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Multiple charged amino acids of *Plasmodium falciparum* SURFIN$_{4.1}$ N-terminal region are important for efficient export to the red blood cell

Ben-Yeddy Abel Chitama$^{a,b,1}$, Shinya Miyazaki$^{b,c,1}$, Xiaotong Zhu$^{b,d,1}$, Wataru Kagaya$^{b,e,f}$, Kazuhide Yahata$^b$, Osamu Kaneko$^{a,b,*}$

$^a$ Leading Program, Graduate School of Biomedical Sciences, Nagasaki University, 1-12-4 Sakamoto, Nagasaki 852-8523, Japan
$^b$ Department of Protozoology, Institute of Tropical Medicine (NEKKEN), Nagasaki University, 1-12-4 Sakamoto, Nagasaki 852-8523, Japan
$^c$ Leiden Malaria Research Group, Parasitology, Leiden University Medical Center (LUMC), Leiden, The Netherlands
$^d$ Department of Immunology, College of Basic Medical Science, China Medical University, Shenyang, Liaoning 110122, China
$^e$ Department of Environmental Parasitology, Graduate School of Tokyo Medical and Dental University, Tokyo Medical and Dental University, 1-5-45 Yushima, Bunkyo, Tokyo 113-0034, Japan
$^f$ Department of Parasitology, Graduate School of Medicine, Osaka City University, 1-4-3 Asahimachi, Abeno, Osaka 545-8585, Japan

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ABSTRACT

*Plasmodium falciparum*, an obligate intracellular protozoan parasite which causes the severe form of human malaria, exports numerous proteins to the infected red blood cell that are important for its survival and of severe pathological effect to its host. These proteins and their export mechanisms are candidates for drug and vaccine development, and among them is the *Plasmodium* SURFIN family of proteins. Previously we showed that the N-terminal region along with the sequence surrounding the transmembrane domain of SURFIN$_{4.1}$ is essential for its export to Maurer’s clefts in the red blood cell cytoplasm. We proposed that this region is recognized by a machinery responsible for protein translocation across the parasitophorous vacuole membrane surrounding the parasite. To understand the export mechanism further, we utilized a fluorescent protein-tagged mini-SURFIN$_{4.1}$ consisting of the minimum essential components for export. Alanine scanning of all charged amino acids within the N-terminal region revealed that replacement of 3 glutamic acid and 2 lysine residues significantly impairs the export efficiency of this protein across the parasitophorous vacuole membrane surrounding the parasite. In addition, N-terminally Myc-tagged mini-SURFIN$_{4.1}$ and mini-SURFIN$_{4.2}$ with similar architectures were detected with anti-Myc antibody at Maurer’s clefts, indicating that elements required for export to Maurer’s clefts are conserved between SURFIN$_{4.1}$ and SURFIN$_{4.2}$, and that N-terminal sequences of these SURFIN members are not cleaved during export. Our results implicate a conserved nature of SURFIN export to the red blood cell, particularly an important role of multiple glutamic acid and lysine residues in the SURFIN N-terminal region.

1. Introduction

Malaria remains a barrier to the economic progress of developing countries, particularly due to *Plasmodium falciparum*, the causative agent of the most severe form of malaria [1]. Of the total malaria cases occurring worldwide, half occur in sub-Saharan African, which also contains a majority of the least developed countries [2]. Malaria is a disease of the red blood cell (RBC), and the pathology is largely due to parasite-encoded proteins that the parasite exports to the RBC cytoplasm and surface [3]. These proteins are involved in RBC remodeling essential to parasite survival, while consequently mediating pathophysiological reaction in the host [4]. In trafficking to the RBC surface majority of these proteins pass through parasite-generated membranous structures called Maurer’s clefts which act as a sorting platforms within the RBC cytoplasm [5].

A conserved N-terminal region sequence, termed the *Plasmodium* EXport EElement (PEXEL) or Vacuolar Targeting Sequence (VTS) motif, was shown to be essential for export of a diverse group of proteins to the RBC cytoplasm. The PEXEL/VTS motif is a pentameric amino acid motif with the consensus sequence RxLxE/Q/D; and identifies a...
reperoire of over 300 Plasmodium predicted exported proteins, termed the exportome [6–8]. During transport within the endoplasmic reticulum (ER), this motif is cleaved by the ER-resident protease plasmin V [9,10]. The exposed amino acid residues (xE/Q/D) after cleavage are essential for export to the RBC cytoplasm [8]. Many studies have identified a second group of exported proteins which lack obvious PEXEL/VTS motifs; and are termed PEXEL negative exported proteins (PNEPs). Examples include SBP1, REX1, REX2, MAHRP1, MAHRP2, MSR6P, and MSR7P [11–16]. The N-terminal regions of some PNEPs such as SBP1, REX2 and MAHRP2 were shown to be essential for export to the RBC [13,15,17], indicating that the N-terminal sequence requirement for RBC export is conserved in PNEPs [8]. The N-terminal regions of PNEPs are potentially cleaved by an unknown mechanism, as shown for REX2 [13]. However, it is not known whether N-terminal processing of PNEPs is required for export to the RBC.

