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Discovery of a novel and conserved *Plasmodium falciparum* exported protein that is important for adhesion of PfEMP1 at the surface of infected erythrocytes

Adéla Nacer,1,2,3§ Aurélie Claes,1,2,3 Amy Roberts,1,2,3 Christine Scheidig-Benatar,1,2,3 Hiroshi Sakamoto,1,2,3† Mehdi Ghorbal,1,2,‡ Jose-Juan Lopez-Rubio,1,2,‡ and Denise Mattei1,2,3,*

1Biology of Parasite-Host Interactions Unit, Department of Parasites and Insect Vectors, Institut Pasteur, 25, Rue du Dr. Roux, Paris F-75015, France.
2INSERM U1201, 25, Rue du Dr. Roux, Paris F-75015, France.
3CNRS ERL9195, 25, Rue du Dr. Roux, Paris F-75015, France.

Summary

*Plasmodium falciparum* virulence is linked to its ability to sequester in post-capillary venules in the human host. *Plasmodium falciparum* erythrocyte membrane protein 1 (PfEMP1) is the main variant surface antigen implicated in this process. Complete loss of parasite adhesion is linked to a large subtelomeric deletion on chromosome 9 in a number of laboratory strains such as D10 and T9-96. Similar to the cytoadherent reference line FCR3, D10 strain expresses PfEMP1 on the surface of parasitized erythrocytes, however without any detectable cytoadhesion. To investigate which of the deleted subtelomeric genes may be implicated in parasite adhesion, we selected 12 genes for D10 complementation studies that are predicted to code for proteins exported to the red blood cell. We identified a novel single copy gene (PF3D7_0936500) restricted to *P. falciparum* that restores adhesion to CD36, termed here virulence-associated protein 1 (*Pfvap1*). Protein knockdown and gene knockout experiments confirmed a role of PfVAP1 in the adhesion process in FCR3 parasites. PfVAP1 is co-exported with PfEMP1 into the host cell via vesicle-like structures called Maurer’s clefts. This study identifies a novel highly conserved parasite molecule that contributes to parasite virulence possibly by assisting PfEMP1 to establish functional adhesion at the host cell surface.

Introduction

*Plasmodium falciparum*, a causative agent of human malaria, is an apicomplexan parasite that infects and develops in host erythrocytes. *Plasmodium falciparum* is the most virulent form of the five *Plasmodium* species that infect man. It is responsible for over half a million deaths annually, principally in children under the age of 5 (WHO, 2013). Several strategies have been implemented in order to reduce the burden of infection such as transmission control and the development of drugs and vaccines (Eastman and Fidock, 2009; Crompton et al., 2014). However, malaria remains a major public health concern. Determining the factors that contribute to parasite virulence is therefore crucial for designing or improving therapeutic interventions.

*Plasmodium falciparum*-mediated pathogenesis has been associated with the parasite's ability to mediate binding of its host cell to the vascular endothelium of several organs, notably the brain leading to complications such as cerebral malaria. The mechanisms that lead to pathology are not well elucidated and include the activation of platelets and endothelial cells leading to increased binding of infected erythrocytes (IE) either directly to the vasculature or via platelet bridges (Miller et al., 2013). Infected erythrocyte adhesion to various host cell receptors such as intercellular adhesion molecule-1, CD36,
gC1-qR/p32 and chondroitin sulphate A is mediated by the main parasite ligand *P. falciparum* erythrocyte membrane protein 1 (PIEMP1; Smith *et al.*, 2013), an antigenically variant surface protein, encoded by approximately 60 var genes expressed in a mutually exclusive manner (Scherf *et al.*, 2008). Specific subsets of var genes are associated with IE sequestration to particular tissues in the placenta (Viebig *et al.*, 2005) or the brain in cerebral malaria patients as determined by post-mortem studies (Tembo *et al.*, 2014). Thus, in-depth insight into parasite proteins that contribute to the PIEMP1-mediated adhesion process could enable intervention strategies to reduce cytoadhesion-associated malaria pathogenesis. A systematic gene knockout study of candidate genes coding for proteins trafficked to the IE illustrated that many helper proteins are involved to traffic PIEMP1 from the parasite to the surface of IE (Elsworth *et al.*, 2014).

