**—To get a comprehensive understanding of the phylogenetic relation and evolutionary background of apicomplexan PATs, 16 apicomplexan proteomes were searched and the retrieved sequences analyzed.

**Data Collection of Putative PATs**—The catalog of putative Asp-His-His-Cys (DHHC) was identified by searching for proteins matching the zf-DHHC HMM profile (dCt region, ~170-amino acid profile corresponding to region encompassing DPG-GRD-TTXE motif), with HMMSEARCH of HMMER 3.1 package (36) using default parameters. The apicomplexan organisms used for this analysis are as follows: Cryptosporidium hominis TU502; Cryptosporidium muris RN66; Cryptosporidium parvum lowali; Babesia bovis T2Bo; Theileria annulata strain Ankara; Theileria parva strain Muguga; P. berghei ANKA; Plasmodium chabaudi chabaudi, Plasmodium cynomolgi strain B; P. falciparum 3D7; Plasmodium knowlesi strain H; Plasmodium vivax Sal-1; Plasmodium yoelii yoelii 17XNL; Neospora caninum Liverpool; T. gondii ME49; Eimeria tenella strain Houghton, and their proteomes were downloaded from EupathDB (37). The hits were then manually checked for the presence of cysteine-rich domain (CRD), an ~50-amino acid region housing the catalytically active DHH(Y)C residues (38). Hits lacking this domain were discarded. The remaining hits were segregated into “canonical” and “pseudo” PATs based on whether the CRD was intact or incomplete. The latter group includes hits that were incomplete/degraded at the N or C terminus of the CRD region, as well as those where the “DH(H/Y)C” motif was substituted. Many of the pseudo-PATs contain the DPG motif and TTXE motifs (13/28) thus arguing that these sequences are either degenerate sequences or represent artifacts of sequence assembly. The catalog of canonical and pseudo-PATs for all the species are listed in Fig. 4. Using the same methodology, putative PATs were identified in the proteomes of closely related free-living protists *Paramecium tetraurelia* and *Tetrahymena thermophila* SB210 and categorized, to enable a comparison of the evolution of the parasitic apicomplexan PATs with their free-living counterparts. The proteomes of *P. tetraurelia* and *T. thermophila* were downloaded from UNIPROT (39). Furthermore, the complement of characterized DHHC PATs from *Saccharomyces cerevisiae* S288c and *Homo sapiens* were extracted from SGD (40) and Swiss-Prot, respectively, to serve as the reference set of PATs.

**Identification of Evolutionarily Related Clusters of DHHC PATs**—The evolutionary relationships between the set of PATs was resolved based on rigorous phylogenetic analyses. Because the PATs in free-living protists are likely to be very similar due to the duplication events, a representative set of PATs was generated by clustering at 40% sequence identity using CD-HIT (41) for use in further analysis. Multiple sequence alignments were generated using Muscle (42) and Probcons (43) for the dCt region (corresponding to the Pfam HMM profile). The CRD region was not used for phylogenetic analyses because it contained poorer signals than the other two sets, presumably due to its smaller size (~50 amino acids in length), according to the likelihood mapping algorithm Tree-puzzle 5.2 (44). The alignment was manually edited using Geneious (created using Bioedit) and trimmed with TrimAl (45) using two cutoffs as follows: automated (which has been shown to be optimized for phylogenetic analyses using maximum likelihood method; Ref. 45) and gap threshold of 0.1 (which removes columns that contain gaps in >90% of sequences). The optimal model of substitution for each of the datasets was determined using ModelGenerator (46). Using the optimal model of substitution and allowing for rate heterogeneity at different sites modeled using γ-distribution, phylogenetic trees were then constructed using two methods, MrBayes and PhyML (47, 48). Bayesian phylogenies were generated using MrBayes for 3 million generations with a burn-in of 80,000 for two runs of four chains each. Maximum likelihood phylogenetic reconstruction was performed using PhyML with 1024 bootstrap replicates. This analysis resolved the PATs into six major clades and 14 subclades. Each of these clusters, including dCt region and full sequence, was then individually analyzed using the same series of steps out-
lined above to study whether the gene tree recaps the species tree or not.

**Analysis and Annotation of PAT Clusters**—The sequences in every cluster were analyzed for conservation of the following sequence motifs: consensus of CRD region; DPG motif, TTXE motif; and presence/absence of inserts between DPG motif and CRD region. Every cluster was also analyzed in terms of conservation of additional domains (wherever was present) and transmembrane helix distribution. Domain information was obtained from Interpro (49), whereas Tied Mixture Hidden Markov Model (TMHMM) (50) was used to predict transmembrane helices for the sequences.