SURFIN4.1 (Surface-associated interspersed protein 4.1) is a PNEP exported to the Mauer’s clefts and is encoded by the surf gene family consisting of 10 members in P. falciparum [18,19]. The N-terminal region (18 amino acids (aa)), transmembrane domain (23 aa), and following short cytosolic region (17 aa) of SURFIN4.1 contain sufficient information for export to Mauer’s clefts [20]; however, specific amino acid residues in the N-terminal region that contribute to this export have not been evaluated. Herein we report the presence of five charged amino acid residues (3 glutamic acid and 2 lysine residues) in the N-terminal sequence of SURFIN4.1 that independently contribute to efficient export of recombinant mini-SURFIN4.1 to Mauer’s clefts. We also show that elements required for export of SURFIN4.1 are conserved in the paralog SURFIN4.2, for which infected RBC (iRBC) surface exposure has been shown [18,21]. We additionally determined that the N-terminal sequences of SURFIN4.1 and SURFIN4.2 are not processed during export to Mauer’s clefts.

2. Materials and methods

2.1. P. falciparum cultivation and transfection

P. falciparum MS822 and 3D7S8 lines were cultured basically as described [20,22]. The parasites were maintained with O– RBCs at 2% hematocrit in RPMI-1640 medium supplemented with 5% heat-inactivated pooled type AB+ human serum, 0.25% AlbuMaxI (Invitrogen, Carlsbad, CA), 200 mM hypoxanthine (Sigma, St. Louis, MO), and 10 μg/mL gentamicin (Sigma).

Transfection of P. falciparum was performed as described [23]. Briefly, 100 μg of plasmids were electroporated to uninfected RBCs (Gene Pulser Xcell; Bio-Rad, Hercules, CA), mixed with P. falciparum-iRBCs and cultured for 3 days without drug. At day 4, 5 nM WR99210 (a gift from Dr. D. Jacobs) was added to the culture for the selection of drug-resistant transfectants. The MS822 line was used to evaluate Myc-mini-SURFIN4.1 and Myc-mini-SURFIN4.2. After transfectants were obtained, the concentration of WR99210 was gradually increased to 20 nM before analysis.

2.2. Plasmid construction

Plasmids were constructed using the Multisite Gateway system (Thermo Fisher Scientific, Waltham, MA) and In-Fusion cloning system (Takara Bio Inc., Japan). The pENT2 plasmids for the Ala replacement (pENT12_SURFIN4.1N4-D34A-T-cyt, pENT12_SURFIN4.1N4-K42A-T-cyt, pENT12_SURFIN4.1N4-D40A-T-cyt, pENT12_SURFIN4.1N4-K49A-T-cyt, pENT12_SURFIN4.1N4- D50A-T-cyt) were modified based on a plasmid to express SURFIN4.1N4-T-cyt [20] by site-directed mutagenesis using the KOD -Plus-Mutagenesis Kit (Toyobo) with oligonucleotide primers listed in supplementary Table 1. The expression plasmids of N-terminal Ala-replaced mini-SURFIN4.1 were constructed by the Gateway Multisite LR recombination reaction with pENT4/1-JpCRT5’ (as a promoter component), pENT23.3 × Ty-GFP, and pCHDR-3/4 plasmids [24]. The open reading frame sequences of all plasmids were confirmed before transfection.

