An upstream open reading frame (uORF) signals for cellular localization of the virulence factor implicated in pregnancy associated malaria

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ABSTRACT

*Plasmodium falciparum*, the causative agent of the deadliest form of human malaria, alternates expression of variable antigens, encoded by members of a multi-copy gene family named var. In var2csa, the var gene implicated in pregnancy-associated malaria, translational repression is regulated by a unique upstream open reading frame (uORF) found only in its 5′ UTR. Here, we report that this translated uORF significantly alters both transcription and posttranslational protein trafficking. The parasite can alter a protein’s destination without any modifications to the protein itself, but instead by an element within the 5′ UTR of the transcript. This uORF-dependent localization was confirmed by single molecule STORM imaging, followed by fusion of the uORF to a reporter gene which changes its cellular localization from cytoplasmic to ER-associated. These data point towards a novel regulatory role of uORF in protein trafficking, with important implications for the pathology of pregnancy-associated malaria.

INTRODUCTION

*Plasmodium falciparum*, the protozoan parasite that causes human malaria remains one of the deadliest pathogens on earth (1). This parasite, transmitted by female Anopheles mosquitoes, causes malaria when it reaches the blood stream and proliferates within infected red blood cells (iRBC). In addition to anemia, the severe pathology caused by *P. falciparum* is linked to its ability to modify the infected erythrocyte with proteins that cause the iRBC to adhere to neighboring uninfected RBCs as well as to several endothelial receptors (2). The main ligand responsible for the cytoadhesive properties of the iRBC are variable surface proteins named PfEMP1 encoded by a multi-copy gene family named var (3). To avoid exposure of its entire antigenic repertoire, the parasite expresses only a single var gene at a time while the rest of the var gene family is transcriptionally silent (4,5). Due to the constant switching of var gene expression, which occurs intrinsically at a very low rate, some parasites avoid the antibody-mediated response against the predominant PfEMP1 and continue the infection, expressing another PfEMP1 variant that is not recognized by the immune system (6). Thus, immune evasion by *P. falciparum* is achieved both by antigenic switches between expression of different PfEMP1 variants and by avoiding clearance by the spleen through cytoadhesion to endothelial receptors and sequestration in deep tissues. It is well established that different PfEMP1 variants have different cytoadhesive properties since they bind different endothelial receptors such as CD36, ICAM1, Chondroitin Sulfate A (CSA) and EPCR (7,8). Thus, some of the severe disease manifestations, such as cerebral and pregnancy-associated malaria, are determined by the adhesion phenotype of the PfEMP1 variant expressed.

CSA is an abundant glycosaminoglycan expressed in the placental intervillous space and is bound by a specific PfEMP1 variant named VAR2CSA (9). The only var gene expressed in a parasite culture selected for binding to CSA is var2csa (10), and parasites in which this gene has been deleted do not adhere to CSA (11). Interestingly, people living in endemic regions that were exposed to continuous *P. falciparum* infections develop antibodies against most of the PfEMP1 repertoire. However, most studies indicate that antibodies against VAR2CSA can only be found in serum
taken from women who were infected during pregnancy and that protection against placental sequestration is acquired over consecutive pregnancies (9,12). Although var genes undergo continuous transcription switching, the PfEMP1 expressed by var2csa is only translated in pregnant women. Similar to other var genes, var2csa transcription is tightly regulated in a mutually exclusive manner, but contains an additional unique mechanism that represses protein translation when expression of VAR2CSA isn’t needed. This mechanism is regulated by an upstream open reading frame (uORF) which is found uniquely in the 5′ UTR of var2csa (13). This uORF is 360bp long and is situated between the transcription start site (TSS) and the downstream open reading frame (dORF) that encodes VAR2CSA. Therefore, the var2csa transcript is bicistronic, with the uORF and the dORF separated by an inter-cistronic region (ICR), and it appears that var2csa mRNA contains both ORFs (13). The molecular mechanism that allows translation of the dORF in the presence of a placenta is unknown, but there is strong evidence that it is controlled by a switch in the efficiency of translation re-initiation of the dORF (14), and expression of an endogenous plasmodium translation-enhancing factor (15). Here, we show that in addition to its role in regulating translation of the var2csa-encoded PfEMP1, this uORF plays a key role in determining the cellular localization of the dORF-encoded protein.