**RESULTS**

**Identification of an IMC-Localized Palmitoyl Aciytransferase**—Three known IMC-associated proteins in *P. falciparum* (*PfGAP45, PfISP1*, and *PfISP3*) displayed a conserved myristoylation and palmitoylation site within the first 20 amino acids that
might be linked to IMC membrane association and may therefore be recognized by an IMC-residing PAT. Recent work in *T. gondii* and *P. berghei* revealed two IMC-localized PATs in each species as follows: *TgDHHC2* (TGME49_278850, homolog of *PfDHHC2*) and *TgDHHC14* (TGME49_293730, homolog of *PfDHHC1*) in *T. gondii* (26, 28) and *PbDHHC3* (PBANKA_092730, homolog of *PfDHHC3*) and *P6DHHC9* (PBANKA_093210, homolog of *PfDHHC9*) in *P. berghei* (28).

We aimed to localize all four of the homologous genes (PFC0160w/*PfDHHC1*, PFF0485c/*PfDHHC2*, PF11_0217/*DHHC3*, and PF11_0167/*PfDHHC9*) to validate their putative IMC localization in *P. falciparum*. We overexpressed the enzymes either as GFP or mCherry fusions, circumventing internal restrictions sites in the genes of interest. We used the *ama1* promoter to mimic the late transcription of most IMC proteins (51). In agreement with this, the transcription of all four PATs appears to be up-regulated to some level in schizonts (51). Localization of the fusion proteins in unfixed parasites revealed that *PfDHHC1*-GFP has a localization pattern resembling the IMC, and shows the IMC-specific dynamics during schizogony (6, 10, 52, 53); it commences as a cramp-like structure, transforming to small ring-shaped formations that toward the end of schizogony expand and are then equally distributed underneath the plasma membrane (Fig. 1A). This IMC localization was supported by the co-localization with the IMC marker GAPM2 (PFD1110w, glideosome-associated protein marker GAPM2) (26, 28) and *gondii* (PFC0160w/*PfDHHC1*, PFF0485c/*PfDHHC2*, PF11_0217/*DHHC3*, and PF11_0167/*PfDHHC9*) to validate their putative IMC localization in *P. falciparum*. We performed a systematic phylogenetic analysis of PATs within the alveolate lineage to gain insights into their evolutionary relationships.

*Phylogenetics Reveals Six Major Families of PATs in the Alveolata*—A recent study used the conserved, yet relatively short DHHC-CRD to construct a phylogeny of PAT proteins derived from several species of Apicomplexa (28). However, due to the shortness of the domain (~77 amino acids), phylogenetic insights were largely confined to comparisons of five families across four species. Here, we applied the hidden Markov model for zf-DHHC (PF01529; a 170-amino acid profile encompassing the DPG, CRD, and TTXE motifs) to retrieve 297 canonical and 28 pseudo (lacking an intact CRD and/or lacking the DH(HY)C motif) PATs from 18 alveolates. Compared with *S. cerevisiae* (seven PATs), apicomplexans tend to possess a larger repertoire of PATs (Fig. 3), with almost double the number in Hemosporidia (7–12) and even more in Coccidia (e.g. 18 in *T. gondii*). Furthermore, the free-living ciliates *P. tetraurelia* and *T. thermophila* show a much larger expansion in their DHHC PAT complement (100 and 38, respectively), likely related to documented duplication events (56, 57). These expansions highlight the potential importance of PATs to alveolate and apicomplexan innovations.

For information on evolutionary relationships between PATs, we performed maximum likelihood and Bayesian phylogenetic analyses on the region encompassed by the zf-DHHC domain (Fig. 4). This analysis allowed us to define six main clades of PATs, with clade 2 further divided into seven further sub-clades. Each clade and subclade is associated with distinct sequence features, including the presence of unique repeats, motifs, and clade-defining residues (Figs. 3 and 4). Three yeast PATs are found to associate with three of these clades (2e, 4 and 6), whereas 16 human PATs are found to cluster with the clades revealed identical dynamics of these two proteins (Fig. 2F), which would be congruent with a potentially specific interaction of these proteins. In pre-sexual gamocyte stages (Fig. 2G), *PfDHHC1*-GFP is not equally distributed within the nascent IMC but accumulates in transversal structures resembling the symmetric meshwork enclosing the gamocyte described previously as the localization of the *Plasmodium*-specific IMC protein MAL13P1.228 (6) that might reflect the sutures of individual IMC plates (55). Given the presence of multiple IMC-localized PATs in *T. gondii* and *P. berghei* and their differing distribution in *P. falciparum*, we performed a systematic phylogenetic analysis of PATs within the alveolate lineage to gain insights into their evolutionary relationships.