A different type of cytoadhesion deficient parasites is linked to a large subtelomeric deletion on the right arm of chromosome 9 of cloned lines such as D10 and T9-96 (Day *et al.*, 1993; Chaiyaroj *et al.*, 1994). D10 lacks 25 genes from this region including the cytoadherence-linked asexual gene 9 (*clag9*), ring exported proteins 1–4 (*rex1, rex2, rex3* and *rex4*), several rifin, two var genes. The export of PIEMP1 in D10 is comparable in parasites with an intact chromosome 9 indicating that the PIEMP1 expressed at the surface of D10 IE lost its virulence feature, namely the adhesion to host receptors (Nacer *et al.*, 2011). This suggests that at least one of the deleted genes of chromosome 9 is important for correct presentation of PIEMP1 at the IE surface. CLAG9 had previously been implicated in cytoadhesion (Trenholme *et al.*, 2000); however, it has been shown that disruption of the *clag9* gene does not impair binding to CD36 suggesting that other genes are involved in this process (Nacer *et al.*, 2011). One promising candidate is the resident Maurer’s cleft protein REX1. Knockout of *rex1* leads to a slight reduction in binding to CD36, and to morphological alterations of the Maurer’s cleft similar to those observed in D10 (Dixon *et al.*, 2011; Nacer *et al.*, 2011). Nonetheless, as binding remains partial, this suggests that other genes absent from chromosome 9 in D10 parasites could be implicated in the correct conformation/presentation of PIEMP1 at the IE surface.

In this work, we complemented 12 genes that are bioinformatically predicted to be exported to the IE. We describe the identification of the virulence-associated protein gene (*Pfvap1*) that restores adhesion of complemented D10 IE to CD36. This gene is unique in *P. falciparum* and its deletion resulted in a 50% to 60% decrease in binding to CD36. We propose a mechanism by which PIVAP1 interacts with PIEMP1.

**Results**

**Identification of a novel gene implicated in cytoadhesion**

We previously showed that non-cytoadherent D10 parasites with a subtelomeric deletion on the right arm of chromosome 9 express PIEMP1 on the IE surface but that the protein has lost its adhesion properties (Nacer *et al.*, 2011). We detected strong surface labelling on IE of D10 and the reference strain for parasite cytoadhesion FCR3 with a pool of hyperimmune human sera (Fig. 1A), known to predominantly recognize PIEMP1 (Chan *et al.*, 2012), thus supporting the notion that the non-binding phenotype of D10 may be due to a conformational defect or incorrect presentation of PIEMP1.

The criteria chosen to identify genes on chromosome 9 with a potential role in cytoadhesion included the presence of export motifs PNEP, PEXEL/VTS (Marti and Spielmann, 2013) and transcription during the intraerythrocytic cycle (www.plasmodb.org). On this basis, 12 genes were chosen to complement D10 (Fig. 1B), none of which have been previously studied. We performed binding assays on purified CD36 receptor coated onto plastic dishes to determine whether episomal expression of a particular candidate could restore adhesion. One out of 12 transfected parasite lines, complemented with PF3D7_0936500 (former ID: PFI1765c), showed a significant restoration of binding to CD36 (20-fold over D10 IE counts). We named this gene *Pfvap1*. Transcription of *Pfvap1* during the intraerythrocytic cycle was confirmed by Northern blot and a 2.3 kb transcript was detected from total RNA of asynchronous cultures of 3D7 and FCR3 (Fig. 1C). *Pfvap1* is expressed in ring stages with timing similar to PIEMP1; maximal expression was detected in highly synchronized ring stage cultures by quantitative real-time polymerase chain reaction (qRT-PCR; Fig. 1D) confirming previous reports (Le Roch *et al.*, 2003; Otto *et al.*, 2010). Homology searches failed to identify other homologues for *Pfvap1* in *P. falciparum* and no orthologues were found in other *Plasmodium* species, suggesting that this single copy gene is specific to *P. falciparum*. With the exception of trafficking motifs, no other known protein domains could be identified in PIVAP1.