**MATERIALS AND METHODS**

### Parasite culture and parasitemia counts

All parasites used were derivatives of the NF54 parasite line and were cultivated at 5% hematocrit in RPMI 1640 medium, 0.5% Albumax II (Invitrogen), 0.25% sodium bicarbonate, and 0.1 mg/ml gentamicin. Parasites were incubated at 37°C in an atmosphere of 5% oxygen, 5% carbon dioxide and 90% nitrogen. Parasite cultures were synchronized using percoll/sorbitol gradient centrifugation as previously described (16,17). Briefly, infected RBCs were layered on a step gradient of 40%/70% percoll containing 6% sorbitol. The gradients were then centrifuged at 12 000g for 20 min at room temperature. Highly synchronized, late stage parasites were recovered from the 40%/70% inter-phase, washed twice with complete culture media and placed back in culture. The level of parasitemia of the resistance experiment (Supplementary Figure S3) was calculated by counting blood smears from two independent parasite cultures stained with Giemsa under a light microscope. The level of parasitemia of the KDEL experiment (Figure 3) was calculated by Flow Cytometry. For flow cytometry, aliquots of 50μl parasite cultures were washed in PBS and incubated 30 min with 1:10 000 SYBR Green I DNA stain (Life Technologies). The fluorescence profiles of infected erythrocytes were measured on CytobFLEX (Beckman Coulter) and analyzed by the CytExpert software.

### Plasmid Construction

The plasmids pVBh (18), pVBbIDh (19) and V2BbIDh (13) were previously described. V stands for a typical var promoter (var7b cloned previously from a Dd2 parasite line (20)) and V2 stands for var2csa promoter. The plasmid pV2Bh was obtained in two steps: First we inserted the AsISI restriction site downstream of the KpnI restriction site by PCR amplification of var promoter region from the plasmid pVBh using 5′-GGGGTACC GGATCCGATTCTGCG TAGAGAAGCCAGAGC-3′ and 5′-GGCATGTTAAC ATAGTCTACG-3′. This fragment was sub-cloned back into pVBh (18) by KpnI/HpaI restriction sites. As a second step, the 5′ UTR of var2csa was amplified from NF54 parasite line using 5′-AAACCGATCGCGGATCC CATTTACAAA-3′ and 5′-ATGTTAACTTTGTCACCA CATTCTAAA-3′ and cloned into the plasmid above by AsISI/HpaI restriction sites. The plasmids pVB-myc and pV2B-myc were obtained by cloning sequence no. 1 (Supplementary Table S1) into the plasmids pVBbIDh (19) and V2BbIDh (13) respectively using HpaI/HindIII. The plasmid pVB+uORF was obtained by cloning sequence no. 2 (Supplementary Table S1) into the plasmid pVBbIDh using EcoRI/HpaI. The plasmid pV2BΔuORF was obtained by cutting out the uORF from the plasmid V2BduORF (14) using NcoI/ClaiI, blunt ends were then generated using Klenow fragment (NEB M0210s) and the construct was finalized by self-ligation. The plasmids pV-GB and pV2-GB were obtained by cloning sequence no. 3 (Supplementary Table S1) into the plasmids pV2BbIDh (19) and V2BbIDh (13) respectively using HpaI/HindIII. The plasmids pV-GB(KDEL) and pV2-GB(KDEL) were obtained by cloning sequence no. 4 (Supplementary Table S1) into the plasmids pVBbIDh (19) and V2BbIDh (13) respectively using HpaI/PmlI. The plasmid pV2mycGB was obtained by cloning sequence no. 5 (Supplementary Table S1) into the plasmid pV-GB using HpaI/PmiI. The plasmid pDuORF-GFP was obtained by cloning sequence no. 6 (Table S1) into the plasmid pV2-GB using KpnI/HpaI. The plasmid pDuORF-GFP was obtained by cloning sequence no. 7 (Supplementary Table S1) into the plasmid pOrf-GFP using ClaI/PmiI.