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![FIGURE 1. Overexpression and localization of PfDHHC1 (PFC0160w), PfDHHC2 (PFF0485c), PfDHHC3 (PF11_0217), and PfDHHC9 (PF11_0167) in late stage parasites. A, expression of PfDHHC1-GFP. Panel A1, Western blot analysis using anti-GFP antibodies. A single protein band of about 100 kDa (expected mass of 100 kDa) was detected in the transgenic but not in the parental parasite line. Panel A2, localization of PfDHHC1-GFP in unfixed parasites showing characteristic IMC dynamics during schizogony as follows: commencing as cramp-like structures (T1) and transforming to small ring-shaped formations (T2) that toward the end of schizogony expand and are then equally distributed underneath the plasma membrane (T3). Nuclei are stained with DAPI (blue). Enlargement of selected areas are marked with a white square and referred to as zoom. Scale bar, 1 μm. Panel A3, co-localization with the IMC marker GAPM2 (anti-GAPM2, red) in fixed cells confirmed PfDHHC1-GFP localization with its distinct biogenesis during schizogony (Fig. 2E). Co-localization with the dual-lipidated PfJP3 (PF14_0578) revealed identical dynamics of these two proteins (Fig. 2F), which would be congruent with a potentially specific interaction of these proteins. In pre-sexual gamocyte stages (Fig. 2G), PfDHHC1-GFP is not equally distributed within the nascent IMC but accumulates in transversal structures resembling the symmetric meshwork enclosing the gamocyte described previously as the localization of the Plasmodium-specific IMC protein MAL13P1.228 (6) that might reflect the sutures of individual IMC plates (55). Given the presence of multiple IMC-localized PATs in *T. gondii* and *P. berghei* and their differing distribution in *P. falciparum*, we performed a systematic phylogenetic analysis of PATs within the alveolate lineage to gain insights into their evolutionary relationships.*
2e, 2a2, and 3–6, suggesting the involvement of clade members in conserved cellular processes.

*PfDHHC1 Is a Member of a Highly Conserved Apicomplexan-specific Clade of PATs*—Across the 14 sub-clades are five that appear apicomplexan-specific (designated clusters 1a, 1b, 2c, 2dY, and 6a, Fig. 3). Cluster 1a, which includes *PfDHHC1* (PFC0160w), is highly conserved, yet specific to apicomplexans, with a single ortholog identified in all species with the exception of *P. yoelii*, in which the ortholog is a pseudo-PAT. However, comparison with its closest ortholog in *P. berghei* indicates that this is likely due to a poorly predicted gene model. Inspection of the sequences indicated conservation over the entire sequence length (data not shown). Furthermore, all members of cluster 1a (as well as 1b) are defined by the presence of ankyrin repeats.
at the N terminus, in addition to a distinct conserved insert between the DPG motif and the CRD region. Interestingly, the Toxoplasma ortholog TgDHHC14 (TGE49_293730) has also been associated with the IMC (26, 28). Cluster 2dY is similarly unique with the distinguishing feature that all members possess, i.e., “DHYC” motifs instead of “DHHC” motifs. Although DHYC-PATs have been identified in yeast (YDR264C/AKR1_YEAST and YOR034C/AKR2_YEAST), the apicomplexan DHYC-PATs represent a distinct clade, suggesting an instance of convergent evolution. From sequence logos constructed for each clade (Fig. 4), we observe that Cys-1 of the CRD region and Asn-40 are invariant, along with the active site residues DH(H/Y). In addition, Trp-31, Gly-37, and Phe-44 are also largely conserved, exhibiting only conservative substitutions. In the case of DHHC-PATs, Cys-4, His-14, Cys-15, Cys-18, Cys-21, and Cys-35 are also invariant; however, for DHYC-PATs, Cys-4, Cys-14, Cys-18, and Cys-21 are not conserved, whereas His-14 is substituted by Tyr-14, suggesting an alternative route for catalysis.