For Myc-mini-SURFIN4.1, pENT12_SURFIN4.1N4-T-cyt was constructed from a plasmid to express SURFIN4.1N4-T-cyt [20] by inserting a DNA fragment encoding double Myc tags using a KOD -Plus-Mutagenesis Kit. For Myc-mini-SURFIN4.2, a DNA fragment encoding the N-terminal region was PCR-amplified from P. falciparum genomic DNA and introduced into the EcoRV site of pB13 plasmid to yield pB13_SURFIN4.2N. pB13 has been made from pB12 [25] by replacing the sequence between Nhel and PstI with the oligonucleotides CATGCAGATCTCTAGTGAGCCGCTCTGCA and GAGGCTACTAGTAGTACTGCTG and by modifying the attB site to attB3 site by site-directed mutagenesis with the oligonucleotides CAAAATGCTGAGCCGCTCTGCA and TATAAAATGGTGTTACTGAGCAGCCTGCA. SURFIN4.2 sequence spanning from the variable region 2 (Var2 region) to a part of the cytoplasmic region was amplified and inserted into the XbaI site of pB13_SURFIN4.2N to yield pB13_SURFIN4.2N. A framshift was then corrected, and the intron and Var2 region were removed, to yield pB13_SURFIN4.2N-T-C. pB13_SURFIN4.2N2Myc-N-T-C was constructed by inserting a DNA fragment encoding double Myc tags and subjected to the Gateway BP reaction followed by a Gateway Multisite LR recombination reaction with pENT41-JpCRT5’ and pCHD43(III) [26].

2.3. Live cell imaging and data analysis

To capture live green fluorescent protein (GFP) signals from transgenic parasites, parasites were synchronized by the 5% D-sorbitol method [27] and allowed to grow until the trophozoite stage. Images were visualized using a fluorescence microscope (Axio Imager Z2; Carl Zeiss, Germany) with a 100×/1.40 oil-immersion lens and captured with a charge-coupled device camera (AxioCam MRm; Carl Zeiss). Images were processed using Adobe Photoshop CS3 (Adobe systems Inc., San Jose, CA).

To quantify the intensity of signals located in the iRBC cytoplasm, 20 images were selected for each transfectant. Area and mean values of the signal were measured from the entire parasite-iRBC (signals from the parasite and in the iRBC cytoplasm), signals in the iRBC cytoplasm, and signals from the nearby external area surrounding the parasite (background) using Fiji [28]. Background signal mean values were subtracted from the test mean values and multiplied by the area values to produce integrated density values for each test area. The proportion of the signals in the iRBC cytoplasm was obtained by dividing integrated signal density values in the iRBC cytoplasm by those from the entire parasite-iRBC. Statistical differences of the proportion of the signals in the iRBC cytoplasm were evaluated by Kruskal-Wallis one-way analysis of variance followed by Dunn’s multiple comparison test using GraphPad Prism6 (GraphPad, San Diego, CA) and graphs were drawn using Microsoft Excel 2016.

2.4. Indirect immunofluorescence assay (IFA)

Thin blood smears on glass slides were air-dried and stored at ~80 °C. Retrieved blood smears were fixed with 4% paraformaldehyde/0.075% glutaraldehyde at room temperature for 30 min, followed by quenching with 50 mM glycine for 15 min at room temperature. The slides were blocked with 10% normal goat serum (Invitrogen) at 37 °C for 60 min and probed with mouse anti-GFP monoclonal antibody (1:250; 3E6, Invitrogen), mouse anti-Ty1 monoclonal antibody (1:500; MAb-054-050; Diagenode, Belgium) or mouse anti-Myc monoclonal antibody (1:1000; 9B11, Cell Signaling Technology, Danvers, MA), and rabbit anti-sBP1 serum (1:2000; a kind gift from T. Tsuboi). After washing with phosphate-buffered saline (PBS), the smears were incubated with PBS containing Alexa Fluor® 488-conjugated goat anti-mouse IgG antibody and Alexa Fluor® 594-conjugated goat anti-rabbit.
IgG antibody (1:500, Invitrogen). Parasite nuclei were stained with 4′,6-diamidino-2-phenylindole (DAPI). Images were captured using a fluorescence microscope and camera and processed as described above.

2.5. Western blotting

Parasite-iRBCs were exposed to 5% D-sorbitol at hours 0 and 6, then harvested at hour 30 to obtain synchronized parasite material. Mature trophozoite/schizont-iRBCs were treated with 0.15% saponin in PBS containing a mixture of protease inhibitors (PI; cOmplete, EDTA-free; Roche, Basel, Switzerland), and the pellets after centrifugation were washed with PBS-PI, and proteins were dissolved in SDS-loading buffer with 2% 2-mercaptoethanol. After passing through a 27G needle to shear parasite DNA, samples were centrifuged and supernatants were collected. Protein from an estimated 1.0 × 10⁵ parasites was subjected to electrophoresis on 5–20% SDS-polyacrylamide gradient mini gels (ATTO, Japan). The protein bands were transferred from gels to PVDF membranes (Millipore, Billerica, MA). The membranes were then probed with mouse anti-Ty1 monoclonal antibody (1:500) or anti-Plasmodium berghei HSP70 mouse monoclonal antibody (obtained from J. Sattabongkot, which cross-reacts with HSP70 orthologs of other Plasmodium spp) [29] followed by a secondary incubation with horse-radish peroxidase (HRP)-conjugated goat anti-mouse IgG antibodies (Promega, Madison, WI). Bands were visualized with Immobilon Western Chemiluminescent HRP substrate (Millipore) and detected using a chemiluminescence detection system (LAS-4000EPUVmini; Fujifilm, Japan). The relative molecular sizes of the proteins were calculated based on molecular size standards (Precision Plus Dual Colour Standards; Bio-Rad).