### Parasite transfection and selection

Parasites were transfected as described (21). Briefly, 0.2 cm electroporation cuvettes were loaded with 0.175 ml of plasmid DNA in incomplete bacterio- cytosme and electroporated using BIORAD GenePulser electroporator. Stable transfectants carrying plasmids with a hDHFR-selectable marker were initially selected on 4 nM WR99210. Stable transfectants carrying plasmids with BSD-selectable marker only (pVBh and pV2Bh) were initially selected on 2 μg/ml blasticidin-S (Invitrogen). In order to obtain parasites carrying large plasmid copy numbers, these cultures were then subjected to elevated concentrations of 4–20 μg/ml blasticidin-S, depending on experimental design.

### Solute permeability measurements

Permeability assays of iRBCs were performed to exclude possible spontaneous resistance to blasticidin S. These assays were done as previously described (22). Briefly, the VB-myc and V2B-myc transfectants were cultivated to the trophozoite stage in standard media with 10μg/mL blasticidin, harvested and enriched to >95% parasitemia using
the percoll/sorbitol method, and resuspended in buffered saline (145 mM NaCl, 20 mM HEPES, 0.1 mg/ml BSA, pH 7.4 with NaOH). Solute uptake was initiated by addition of 20 volumes of 280 mM sorbitol or 145 mM phenyltrimethylammonium chloride (PhTMA-Cl) in buffer (20 mM Na-HEPES, 0.1 mg BSA, pH 7.4) at 37°C with transport inhibitors as indicated (Supplementary Figure S3). Osmotic lysis, which results from net sorbitol or PhTMA-Cl uptake, was continuously recorded by measuring the transmittance of 700 nm light through the cell suspension (hematocrit, 0.125%, DU spectrophotometer with Peltier temperature control). The osmotic lysis halftime, the time required for half-maximal cell lysis, is inversely proportional to solute permeability and was determined by linear interpolation.

**Genomic DNA extraction, RNA extraction and cDNA synthesis**

Genomic DNA was extracted as described (18). Briefly, iRBCs from a 20 ml culture were lysed with saponin and the parasite pellet was washed with PBS and taken up in 200 μl TSE buffer (100 mM NaCl, 50 mM EDTA, 20 mM Tris, pH 8), 40 μl of 10% SDS and 20 μl 6M NaClO4. This suspension was rocked for twelve hours and genomic DNA was extracted with phenol/chloroform. The resulting DNA was taken up in 100 μl D2O. RNA was extracted from synchronized parasite cultures at 20–24 h after percoll/sorbitol gradient centrifugation (16,17). RNA was extracted with the TRIZOL LS Reagent® as described (23) and purified on PureLink column (Invitrogen) according to manufacturer’s protocol. Isolated RNA was then treated with DNase I and centrifuged at 4°C to discard any membrane residuals. The resulting DNA was quantified using RT-qPCR of gDNA and comparing the ΔCt of the single copy housekeeping gene arginine-tRNA ligase (PF3D7_1218600).

**mRNA stability assay**

Parasite cultures were synchronized to late ring stages (20 h post percoll/sorbitol gradient centrifugation). The same amount of parasites were grown in the presence of 10 μg/ml blasticidin in 12-well plates, 1.5 ml culture in each well. 20 μg/ml actinomycin D (ActD, TOKU-E) was added to cultures to inhibit transcription and samples were collected immediately and at 5, 15, 30, 60 and 120 min post addition of ActD. Cultures were collected by washing in PBS. Cell pellets were immediately dissolved in 750 μl TRIZOL LS Reagent® and followed by RNA extraction and cDNA synthesis as described above.