FIGURE 2. PIDHHCl is an IMC localized palmitoyl acyltransferase. A, schematic representation of the GFP replacement of the endogenous 3′ end of PIDHHCl creating a 3D7-PIDHHCl-GFP cell line. The vector encompasses the selection cassette (black), and 1 kb of the PIDHHCl gene (gray) was fused to GFP (green) accompanied by the 3′ UTR of PbdT (orange) without any promoter. Integration of the vector takes place by homologous recombination (cross) into the PIDHHCl locus creating a full-length PIDHHCl-GFP fusion under the control of the endogenous promoter. B, integration was confirmed by PCR using two different primer combinations on gDNA. One primer set (red) hybridizes in a PIDHHCl region upstream of the integration and in the coding region of GFP. This primer combination can only amplify a 1.7-kb DNA fragment after recombination took place. The other set (blue) amplifies 1.1 kb of PIDHHCl in the parental as well as in the transgenic parasite line. Control indicates PCRs with the red primer set in the absence of parasite DNA. C, expression of the transgene from the endogenous locus was shown by Western blot analysis using anti-GFP antibodies (upper panel) resulting in a fusion protein protein of ~100 kDa (calculated mass of 100 kDa) using late stage parasite material. Anti-GAPDH was used as a loading control. D, stage-specific expression pattern of PIDHHCl-GFP using synchronized 3D7-PIDHHCl-GFP parasite material (harvested after 8, 16, 24, 32, 40, and 48 h) and anti-GFP antibodies. Antibodies directed against the IMC marker protein GAMP2 were used as a stage-specific control. Anti-GAPDH antibodies were used as loading control. E, PIDHHCl-GFP was localized in unified late stage parasites showing characteristic IMC dynamics during schizogony (T1 to T3). Nuclei stained with DAPI (blue). Enlargement of selected areas are marked with a white square and referred to as zoom. Scale bars, 1 μm. F, co-localization of PIDHHCl-GFP and PISp3.MCherry. PISp3.MCherry was episomally co-expressed and shows identical spatial distribution compared with PIDHHCl- GFP. G, PIDHHCl-GFP distribution within the nascent IMC during gametocytogenesis (stage IV and V). These symmetric sutures of the IMC are expanding with ongoing growth of the IMC vesicles and maturation of the gametocytes (stage IV and V). DIC, differential interference contrast. Scale bars, 1 μm.

FIGURE 3. Phylogenetic analyses of PATs from Apicomplexa, free-living protists, yeast, and human. The phylogenetic tree reconstructed using MrBayes based on dCt region of all canonical PATs from apicomplexa, yeast, and human, and nonredundant representative set from free-living protists. The clusters with posterior probabilities greater than ~0.6 are shown as clades, with posterior probabilities and bootstrap replicates indicated at the branching point. Clusters consisting only of apicomplexan members are shown in orange, and clusters consisting of sequences from both apicomplexan organisms and the free-living protists are shown in cyan. The yeast PATs that are orthologs of specific clusters are labeled in brown, and human orthologs are shown in blue. Features characteristic of a cluster are indicated alongside the cluster.

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217870 and PBANKA_093210, have previously been shown to localize to the ER and the IMC, respectively, compared with the apical membrane/food vacuole for PIDHHCH9 (28). Given the disparity in localization of these orthologs, we speculate that after their emergence the members of this clade have undergone adaptive radiation resulting in the evolution of lineage-specific functions.

Outside the apicomplexan-specific clusters, we note several instances of alveolate-specific clusters (2a1, 2b, 2dH, and 5a, a subclass of 2dH) lacking the TTXE motif, see Fig. 3). Intriguingly, with the exception of 2dH, members from these clusters appear to have been lost in the hemosporidians. Cluster 2dH is noteworthy as orthologs are present for all alveolates except N. caninum, and therefore likely represent a conserved alveolate innovation.

For the most part, lineage-specific duplications within subclades are rare with the exception of duplicates in T. gondii in clades 2dH, 5, and 6a, although piroplasmids contain two copies of PATs in cluster 6b. Finally, clusters 2a2, 2e, 3–5, and 6b appear conserved outside the alveolates with yeast and human orthologs, offering the opportunity to gain insights into func-
tions of cluster members. This is of particular relevance with two of the clades, 2e and 3, that contain the non-IMC membrane-localized PATs PF11_0217/PfDHHC3 and PFF0485c/PfDHHC2, respectively.