**PIVAP1 partially restores binding to CD36**

Complementation of D10 parasites with *Pfvap1* (*D10*<sup>Pfvap1</sup>) resulted in a significant increase of binding to CD36 over background values observed for D10 (Fig. 2A). *D10*<sup>Pfvap1</sup> adhesion values, when compared with previously selected CD36 laboratory lines FCR3 and 3D7 (Fig. 2B), suggest that only partial restoration of adhesion in D10 parasites was achieved. Wild-type D10 parasites do not exist, and thus, the binding capacity of this parasite before
Fig. 1. Complementation analysis identified a novel gene that restores adherence in D10.
A. Synchronized FCR3 and D10 parasites (30–35 h) were labelled with a pool of hyperimmune human sera followed by an anti-human IgG Alexa 488 secondary antibody (green). Fluorescence was measured by flow cytometry and imaged by fluorescence microscopy (inset).
B. Schematic representation of the chromosome 9 deletion in *P. falciparum* D10 (D10) compared with *P. falciparum* 3D7 (3D7). Complementation of D10 using episomal expression of candidate genes (shown in back boxes).
C. A transcript of ∼2.3 kb was detected in asynchronous cultures of *P. falciparum* FCR3 and 3D7.
D. qRT-PCR analysis of synchronized FCR3 parasites showed maximal expression in ring stage parasites. Y-axis relative copy number normalized against seryl transferase and inositol. X-axis age of parasites (hours post-invasion).
chromosome 9 deletion cannot be assessed. We generated a Ty1-tagged \textit{Pfvap1} gene and showed that the addition of the tag to the C-terminus did not change the restoration of adhesion levels (Fig. 2A).

Complementation was confirmed by Southern blot of D10\textsuperscript{Pfvap1} genomic DNA digested with Dral and XmnI hybridized with a \textit{Pfvap1} probe where fragments of expected sizes were observed (Fig. 2C).

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**Fig. 2.** \textit{PfVAP1} partially restores binding to CD36.

A. Complementation of D10 with \textit{PfVAP1} (episome) leads to a > 20-fold increase in binding.

B. Binding of FCR3, 3D7 and D10 to purified CD36. Binding assays performed in duplicate wells spotted with either CD36 or BSA. Error bars (SD) are shown. FCR3 (\(n = 5\)), 3D7 (\(n = 2\)), D10 (\(n = 6\)) and D10\textsuperscript{Pfvap1} (\(n = 10\)) replicates. D10 complemented with \textit{PfVAP1} or \textit{PfVAP1-Ty1} binds similarly to CD36. Binding assays were performed following four rounds of panning on CD36 purified CD36 or BSA. Error bars represent the standard deviation.

C. Diagram showing the position of restriction sites for Dral and XmnI used for digestion of the genomic DNA of complemented D10\textsuperscript{Pfvap1}.

D. Diagram confirming the complementation of D10 with \textit{Pfvap1}. Genomic DNAs were digested with Dral (lane 1), XmnI (lane 2), Dral + XmnI (lane 3) or not digested (lane 4). Complemented D10\textsuperscript{Pfvap1} parasites, D10 and FCR3 are shown. The absence of a reaction on D10 confirmed the specificity of the probe.

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Pfvap1 has a typical exported protein gene organization with a PEXEL motif in the beginning of the second exon (Sargeant et al., 2006), encoding for a protein with a predicted molecular weight of 31.1 kDa with two predicted C-terminal transmembrane domains (Fig. 2D). A single nucleotide polymorphism resulting in a non-synonymous amino acid substitution at position 117 (T for K) has been documented and occurs in 8 out of 10 of the laboratory strains examined (Aurrecoechea et al., 2009).

PfVAP1 is exported to the red blood cell (RBC) membrane

We generated D10 parasites complemented with Ty1-tagged PfVAP1 to investigate the subcellular localization and temporal expression profile. PfVAP1 is synthesized by ring stage parasites (from 8–11 h post-invasion) onwards. In D10Pfvap1 early trophozoites PfVAP1 is present in the parasitophorous vacuole (PV) where it partially co-localizes with PfEMP1 (PV) and appears to localize in distinct regions, separate from PfEMP1 at the erythrocyte membrane (fourth row). Scale bars = 5 μm.