**Western blot**

To collect parasite proteins, infected RBCs were lysed with saponin, then parasites were washed twice with PBS and collected in Laemmli sample buffer (Sigma). The same protein amounts of denatured samples were subjected to SDS-PAGE (gradient 4–20%, Bio-Rad) and electroblotted to a nitrocellulose membrane. Immunodetection was carried out using rabbit anti-BSD (EMCbiosciences, 1:1000), mouse anti-c-Myc antibody (Santa-Cruz, 1:300), mouse anti-GFP (Life Technologies, 1:1000), rabbit anti-aldolase (Roche, 1:1000) or rabbit polyclonal anti-aldolase (1:2000) (26). The secondary antibodies used were antibodies conjugated to Horseradish Peroxidase (HRP), goat anti-rabbit and goat anti-mouse (Jackson ImmunoResearch Laboratories, 1:10 000). The immunoblots were developed in EZ/ECL solution (Israel Biological Industries).

**Cell fractionation of infected erythrocytes**

Parasite cultures were lysed on ice by saponin and centrifuged at 5000 rpm for 5 min. The parasite pellet was washed three times with ice-cold PBS containing protease inhibitors, resuspended in 500 μl H2O and lysed by three ‘thaw and freeze’ cycles at −80°C. Samples were centrifuged at 14 000 rpm for 15 min at 4°C and then separated into soluble and pellet fractions. The soluble fraction was centrifuged again at 4°C to discard any membrane residuals and concentrated using Amicon Centrifugal filters (Amicon Ultra 0.5 ml centrifugal filters, 3K, Millipore) at 14 000 rpm for 30 min at 4°C. The pellet fraction was washed three times with ice-cold PBS containing protease inhibitors (Roche). Both fractions were mixed to the same volume with sample buffer and subjected to western blot analysis.

**Immunofluorescence assay and live cell imaging**

IFA was performed as described before (26) with few modifications. Briefly, 1 ml of parasite culture at 5% parasitemia was washed with PBS and re-suspended in a fresh fixative solution (4% Paraformaldehyde (EMS) and 0.0075% glutaraldehyde (EMS) in PBS). Fixed parasites were treated with 0.1% Triton-X 100 (Sigma) in PBS, then blocked with 3% BSA (Sigma) in PBS. Cells were then incubated with primary antibodies used at the following dilutions: rabbit anti-BSD (EMCbiosciences, 1:1000), mouse anti-Actin (MR4, 1:1000), rabbit anti-GFP (Life Technologies, 1:300). Samples were washed and incubated with Alexa Fluor 555 goat anti-rabbit (Life Technologies, 1:500), Alexa Fluor 568 goat anti-Mouse (1:500, Life Technologies) or Alexa Fluor 488 goat anti-rabbit (Life Technologies, 1:500) secondary antibodies. Samples were washed and laid on ‘PTFE’ printed slides (EMS) and mounted in ProLong Gold antifade.
Stochastic Optical Reconstruction Microscopy (STORM) imaging and analysis

Parasite cultures were saponin lysed and washed three times with fresh PBS. Parasites were then re-suspended on ice with fresh fixative solution (4% paraformaldehyde (EMS) in PBS) for 1 h. Fixed parasites were allowed to air dry in an eight well-chambered cover glass system 1.5H (In Vitro Scientific) for 2–3 h at room temperature and covered with glucose (30 mM in PBS) to quench unreacted aldehyde groups. Samples were washed three times with PBS and permeabilized with 0.1% Triton-x 100 (PBS) for 10 min at room temperature, washed with PBS and blocked with 10% horse serum (Biological Industries) containing 3% BSA (IgG free, Sigma) and 0.025% Tx-100 in PBS for 1 h at room temperature. Primary antibodies, rabbit anti-GFP (1:300, life technologies) and mouse anti-β-actin (MR4, 1:750) were diluted in 5% horse serum containing 0.025% Tx-100 and applied for 1 h at room temperature, followed by five washes in PBS containing 0.025% Tx-100. Secondary antibodies, Donkey anti-Mouse Alexa647 (1:300, Jackson ImmunoResearch) and Goat anti-Rabbit Alexa568 (1:300, life technologies) were applied for 1 h in 5% horse serum containing 0.025% Tx-100 in PBS at room temperature, followed by five washes with PBS containing 0.025% Tx-100. Parasite nuclei were labelled with YOYO-1 (1:300, Life Technologies) for 20 min at room temperature, followed by 3 washes with PBS. STORM was performed by a Nikon Eclipse Ti-E microscope with a CFI Apo TIRF × 100 DIC N2 oil objective (NA 1.49, WD 0.12 mm). Multi-channel calibration was performed prior to data acquisition using a PlanApo 100× oil NA = 1.5, WD = 130 μm lens on a Teflon coated slide (EMS), covered with cover slips and visualized immediately. Fluorescent images were obtained using a Nikon Eclipse Ti-E microscope equipped with a CoolSNAP Myo CCD camera. Images were processed using the NIS-Elements AR (4.40 version) software.