Palmitoylation of the IMC Localized PfISPs—The two members of the ISP family in P. falciparum, PfISP1 and PfISP3, homologs of TgISP1 and TgISP3 (10, 11), encode N-terminal myristoylation and palmitoylation motifs. The N-terminal 20 amino acids of both proteins are sufficient for IMC membrane association (15), suggesting that they are recognized and modified by the IMC-residing PATs.

First, we verified the role of N-terminal myristoylation and palmitoylation motifs for IMC membrane recruitment by expressing and analyzing the subcellular distribution of mutants by microscopy (Figs. 5 and 6). Expression of full-length PfISP3-GFP shows a typical IMC dynamic during schizogony (Fig. 5A) and co-localizes with the IMC markers GAP45 and GAPM2 (Fig. 5, panels A2 and A3). Alanine substitutions of the myristoylation or palmitoylation motifs led to the expected cytosolic distribution of the mutant proteins, underlining the synergistic effect of myristoylation and palmitoylation for membrane association (Fig. 5, B and C). Noteworthy, single point mutations of either C5 or C6 in PfISP3 do not interfere with IMC membrane localization, which might point toward a redundancy of the palmitoylation sites (Fig. 5, D and E). Cytosolic distribution of PfISP3C5AC6A-GFP and PfISP1C7AC8A-GFP was additionally verified by solubility assays. Although overexpressed PfISP3-GFP is mainly in the carbonate fraction, confirming its membrane association, both mutants are exclusively found in the hypotonic fraction, resembling the cytosolic GAPDH control. (Fig. 5F). The pertinent alanine substitutions (20 PISP1G2A-GFP and 20 PISP1C7AC8A-GFP) were also introduced in PfISP1 and resulted in similar phenotypes (Fig. 6, A–C).

Second, to confirm the predicted palmitoylation in vivo, parasites expressing either the wild type sequence (20 PISP3-GFP and 20 PISP1-GFP) or the corresponding cysteine/alanine substitutions (20 PISP3C5AC6A-GFP and 20 PISP1C7AC8A-GFP) were used in acyl biotin exchange assays (21, 58). This assay substitutes thioester-linked acyl groups with biotin that can subsequently be used for affinity purification. These experiments showed that both PfISPs (20 PISP3-GFP and 20 PISP1-GFP) are biotinylated and are enriched in the fraction that is expected to contain all S-acylated proteins assayed by this method. The corresponding mutants without
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FIGURE 5. Role of N-terminal acylation for IMC membrane association of PfISP1 and acyl biotin exchange assays. A, overexpression of PfISP1-GFP showing characteristic IMC dynamics as PfISP3-GFP. Putative myristoylation and palmitoylation sites are highlighted in light blue (G) or green (C,C). B and C, mutation of either one of the acylation motifs (PfISP1GGA-GFP and PfISP1GACA-GFP) resulted in a cytosolic variant shown by microscopy. Nuclei are stained with DAPI (blue). Enlargement of selected areas are marked with a white square and referred to as zoom. Scale bar, 1 μm. D, PfISP3-GFP (upper panel) and 20PfISP3GACA-GFP (lower panel) fusion proteins are present in the 2 aliquots that were incubated with biotinylation reagent with or without hydroxylamine after N-ethylmaleimide treatment (Loading). After elution from the NeutrAvidin beads (Avidin beads), only 20PfISP3-GFP but not the cysteine mutant 20PfISP3GACA-GFP is enriched in the hydroxylamine (+HA)-treated sample when compared with the untreated sample (−HA). E, PfISP1-GFP, PfISP1-GFP fusion proteins are present in the 2 aliquots that were incubated with biotinylation reagent with or without hydroxylamine after N-ethylmaleimide treatment (Loading). After elution from the NeutrAvidin beads (Avidin beads), only PfISP1-GFP but not the cysteine mutant PfISP1GACA-GFP is enriched in the hydroxylamine (+HA)-treated sample when compared with the untreated sample (−HA). DIC, differential interference contrast.