Fig. 3. Localization of PfVAP1 in complemented D10 parasites. Ty1-tagged D10Pfvap1 parasites were labelled with anti-Ty1 monoclonal antibody to determine the localization of the protein. In early trophozoites PfVAP1 is associated with the parasitophorous vacuole (PV) as determined by double labelling with PfExp1. It is then trafficked via the Maurer’s clefts (PfSBP1) to the infected erythrocyte membrane. In the Maurer’s clefts, PfVAP1 co-localizes with PfEMP1 (third row) and appears to localize in different regions, separate from PfEMP1 at the erythrocyte membrane (fourth row). Scale bars = 5 μm.
the PV until trophozoite stages and then the protein associates with the Maurer’s clefts (24–36 h) as shown by labelling with PfSBP1 antibodies. PfEMP1 and PfVAP1 transiently co-localize in the Maurer’s clefts suggesting that they interact in this particular compartment or that they are co-trafficked (Fig. 3; Supporting Information Fig. S1), they then appear to ‘dissociate’ when they reach the RBC membrane as very little overlap is evident in schizonts (Fig. 3). Similar to PfEMP1, PfVAP1 gives a punctate labelling pattern at the RBC membrane in schizonts (Fig. 3).

PfVAP1 plays a major role in cytoadhesion of FCR3-IE

To directly assess the role of PfVAP1 in adhesion in the FCR3 reference strain, we used a protein knockdown approach in which Pfvap1 was cloned in frame with an E. coli dihydrofolate reductase (DHFR) destabilization domain (Muralidharan et al., 2011). The original pGDB2 vector was modified by the insertion of a human DHFR (hDHFR) cassette to transfect parasites without prior resistance to WR9220, necessary for the maintenance of the stable protein with trimethoprim (TMP; Fig. 4A; Supporting Information Table S1). In addition, we replaced GFP by a haemagglutinin A tag (HA-tag). Mutant parasites were selected with blasticidin-S (BSD) and following integration by single crossover homologous recombination, clones were obtained by limiting dilution and validated using specific PCR primer combinations to show integration into the genome (Fig. 4B; Supporting Information Table S1). In recombinant parasites, binding to CD36 was reduced by at least 50% upon removal of the stabilizing drug TMP indicating a significant role of PfVAP1 in this process (Fig. 4C). FCR3 transfected with the empty vector in the presence of TMP was comparable with the wild type, indicating no effects of drug treatment on binding (data not shown).

Western blot analysis of the HA-tagged FCR3Pfvap1-DD in the presence or absence of TMP using anti-HA antibodies shows that protein expression was abrogated in the conditional protein knockdown (Fig. 4D). However, since minimal amounts of protein still may mediate residual adhesion, we generated knockout FCR3Pfvap1Δ parasites by double homologous recombination (Fig. 5A; Supporting Information Table S1) replacing the entire genomic sequence of Pfvap1 with hDHFR using the recently developed CRISPR-Cas9 system (Ghorbal et al., 2014). Knockout parasites emerged 3 weeks after transfection. Cultures were treated with Ancotil to remove episomes and two independent clones (B6 and C7) were selected for further analysis by PCR and sequencing to confirm the integration of the hDHFR cassette and the absence of the gene (Fig. 5A; Supporting Information Table S1). Binding to CD36 was reduced by 55% (B6) and 40% (C7). After two rounds of panning on CD36, both clones retained similar adhesion levels (40% for B6 and 50% for C7; Fig. 5B), suggesting that maximal binding inhibition attributable to PfVAP1 was reached and had been obtained in the FCR3 PfVAP1 knockdown parasites.

To ascertain that the observed decrease in binding levels in FCR3Pfvap1Δ were not due to off-target events, such as a concomitant deletion of KAHRP as has been reported for the complete rex1 knockouts in 3D7 (Dixon et al., 2011), we verified targeting of key proteins to different cellular compartments. Immunolabelling of FCR3Pfvap1Δ indicated that these parasites have KAHRP at the IE membrane and trafficking of PfEMP1 was comparable with FCR3 (Fig. 5C).

Trypsin resistance of PfEMP1 was reported in non-cytoadherent D10 parasites (Nacer et al., 2011). We investigated whether mutant FCR3Pfvap1Δ parasites show changes in trypsin sensitivity compared wild-type FCR3. The PfEMP1 of both B6 and C7 FCR3Pfvap1Δ clones were sensitive to trypsin digestion in a manner comparable with FCR3 as judged by the appearance of a shortened intracellular PfEMP1 fragment in Western blot analysis (Fig. 6).