Immuno-electron microscopy

Sample preparation for Immuno-electron microscopy was done as described (29) with few modifications. Samples were immersed in 1-Hexadecane and then high-pressure frozen in a high pressure freezing machine (HPM010, BAL-TEC). Samples were then freeze-substituted in dry acetone containing 0.1% uranyl acetate and 0.1% glutaraldehyde for 30 h at –90°C in a freeze substitution machine (Leica EM AFS). Samples were slowly warmed to ~30°C, washed in dry ethanol, and infiltrated with increasing concentrations of Lowicryl HM-20 resin. Samples were UV-polymerized at ~30°C, sectioned and deposited on formvar-coated nickel 200 mesh grids. Grids were treated with 0.5% blocking solution (0.5% gelatin, 0.5% BSA) for 20 min and then incubated with anti-GFP antibody in blocking solution (ab6556, Abcam) for 2 h at RT. Grids were rinsed with 0.1% glycine in PBS and incubated with 10 nm gold conjugated goat anti-mouse (EMS) for 30 min. Grids were washed with PBSX1 and DDW. Samples were visualized using an FEI Tecnai T-12 (FEI Company, Eindhoven, the Netherlands).

RESULTS

We have previously shown that it is possible to silence the entire var gene family and erase its epigenetic memory using promoter titration (18,30). In this technique, one can select for increasing numbers of active var promoters on episomes that outcompete activation factors needed for transcription of the endogenous var genes and effectively silence the entire gene family. Surprisingly, we were unable to completely silence and erase the epigenetic memory of a clonal population expressing var2csa using promoter titration with a typical var promoter on the competing pV Bh episomes (Supplementary Figure S1). We therefore decided to perform promoter titration with plasmids that express the drug selectable marker bsd (which encodes the deaminase that confers resistance to blasticidin-S) by a var2csa promoter. We used two approaches to perform promoter titration: first, we used the plasmid pV2 Bh to directly select for active episomal var2csa promoters using blasticidin; and second, we used the plasmid pV2 Bh D, in which the silencing effect of var intron was disabled (31), and therefore it is possible to select first for stable episomal transfection.
using hDHFR as selectable marker and activate the var2csa promoter only after the establishment of parasite populations that stably carry the episomes (see plasmid maps in Supplementary Figure S2). For comparison, we used parasites which were transfected with similar plasmids except that a typical Up/C var promoter was used instead of the var2csa promoter (in pVBh and pVBbIDh respectively). As a first step, we wanted to confirm that increasing drug concentrations would increase bsd expression as previously observed (18,32–35). As expected, in the parasites that expressed the typical var promoter, increasing drug concentration was associated with an increase in the mRNA levels from the activated episomal var promoter (Supplementary Figure S2A and B). However, the parasites expressing bsd from the var2csa promoter did not elevate mRNA levels in response to increase selection pressure (Supplementary Figure S2C and D). To exclude that the different response in mRNA levels between the two promoter types was due to differences in RNA stability we incubated both parasite lines, expressing the two different promoter types, with the transcription inhibitor Actinomycin D (ActD) and compared the rate of RNA decay. There was no difference in the rate of RNA decay indicating that RNA transcribed by the two promoter types is equally stable (Supplementary Figure S2E and F).