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FIGURE 6. Role of N-terminal acylation for IMC membrane association of PfISP3 and PfISP1 and acyl biotin exchange assays. A, overexpression of PfISP1-GFP showing the same characteristic IMC dynamics as PfISP3-GFP. Putative myristoylation and palmitoylation sites are highlighted in light blue (G) or green (C,C). B and C, mutation of either one of the acylation motifs (PfISP1GGA-GFP and PfISP1GACA-GFP) resulted in a cytosolic variant shown by microscopy. Nuclei are stained with DAPI (blue). Enlargement of selected areas are marked with a white square and referred to as zoom. Scale bar, 1 μm. D, PfISP3-GFP (upper panel) and 20PfISP3GACA-GFP (lower panel) fusion proteins are present in the 2 aliquots that were incubated with biotinylation reagent with or without hydroxylamine after N-ethylmaleimide treatment (Loading). After elution from the NeutrAvidin beads (Avidin beads), only PfISP3-GFP but not the cysteine mutant 20PfISP3GACA-GFP is enriched in the hydroxylamine (+HA)-treated sample when compared with the untreated sample (−HA). E, PfISP1-GFP, PfISP1-GFP fusion proteins are present in the 2 aliquots that were incubated with biotinylation reagent with or without hydroxylamine after N-ethylmaleimide treatment (Loading). After elution from the NeutrAvidin beads (Avidin beads), only PfISP1-GFP but not the cysteine mutant PfISP1GACA-GFP is enriched in the hydroxylamine (+HA)-treated sample when compared with the untreated sample (−HA). DIC, differential interference contrast.

Lysine but Not Arginine within Close Proximity of the Palmitoylation Sites of PfISP3 Prevents IMC Membrane Targeting—The distribution of 20PfISP31–20A-GFP points toward a crucial role of the first 10 amino acids for IMC targeting (Fig. 7A). This was validated by the expression and localization of the first 10 amino acids of PfISP3 fused with GFP (10PfISP3-GFP). Consistent with the previous findings10PfISP3-GFP is localized to the IMC (Fig. 7B). Next, the only charged residue (Asp-10) in the 10 amino acids was exchanged either to a neutral glycine (Fig. 7C) or to a positively charged arginine (Fig. 7D) or lysine (Fig. 7E). Although neither the exchange with glycine (10PfISP3D10G-GFP) nor arginine (10PfISP3D10K-GFP) interfered with correct targeting, the substitution with a lysine (10PfISP3D10K-GFP) led to a re-direction of the GFP fusion to the periphery of the nascent merozoites, resembling plasma membrane localization.

The aberrant localization of 10PfISP3D10K-GFP was confirmed by co-localization using the IMC marker protein GAPM2 (Fig. 7E). Interestingly, previous work showed an essential role for an...
arginine and a lysine in close proximity downstream of the palmitoylated cysteine for rhoptry membrane targeting of \( \text{PfARO} \) (15). To probe into a putative discriminative role of an arginine in combination with a lysine, we attempted to re-direct \( \text{PfISP3-GFP} \). As expected, the exchange of Asn-9 with aspartate (\( \text{PfISP3N9D-GFP} \)) did not change IMC localization (Fig. 8A), and the exchange of Asn-9 and Asp-10 with lysines (\( \text{PfISP3N9K10L-GFP} \)) re-directed the protein to the periphery of the merozoites (Fig. 8B). Nevertheless, the substitution of Asn-9 and Asp-10 with an arginine and lysine, respectively