3. Results

3.1. Identification of SURFIN₄₋₁ N-terminal amino acid residues that are important for efficient export to the iRBC cytoplasm

To evaluate the influence of charged amino acid residues within the SURFIN₄₋₁ N-terminal region on export to the iRBC cytoplasm, we utilized sequences consisting of amino acid residues 34–50 (termed N₄ region), which was found to contain sufficient information for export to Maurer’s clefts [20]. Initially we used the MS822 parasite line to express a panel of recombinant mini-SURFIN₄₋₁ proteins consisting of the N₄ region fused to the transmembrane domain and a short cytoplasmic region (N₄TC). In these proteins, six negatively charged amino acid residues (D₃₄A, D₃₇A, E₃₈A, D₄₀A, E₄₅A, and E₅₀A) and three positively charged amino acid residues (K₃₅A, K₄₂A, and K₄₉A) were independently replaced to Ala (Fig. 1A). To monitor protein export we fused a Ty1 peptide tag and GFP to the C-terminus (Fig. 1A). Western blot analysis confirmed that all recombinant mini-SURFIN₄₋₁ proteins
were expressed in a similar quantity and fashion with a few extra smaller bands of possible proteolysed products (Fig. 1B). Live cell imaging for MS822-background transfectants expressing modified SURFIN_{4,1}N4-T-Cyt (N4TC) protein. Scale bar = 5 μm. (C) Quantification of signal in the iRBC cytoplasm. Significance was evaluated by Kruskal-Wallis one way analysis of variance followed by Dunn’s multiple comparison test (p values are indicated). Twenty infected RBCs were evaluated for each line. (D) Amino acid residues for which Ala replacement made a significant loss of signal export are shown with enlarged letters. Blue and red letters indicate negatively and positively charged amino acids, respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Fig. 2. The effect of Ala replacements within the SURFIN_{4,1} N-terminal region for export in the P. falciparum line 3D7S8. (A) Western blot of N4TC variants with mouse anti-Ty1 antibody. HSP70 was used as a loading control marker. Arrow indicates the expected size of N4TC protein. (B) Live cell imaging of MS822-background transfectants expressing modified SURFIN_{4,1}N4-T-Cyt (N4TC) protein. Scale bar = 5 μm. (C) Quantification of signal in the iRBC cytoplasm. Significance was evaluated by Kruskal-Wallis one way analysis of variance followed by Dunn’s multiple comparison test (p values are indicated). Twenty infected RBCs were evaluated for each line. (D) Amino acid residues for which Ala replacement made a significant loss of signal export are shown with enlarged letters. Blue and red letters indicate negatively and positively charged amino acids, respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

were expressed in a similar quantity and fashion with a few extra smaller bands of possible proteolysed products (Fig. 1B). Live cell imaging for MS822 line transfectants revealed that E38A, K42A, E45A, K49A and E50A showed sparse signal in the iRBC cytoplasm (Fig. 1C) and had significantly lower signal in the iRBC cytoplasm per total signal compared to N4TC which served as a background (p < .0001; Fig. 1D). Signal in the iRBC cytoplasm of D34A, K35A, D37A, and D40A replacements were similar to N4TC (Fig. 1C) and no significant difference was detected for the signal in the iRBC cytoplasm per total signal compared to N4TC (Fig. 1D).