Discussion

We have identified a novel gene implicated in the major virulence process of P. falciparum, the adhesion of IE to host endothelial cells. Previously, we had observed the complete loss of parasite adhesion despite apparent normal expression levels of PfEMP1 at the IE surface. Since this phenotype is linked to the deletion of a large subtelomeric fragment on chromosome 9 in several distinct parasites lines, our screen for candidate molecules focused on genes that were lost on the truncated chromosome 9 of D10 parasites. Our complementation strategy and screening for restoration of adhesion led to the discovery of a single candidate gene out of 12 tested, which showed a clear increase in the levels of adhesion (D10 = 12.3 IE mm⁻², D10Pfvap1-Ty1 = 255 IE mm⁻² and D10Pfvap1 = 293 IE mm⁻², i.e. 20-fold and 25-fold increase, respectively, for D10Pfvap1-Ty1 and D10Pfvap1). The screen revealed a protein that has never before been investigated or associated to parasite adhesion. Comparative genome analysis showed that Pfvap1 is only found in P. falciparum and despite its subtelomeric location adjacent to highly variable gene families, it is highly conserved between different parasite lines. The fact that this protein is exported together with PfEMP1 via Maurer’s clefts to the host membrane suggests that it has evolved to contribute in the process of functional cytoadhesion at the surface of IE.
PfEMP1 trafficking to the surface does not depend on the deleted genes in D10. Our observation that PfVAP1 complementation does restore the adhesion phenotype adds an important new piece to this complex mechanism. How PfVAP1 contributes to this process is still unknown but we speculate that it may occur at different levels. PfEMP1, after having been exported to the IE cytoplasm via the *Plasmodium* translocon of exported proteins, may need chaperones to refold into a functional adhesion molecule (Elsworth et al., 2014). Although the amino acid sequence of PfVAP1 does not reveal any homology to chaperones, it may interact directly with PfEMP1 to allow proper refolding. Alternatively, PfVAP1 and PfEMP1 may form a heterodimer at the surface of IE that stabilizes a PfEMP1 conformation needed for functional display to host receptors. A third

**Fig. 4.** Knockdown of PfVAP1 reduces binding to CD36.  
A. Schematic representation of the knockdown (KD) strategy. Primer locations are indicated by arrows and numbered (e.g. p1). Expected amplicons are shown for the different primer pairs used.  
B. Integration of the PfVAP1-DD-HA in FCR3. Genomic DNA was used for PCR with the indicated primer pairs. Genomic DNA from parasites obtained from a control transfection with empty plasmid is shown.  
C. Binding of FCR3<sup>PfVAP1-DD</sup> KD, FCR3<sup>PfVAP1-DD</sup> KD parasites were cultured in the presence or absence of the drug TMP that stabilizes the degradation domain. A 50% reduction in binding was observed in FCR3<sup>PfVAP1-DD</sup> parasites without TMP. Mean binding of infected erythrocytes (IE) to purified CD36 and BSA is shown. Error bars = ± standard deviation, n = 4 replicates). No effect of TMP on binding was detected using vector-transfected FCR3 parasites (data not shown). The inset shows Giemsa staining of FCR3<sup>PfVAP1-DD</sup> (+/− TMP) bound to CD36 or the BSA negative control showing the effect of PfVAP1 in adhesion.  
D. Western blot of FCR3<sup>PfVAP1-DD</sup> parasites cultured in the presence (+) or absence (−) of TMP probed with anti-HA. A band of the expected molecular weight (52 kDa) corresponding the PfVAP1-HA fusion protein is observed in the presence of TMP that is lacking in the absence of TMP, confirming the degradation of the protein.

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The possibility is that posttranslational modifications such as phosphorylation, which have been shown to reduce cytoadhesion, have been modified in PfVAP1 mutant parasites (Hora et al., 2009). Further experiments are needed to obtain in-depth insight into the role of PfVAP1 in surface adhesion. The possibility that PfVAP1 is exposed on the IE surface could not be confirmed experimentally as surface labelling with the Ty-1 antibodies did not result in surface signals using immunofluorescence assay (IFA) on live IE. Failure of these antibodies to label surface-exposed regions of PIVAP1 may be due to the location of the C-terminus at the inner IE membrane.