FIGURE 7. Mutational analysis of the minimal sequence requirements for IMC association of PfISP3. A and B, expression and localization of the 20-amino acid N terminus with the last 10 (\( \text{PfISP3}_{11-20\alpha}-\text{GFP} \)) amino acids substituted with alanines. Panel A1, \( \text{PfISP3}_{11-20\alpha}-\text{GFP} \) was localized in unfixed parasites and showed an identical dynamic as the wild type protein. Panel A2, Western blot analysis of the expression using anti-GFP antibodies. B, first 10 amino acids of PfISP3 (\( \text{PfISP3}-\text{GFP} \)) are sufficient for IMC localization (panel B1). Panel B2, Western blot analysis of the expression of PfISP3-GFP. C and D, mutation of the sole charged residue D10 into a neutral glycine (\( \text{PfISP3D10G-GFP} \), panel C1) or a positively charged arginine (\( \text{PfISP3D10R-GFP} \), panel D1) does not interfere with IMC targeting. Western blot analysis (panels C2 and D2) of GFP fusion protein expression. E, substitution of D10 with a positively charged lysine re-directs the GFP fusion protein to the periphery of the nascent merozoites. The zoom highlights the additional association with the food vacuole membrane characteristic for plasma membrane proteins (panel E1). Co-localization was with the IMC marker GAPM2. Nascent IMC (anti-GAPM2, red) is clearly distinguishable from the peripherally associated \( \text{PfISP3D10K-GFP} \) in early stages (T1) and congruent in late stages (T3). Western blot analysis (panel E3) of \( \text{PfISP3D10K-GFP} \). DIC, differential interference contrast.
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FIGURE 8. Re-direction of the IMC 10PfISP3-GFP fusion protein to the rhoptry membrane. A, substitution of Asn-9 with Asp (10PfISP3N9D-GFP, A) does not interfere with correct IMC recruitment of the GFP fusion protein (panel A1). Nuclei were stained with DAPI (blue). Panel A2, Western blot analysis using anti-GFP antibodies. B and C, substitution of N9 and D10 with lysine and its expression in the parasite (10PfISP3N9KD10K-GFP) leads to a peripheral localization (panel B1). The zoom highlights the additional association of this mutant protein with the food vacuole membrane. Panel B2, Western blot analysis using anti-GFP antibodies. Panel B3, co-localization with the IMC marker GAPM2. Nascent IMC (anti-GAPM2, red) is clearly distinguishable from the peripherally associated 10PfISP3D11K-GFP in early stages (T1) and congruent in late stages (T3). C, in contrast, the substitution of N9 with arginine (10PfISP3N9RD10K-GFP) re-directs the fusion protein to cell periphery (zoom 1) and the apical pole (zoom 2, panel C1). Panel C2, co-localization with the IMC marker GAPM2. Nascent IMC (anti-GAPM2, red) is clearly distinguishable from the apical 10PfISP3N9RD10K-GFP in early stages (T1) and late stages (T3). Panel C3, Western blot analysis using anti-GFP antibodies. DIC, differential interference contrast.

(10PfISP3N9RD10K-GFP), led to a partial re-direction of the GFP fusion to a structure that co-localizes with the rhoptry marker RALP (Fig. 8C).

PFISP1 Uses the Same N-terminal Signature for IMC Targeting as PfISP3—Given the similar N-terminal sequence signature of both PfISP proteins, we probed into the minimal sequence requirements for IMC-specific membrane recruitment of PfISP1 (Fig. 9). First, we expressed minimal targeting sequences by using only the first 11 (including the negative charged residue) or the 10 amino acids of PfISP1. As expected, based on the previous results from PfISP3, both constructs, 11PfISP1-GFP and 10PfISP1-GFP, directed GFP to the IMC (Fig. 9, A and B). Again, the introduction of a lysine in this short targeting sequence (11PfISP1D11K-GFP) aborted IMC trafficking and led to an aberrant membrane association (Fig. 9C).

DISCUSSION

The specific recruitment of proteins to endomembrane systems such as the IMC is a prerequisite for their physiological role. The IMC has three main functions as follows: (i) it plays a major role in motility and invasion; (ii) it confers stability and shape to the cell; and (iii) it provides a scaffolding framework during cytokinesis. Therefore, it is not surprising that it harbors a structurally, phylogenetically, and functionally diverse proteome. To date, about 30 IMC proteins have been identified so far in Apicomplexa, with the majority being associated with the cytosolic face of the IMC (3). Most of these peripheral membrane proteins display predicted palmitoylation motifs either in combination with an N-terminal myristoylation motif (e.g. PfISP1, PfISP3, and GAP45) or without (e.g. alveolins).

Palmitoyl acyltransferases embedded in distinct membranes are thought to play an important role for membrane-specific
recruitments of these proteins. Here, we show that PfDHHC1 is, like its homolog TgDHHC14 in T. gondii (26), IMC-localized. Future work has to address the localization of PfDHHC2, PfDHHC3, and PfDHHC9. Although previous analyses reveal PfDHHC1 to be a member of a conserved clade (28), our comparative analyses involving ciliates establishes that this PAT is an apicomplexan-specific innovation.

By augmenting the PAT repertoire in apicomplexans with those from alveolates, yeast and human, our study enables us to differentiate those PATs representing alveolate and apicomplexan-specific innovations from those mediating core conserved eukaryotic functions. Furthermore, our phylogenetic analysis defines six distinct clades of PATs, supported by the presence of distinguishing molecular signatures. In general, PATs exhibit few lineage-specific innovations and are largely conserved across many species. This conservation, when considered in combination with the presence of clade-specific residues in the catalytic domain, argues that alveolates and apicomplexans have recruited this ubiquitous family of enzymes to perform a central role in their biology.