To confirm these results we further examined the influence of amino acid residues in the N4 region using 3D7S8-background transfectants; focusing in part on amino acid residues that by the replacement to Ala showed significant reduction of the signal in the iRBC cytoplasm per total signal in MS822-background. We used two different lines to add more confirmatory value to our findings. Western blot analysis indicated that all generated recombinant mini-SURFIN_{4,1} proteins were expressed with a similar quantity and fashion with a few extra smaller bands of possible proteolysed products (Fig. 2A). Live imaging indicated that D37A, for which reduction was not observed in the MS822-background transfectant, showed signals in the iRBC cytoplasm similar to N4TC (Fig. 2B) and the signal in the iRBC cytoplasm per total signal was not significantly different from that of N4TC (Fig. 2C). Five replacements that reduced the export efficacy in MS822-background (E38A, K42A, E45A, K49A, and E50A) again showed sparse signal in the iRBC cytoplasm (Fig. 2B) and the ratios of the iRBC cytoplasm per total signal was significantly lower than that of the 3D7S8-background N4TC parasite (Fig. 2C). Amino acid residues for which Ala replacement resulted in a significant loss of signal export to iRBC cytoplasm in both MS822 and 3D7S8 parasite lines are shown in the Fig. 2D.
3.2. N-terminal sequences of SURFIN4.1 protein was not cleaved during export to Maurer’s clefts

To assess whether the SURFIN4.1 N-terminal sequence is cleaved during export to Maurer’s clefts, we generated a transfectant expressing N-terminally Myc-tagged and C-terminally Ty1 and GFP-tagged mini-SURFIN4.1 consisting of a short N-terminal region, transmembrane domain, and the adjacent short cytosolic region (Fig. 4A). Both anti-Ty1 and anti-Myc antibodies detected a punctate pattern of expression in the iRBC cytoplasm which co-localized with the Maurer’s cleft marker protein SBP1 (Fig. 4B), indicating that the N-terminal sequence of SURFIN4.1 is not processed during export to Maurer’s clefts.

3.3. Regions required for export to Maurer’s clefts are conserved between SURFIN4.1 and SURFIN4.2

To determine if regions of SURFIN4.1 required for export to Maurer’s clefts are conserved in another SURFIN family member, we generated, termed Myc-mini-SURFIN4.2, a transfectant expressing N-terminally Myc-tagged and C-terminally Ty1 and GFP-tagged mini-SURFIN4.2 whose composition is the same as Myc-mini-SURFIN4.1; specifically, containing the N-terminal region (51 aa), transmembrane domain (23 aa) with the adjacent short cytosolic region (20 aa) (Fig. 4A). IFA with anti-Ty1 or anti-Myc antibodies showed that both signals co-localized with SBP1 (Fig. 4C), indicating that Myc-mini-SURFIN4.2 is exported to Maurer’s clefts and at that location Myc-mini-SURFIN4.2 retains the N-terminal Myc sequence. This result indicates that the sequence information for export to the RBC is conserved between SURFIN4.1 and SURFIN4.2 and confirms that the processing of the N-terminal sequence of SURFIN is not required for this process.

4. Discussion

Protein export to the iRBCs plays an essential role for malaria parasite development and proliferation within the RBC and consequently virulence in the human host. In this study we demonstrate the following: i) at least five charged amino acid residues in the N-terminal region of SURFIN4.1 (E38, K42, E45, K49, and E50) are involved in its efficient export to the iRBC cytoplasm, ii) regions required for export are conserved between SURFIN4.1 and SURFIN4.2 and iii) processing of the N-terminal region of SURFIN4.1 and SURFIN4.2 is not required for export to the iRBC cytoplasm.

The importance of the charged amino acids for export of P. falciparum-encoded proteins to the iRBC cytoplasm has been shown in other studies; for example, the role of Glu has been also observed for export of another PNEP, REX2, whereby mutation of Glu at position 7 (MKMYLAIFSSGK) was sufficient to abolish export of this protein to the iRBC cytoplasm [13]. Ala replacement of Glu residues regarding export of PHsp70x (SNNAEES) was also found to be essential as it led to the accumulation of this protein in the parasitophorous vacuole (PV) [30]. In the case of SBP1, Ala replacement of the N-terminal DEPTQLDAVP (amino acid positions 16–26), which contains three negatively charged residues, stopped export of this protein [17]. A net negative charge of the N-terminal region at a neutral pH was proposed to be important for SBP1 export to Maurer’s clefts (pl of N-terminal 35 amino acid residues of SBP1 is 4.50) [17]. The pl of the N-terminal 50 amino acid residues of SURFIN4.1 and N-terminal 18 amino acid residues of mini-SURFIN4.1 N4 region are 4.27 and 4.30, respectively. Ala replacement at E38, E45, and E50 in the mini-SURFIN4.1 N4 region all only slightly increased the pl (4.44) but had a strong negative impact on export. We found that K42A and K49A, both of which decreased the pl to 4.02, also reduced the protein transport efficacy, suggesting that rather than a net negative charge the correct positioning of negatively and positively charged amino acid residues are critical for mini-SURFIN4.1 export to the iRBC cytoplasm. The structure was recently resolved for a translocon on the parasitophorous vacuole, termed Plasmodium translocon of exported proteins (PTEX) [31]. Because PTEx translocates both PEXEL-containing proteins and PNEPs [32,33], these charged amino acid residues may interact directly with PTEx or may be responsible for a correct conformation to be recognized by this translocation machinery. Future computational modeling of the interaction between PTEx and mini-SURFIN4.1 may provide insight into how SURFIN4.1 and PNEPs are recognized by the translocon.