Our protein knockdown and gene knockout data demonstrate that mutant PIVAP1 FCR3 parasites show reduced binding to CD36 (between 50% and 60% reduction). This suggests that other genes located on chromosome 9 modulate the binding capacity in adherent parasites. A promising candidate is rex1, which encodes for a Maurer's cleft resident protein, and which

Fig. 5. FCR3Pfvap1Δ knockout parasites show a reduction in cytoadhesion.
A. Diagram depicting the cas9/CRISPR strategy (Ghorbal et al., 2014) used for disruption of Pfvap1 in FCR3 parasites. Only the pL7 plasmid carrying the Pfvap1-specific guide RNA and homology boxes is shown. The position of primers is shown (arrows) and each primer (p) is numbered. The expected amplicon sizes for PCR verification are shown. Gene replacement with hDHFR was verified by PCR (left). In FCR3Pfvap1Δ parasites, part of the 5′UTR sequence is lost (p3), the absence of this sequence used to select two KO clones (right).
B. Binding of FCR3Pfvap1Δ knockout clones panned on CD36. A 40% reduction in binding was observed in FCR3Pfvap1Δ clone B6 (dark grey). Binding to CD36 was reduced by 50% in the C7 clone (light grey). Mean binding of IE mm−2 to purified CD36 and BSA is shown. Error bars = ± standard deviation, n = 3 replicates.
C. FCR3Pfvap1Δ parasites export PfEMP1 and KAHRP in a manner comparable with wild-type FCR3 parasites, suggesting that there is no impairment or retention of these proteins in mutant parasites. Scale bar = 5 μm.
is also located on chromosome 9. Indeed, disruption of REX1 results in a partial loss of binding to CD36 (Dixon et al., 2011). Unlike the rex1 knockout or Hsp101 protein knockdown (Beck et al., 2014), PIEMP1 appears to be correctly trafficked in FCR3\(^{Pfvap1\Delta}\) knockout parasites.

Genes located on the right arm of chromosome 9 have also been implicated in gametocytogenesis and a number of them are predicted to be expressed in gametocytes, a stage that loses the expression of PIEMP1. Various transcriptomic data sets (www.plasmodb.org) suggests that Pfvap1 is also transcribed in gametocytes suggesting that the protein could have distinct functions depending on the life-cycle stage during which it is expressed.

In conclusion, we have identified a novel and unique \(P. \text{falciparum}\) protein with a major role in cytoadhesion named PIVAP1. Further characterization of PIVAP1 is warranted to determine exactly how this protein functions and interacts with PIEMP1 to clearly elucidate its role in binding.

**Experimental procedures**

**Parasite cultures and transfections**

*Plasmodium falciparum* D10 (derived from FC27), FCR3 and 3D7 were cultured in RPMI 1640 (GibcoBRL, Life Technologies), 5% Albumax I (GibcoBRL, Life Technologies) supplemented with 0.2 mM Hypoxathine (C.C.pro GmbH) as described (Cranmer et al., 1997). All parasite lines were regularly selected by gelatine flotation (Plasmion) and tested for the presence of mycoplasma (Rowe et al., 1998). Parasites were enriched by gelatin flotation (Plasmion) and resuspended in binding medium (RPMI 1640 supplemented with 25 mM HEPES pH 6.8) at 5 \(\times\) 10^7 IE ml^-1. Assays were performed as described (Buffet et al., 1999) with 20 \(\mu\) l of purified CD36 (10 \(\mu\) g ml^-1) immobilized on plastic Petri dishes or culture flasks, overnight at 4°C. For binding assays, control spots were coated with 1% BSA (fraction V, Sigma). Binding assays dishes were fixed with 2% glutaraldehyde, Giemsa stained for 10 min and quantified by microscopy.