Although considerable efforts were undertaken to identify specific sequence requirements for substrate specificity of PATs, a clear consensus sequence for individual PATs has not yet been identified. On the contrary, experimental data suggest overlapping substrate specificity between enzymes (23, 35, 59, 60), leaving us with the question of how peripheral membrane proteins are recruited to specific membrane systems such as the IMC in the malaria parasite. In a previous study, we tracked 17 proteins predicted to be myristoylated and palmitoylated in the first 20 N-terminal amino acids in the malaria parasite (15). The use of this short sequence excluded putative interference with other determinants for membrane specificity like protein-protein interaction or structural features and focused on primary sequence motifs. Interestingly, although most of the corresponding GFP fusion proteins were trafficked to the parasite plasma membrane, three were sorted to the apical organelles and two, PfISP1 and PfISP3, were trafficked to the IMC. Both groups use a different molecular signature. Although rhoptry membrane attachment depends on positively charged residues in addition to an arginine two amino acids apart from the palmitoylated cysteines, a positively charged lysine appears to be the only exclusion criterion for an IMC localization (Figs. 8 and 9). In agreement with this, all experimentally localized rhoptry fusion proteins (PF08_0062, PFL1110c), including PfARO (PFD0720w), display an arginine residue two amino acids apart from the palmitoylated cysteines (15). It will be highly interesting to dissect the discriminative parameters between IMC and plasma membrane-localized proteins.

To help define substrate specificity, we generated a structural model of the CRD domain (Fig. 10). The structure features two lobes, with one lobe containing a set of conserved Cys and His residues that are spatially close and may therefore form the
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Figure 10. Structural model for the CRD region. Based on PDHHC7 (Pf3D7_0528400, 2DH), a model was generated using I-TASSER (65), the best performing web server for protein structure prediction according to the community-wide Critical Assessment of Protein Structure Prediction (CASP) experiment (66). The fit of the sequence to the modeled structure was verified using PROSA (67), which returned a z-score of −4.54, comparable with those observed for experimental structures of this size. The conserved residues of the CRD region are shown as sticks. The different residues are colored as follows: DHHHC motif in blue, invariant Cys-1 and Asn-40 in magenta, highly conserved Cys and His residues in orange, and the largely conserved Gly-37, Trp-31, and Phe-44 in white. The residues are marked according to the number shown in the consensus sequence logo for the CRD region in Fig. 3. The backbone of cluster defining residues for the PATs of interest are shown as red sticks.

zinc-binding pocket. The zinc finger domain is a short independent domain present in many proteins, consisting of a conserved mosaic of cysteine and histidine residues, a characteristic feature of the CRD domain of DHHHC PATs, which requires zinc binding to stabilize its structure (63). Recently, zinc binding has been experimentally shown to be necessary for stabilizing the structure of the CRD domain in a yeast DHHHC PAT, the Swf1 protein (64). Within the other lobe, the invariant Asn-40 appears to form a hydrogen bond with the active site His-27, whereas the conserved Gly-37, located close to His-27 at a constrained turn, is likely important for packing. The conserved Trp-31 and Phe-44 may interact through their aromatic side chains, helping to maintain orientation of the nearby active site residues. Outside these conserved residues, we identified a number of cluster-defining residues that may serve to define substrate specificity. We note that different clades of PATs are associated with distinct sets of clade-specific residues and motifs (Fig. 10). Interestingly, both PDHHC1 and PDHHC9, members of apicomplexan specific groups, feature a clade-defining residue close to the active site of the CRD (Asp-33 and Gly-33, respectively). We speculate that these residues mediate discriminatory roles for the DHHHC catalytic site of these two proteins. In addition, we also show that PDHHC9 and PDHHC2 (PF0485c) feature clade-defining residues close to the likely zinc-binding pocket (Glu-11 and Ser-17, respectively), which may have implications for the function of this site. For example, Glu-11 may be involved in interactions with positively charged side chain residues associated with substrates targeted to the plasma membrane (e.g. via the formation of stabilizing salt bridges). Finally, although PDHHC3 (PF11_0127) does not feature any conserved clade-defining residues in the CRD region, we do note that cluster 3 members differ from all other PATs as the only group without the well-conserved Gly of the DPG motif, present next to the second transmembrane helix. This might have potential consequences for its embedment into the membrane.

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