Marti et al. (2004) demonstrated Maurer’s cleft transport of a mini-PiEMP1 consisting of the N-terminal aa 1–32 (MGPPSTAPDYSS-AKDKEELDGGQVHDKVK, “PEXEL-like” residues are underlined)
Fig. 4. Myc-tag fused to the N-terminus of mini-SURFIN$_{4.1}$ and mini-SURFIN$_{4.2}$ proteins are detected in Maurer's clefts. (A) Schematics of Myc-mini-SURFIN$_{4.1}$ and Myc-mini-SURFIN$_{4.2}$ proteins. Myc tag sequences are underlined with red colour and predicted transmembrane domain (T) are boxed. N-terminal regions (N, 49 aa) of SURFIN$_{4.1}$ and SURFIN$_{4.2}$ were used. (B and C) IFA images of the transfectants expressing Myc-mini-SURFIN$_{4.1}$ (B) or Myc-mini-SURFIN$_{4.2}$ (C) with anti-Ty1 or anti-Myc antibodies co-stained with anti-SBP1 Maurer's cleft marker antibody. Scale bar = 5 μm. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
combined with the transmembrane and cytoplasmic domains (aa 1913–2360) of PEMP1 (PF3D7_1240600). This is a similar domain composition to our mini-SURFIN proteins. For the mini-PEMP1 the simultaneous Ala replacement of K7, I19, and D32 (MGPSTPDYSS-AKDAEALAKIGQVHDVKKV), which alters the PEXEL-like motif, abolished export [34]. Because PEMP1 does not possess an N-terminal ER-signal sequence, the proposed “PEXEL-like” motif is likely recognized by PTEX in a similar fashion to our mini-SURFIN proteins. If this is the case, other charged amino acids in the PEMP1 N-terminal region (K14, D15, E18, K22, D28, K30 and K42) may also be critical for its export. We speculate that the “PEXEL-like” motif in the PEMP1 N-terminal is not cleaved, based upon the lack of processing of the N-terminal SURFIN-derived sequences in mini-SURFINS.

Because we focused on the signal detected within Maurer’s clefts in the iRBC cytoplasm, we did not assay possible inhibitions at prior trafficking steps, such as the stability of the proteins in their transit from the ER to incorporation in Maurer’s cleft structures. IFA signal can be seen around the parasite nucleus even in the transfecant with parental construct N4Tc (e.g. Fig. 2A), suggesting that a portion of the recombiant protein is retained in the ER at varying levels in all transfecants. However, all transfecants, including five transfecants with reduced signal in the iRBC cytoplasm, showed IFA signal surrounding the parasite, indicating that proteins were able to reach the parasite plasma membrane. We interpret that translocation across the parasitophorous vacuole membrane is disturbed and mini-SURFIN1,4 were retained on the parasite plasma membrane in the five transfecants with reduced signals in the iRBC cytoplasm, likely due to insufficient recognition by PTEX.

Mini-SURFIN1 and mini-SURFIN4.2 proteins containing N-terminal Myc tags were detected within Maurer’s clefts, indicating that N-terminal processing of SURFIN sequence is not required for export of these proteins to the iRBC cytoplasm. To our knowledge this is the first report with unambiguous data showing that the N-terminal region is not cleaved during this process for PNEPs. Thus, although another PNEP, REX2, was shown to be processed at its N-terminal region like PEXEL-dependent proteins [13], the N-terminal processing is clearly not a general feature of PNEPs.

In summary, we characterized the N-terminus of P. falciparum P. falciparum and P. knowlesi, which are associated with the erythrocyte skeleton, Mol. Biochem. Parasitol. 156 (2008) 1–10, https://doi.org/10.1016/j.molbiopara.2008.06.005.


25. K. Deitsch, C. Driskill, T. Wellemes, Transformation of malaria parasites by the