**Immunodetection**

A pool of human hyperimmune sera (PIAG, Bouharoun-Tayoun and Drulhe, 1992), guinea-pig anti-PIEMP1 acidic transmembrane segment (ATS) (Wickert et al., 2003), mAb89 anti-KAHRP (Taylor et al., 1987), mAb1C11 anti-PfHSP70 (Mattei et al., 1989), anti-PfSBP1 (Bilsnick et al., 2000) and mouse anti-PfExp1 (Günther et al., 1991) were used to detect parasite antigens. Tag-specific antibodies were mouse hybridoma supernatant Ty1 mAb BB2 (Bastin et al., 1996) and rabbit anti-HA (Abcam ab8110). Fluorochrome-conjugated secondary antibodies were purchased from Life Technologies. Alcaline phosphatase and horseradish peroxidase secondary antibodies were from Promega and GE Healthcare respectively.

**DNA constructs**

**Complementation.** The 6.4 kb expression vector was derived from pLN-ENR-GFP (Nkrumah et al., 2006) by replacing the ENR-GFP by selected genes (see Fig. 1). The genes were amplified by PCR from FCR3 genomic DNA (gDNA) and cloned into the FseI/AscI sites of the pComp-BSD plasmid. Construction with the FseI/AscI sites of the pComp-BSD plasmid. Construction with

**Conditional protein knockdown.** The pGDB2-WR plasmid was generated from pGDB (Muralidharan et al., 2011) by insertion of a HDHFR cassette to use with parasites susceptible to WR99210 or TMP. FCR3 gDNA amplified by PCR was cloned into the
AfII/XhoI sites and transfected FCR3 parasites selected by adding BSD. Integration was verified by PCR and clones obtained by limiting dilution.

Knockout of PfVAP1. The cas9/CRISPR genome editing system recently adapted to P.falciparum was used as described (Ghorbal et al., 2014) to knockout the PfVAP1 gene. In brief, homology regions in the 5′ and 3′ UTRs were selected to ensure complete replacement of the coding sequence and cloned into the pL7 plasmid on either side of the hDHFR selectable marker. The 555 bp 5′ UTR homology region was amplified from FCR3 DNA and cloned between the SpeI/AflII sites of the pL6 plasmid by In-Fusion reaction (In-Fusion HD cloning kit Clontech Laboratories). Similarly, the 477 bp 3′ UTR homology region was amplified by PCR and cloned into the EcoRI/NcoI sites. Single gRNA sequences were selected generating two plasmids. The first sequence corresponded to 20 bp in PfVAP1 coding sequence (5′-ACATATTAGTTTCAACC-3′) located at the beginning of the second exon (ATG + 327 bp), approximately halfway between the two homology regions. The second gRNA sequence (5′-TTTGTAGGTGTAACATAAC-3′) was selected towards centre of the coding sequence (ATG + 539 bp). After transfection with the two plasmids, parasites were obtained in 16 days, in two independent transfections. Parasites were treated with Ancotil to remove residual episomes, and were cloned by limiting dilution. After PCR verification, two FCR3 PfVAP1 clones (B6 and C7) were selected for further phenotypic studies.

Northern and Southern blots

Total RNA from 3D7, D10 and FCR3 was isolated and processed as previously described (Kyes et al., 1999; 2000). TRIzol (TRIzol reagent, Ambion Life Technologies) was added to harvested ring-, trophozoite- and schizont-IE. Parasite DNA was extracted (Gene Elute Mammalian Genomic DNA mini prep kit, Sigma-Aldrich) digested with Drai/Xmnl (BioLabs), and processed (Viebig et al., 2005). Nucleic acids were transferred onto Hybond N + (Amersham) and hybridized with a 32P-labelled PfVAP1 probe (Megaprime Labelling System, GE Healthcare). Membranes were washed with 2X sodium saline citrate (SSC), 0.1% sodium dodecyl sulfate (SDS) at 60°C and exposed with BioMax MS films and Transcreen HE (Kodak) at −80°C.

Parasite protein preparation

For the detection of tagged PfVAP1, IE were collected and saponin-lysed. Parasites were washed three times in PBS containing protease inhibitors and the pellet resuspended to obtain 10^6 IE µl−1. Samples were run on 10% Tris-Bis gels (BioRad) in MOPS Buffer (BioRad) and transferred onto nitrocellulose membranes (iBlot, Life Technologies). Membranes were blocked in 5% milk/0.05% tween 20 (Promega) in PBS (PBS/milk/tween) for 30 min at room temperature and incubated overnight with either mouse anti-Ty1 (1:100) or rabbit anti-HA (1:250). Following three washes with PBS/milk/tween, alkaline phosphate-conjugated secondary antibodies were added for 1 h at room temperature.

Trypsin cleavage assays

Infected RBCs were enriched by gelatin flotation and incubated for 15 min with either 0, 10 or 100 µg ml−1 of trypsin (Baruch et al., 1996). Following three washes in PBS containing protease inhibitors, cells were either incubated with pooled hyperimmune serum (1:5) or proteins extracted with 1% Triton X-100 (v/v) followed by 2% SDS. Extracts were resuspended in sample buffer, heated, separated on 10% Bis-Tris gels (BioRad), and transferred onto nitrocellulose membranes (iBlot, Life Technologies). Detection of PfEMP1 was performed by incubation with a guinea pig anti-ATS antibody (1:400) followed by guinea pig alkaline phosphatase (AP)-conjugated secondary antibody. Equal protein loading was verified with the mAb1C11 anti-Hsp70 antibody (1:2000) and revealed with NBT/BCIP following incubation with an anti-mouse AP-conjugated secondary antibody.

qRT-PCR

To confirm the intraerythrocytic gene expression profile of PfVAP1, parasites were synchronized by plasmion and sorbitol to obtain a 5 h window. Parasites were harvested every 12 h during the course of a single cycle. Total blood was washed once in PBS and TRIzol immediately added. Samples were then stored at −80°C until further processing. RNA was extracted, precipitated, treated with DNase (Ambion), reverse transcribed (SuperScript II Reverse Transcriptase – Life Technologies), and diluted to 300 ng µl−1 in nuclease-free water for qRT-PCR. Expression was analysed using the primers Exon1-F: 5′-GT TTCTTTATTTTCTTACTAC-3′; Intron-R: 5′-AAAGCACTTTTCTATAC-3′; Intron-F: 5′-CCCTATCCAAATGTAAC-3′; Intron-R: 5′-GCTATATTTTTGTAGAT3′. Control genes used were seryl transferase (Salanti et al., 2003), inositol (Mancio-Silva et al., 2013) and aldolase (Knapp et al., 1990).

Immunofluorescence

PfVAP1 was localized using anti-Ty1 or anti-HA tag antibodies. Parasites were spotted onto multiwell slides and air-dried (Hinterberg et al., 1994). To further define the localization of PfVAP1, double labelling IFA with PfEMP1 anti-ATS or rat anti-PfSBP1 Maurer’s clefts markers were performed as already described (Nacer et al., 2011).

Flow cytometry

Parasites were synchronized (within a 5 h window) by plasmion and sorbitol and 30–35 h post-invasion schizonts were incubated with the human sera for 1 h at room temperature. After three washes with PBS, samples were reacted 1 h with anti-human Alexa 488 (1:100) (Life Technologies). Nuclei were stained 10 min with DAPI, washed, and the samples were diluted to 500 cells µl−1 for reading by flow cytometry (Fortessa). For the quantification of CD36 labelling, parasites were obtained as described above and incubated for 1 h at RT with 10 µg ml−1 chimeric CD36-human IgG in suspension. Cells were washed in PBS prior to the addition of mouse anti-human IgG Alexa 488-conjugated secondary antibody. Nuclei were stained with SYTO 61 (Life Technologies) and samples were analysed on a Guava flow cytometer (BD Biosciences). Data were compiled and analysed using FlowJo (version 10).

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Conflict of interest
The authors declare no conflict of interest.

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**Supporting information**

Additional Supporting Information may be found in the online version of this article at the publisher’s web-site:

**Fig. S1.** PIVAP1 and PIEMP1 co-localize in the Maurer’s clefts in complemented D10 parasites. Ty1-tagged D10*PfVAP1* parasites were labelled with anti-Ty1 monoclonal antibody (red), guinea pig anti-PIEMP1 (green), and rat anti-PISBP1 (magenta), nuclei were stained with DAPI (blue). PIVAP1 associates with PIEMP1 in Maurer’s clefts (PISBP1). Scale bar = 5 µm.

**Table S1.** List of oligonucleotides used in this study.