We then tested whether the increase in drug pressure had forced the parasites to express more protein, assuming that the similar growth rate of both lines (Figure 1A) implies that they make more of blasticidin-S-deaminase (BSD) to de-toxify blasticidin S and overcome the increase in its concentration. To our surprise, while the increase in selection pressure for activation of typical var promoters resulted in both elevated levels of mRNA and a gradual increase in protein expression, no protein was detected by western blot in parasites expressing the typical var promoter. The protein was easily observed in the parasite cytoplasm via plasmoidal surface anion channel (PSAC). To reduce uptake, the parasite may either silence or switch expression of the two clag3 genes that determine PSAC activity (36–39). Low concentrations of blasticidin (<0.4 μg/ml) selects for clag3.1 expression whereas resistance to higher concentrations of blasticidin (up to 2 μg/ml) results from silencing of both clag3 genes (38). However, here both the pre-transfected lines as well as the parasites carrying the active var2csa promoters continued to express clag3.1 gene even when selected with very high concentrations of 10 μg/ml blasticidin (Supplementary Figure S3B, D, F, H). We also performed PSAC permeability assays with two organic solutes, sorbitol and phenyltrimethy lammonium (PhTMA) that use distinct routes through this channel (40). These measurements revealed preserved channel-mediated permeability in both the VB-myc and V2B-myc lines (Supplementary Figure S3 I-L); halftimes of 12.5 ± 0.7 in sorbitol (n = 6 trials for each line) and 5.4 ± 0.5 min in PhTMA (n = 5 trials each); P = 0.7 for comparisons between the two lines in each solute); these values contrast with markedly greater halftimes indicative of reduced permeability in parasites resistant to blasticidin without BSD expression (106 and 22 min for sorbitol and PhTMA, respectively (36)). Block by the potent PSAC inhibitor ISG-21 was also unchanged and indistinguishable in the two transfectedants (Supplementary Figure S3M–N); ISG-21 was less effective against PhTMA uptake than sorbitol in both lines, as previously reported (40). These findings clearly demonstrate that both parasite populations that express bsd, under either the typical var promoter or the var2csa promoter, are equally resistant to osmotic lysis (Supplementary Figure S3I–N), and excludes altered host cell permeability as the mechanism of blasticidin resistance in parasites expressing bsd by the var2csa promoter.

We hypothesized that the uORF found within the 5′ UTR of var2csa could account for the different RNA and protein phenotypes we observed, since this is the main regulatory element that is different between the upstream region of typical var genes and var2csa. To test this hypothesis, we created two additional constructs. In the first we inserted the uORF into the 5′ UTR of a regular var promoter (Figure 2A, left), and in the second we have deleted the uORF from the 5′ UTR of var2csa (Figure 2A, right). We then subjected the transfected parasites to increasing concentrations of blasticidin, and measured steady-state RNA as well as BSD protein levels. In parasites carrying the episomes in which the uORF was removed from var2csa 5′ UTR, we observed an increase in steady state RNA levels that was associated with elevated protein levels detected by Western blot as we increased drug concentration (Figure 2C, E). In addition, the protein was easily observed in the parasite cytoplasm (Figure 2F, upper panel). On the other hand, in parasites that carried the episomes in which the uORF was inserted into a typical var 5′ UTR, at a similar location where it resides in the original var2csa promoter, mRNA levels did not elevate in response to the increase in drug pressure and more strikingly, the protein was undetectable by Western

\[ \text{var2csa} \]
Figure 1. The protein expressed by an activated var2csa promoter is not detected. (A) Growth curves of parasites expressing bsd by typical var and var2csa promoters show that they grow at similar rates on increasing blasticidin pressure. (B) Western blot analyses of BSD expressed by either var or var2csa promoters and selected using increasing levels of blasticidin. (C) Western blot analyses of BSD expression by a var2csa promoter or a var promoter (D) at different time points in the parasite’s life cycle. (E) Quantification of BSD expression by a var2csa promoter or a var promoter at different time points in the parasite’s life cycle using densitometry analysis of three independent experiments. **P < 0.01. (F) Western blot analyses of BSD-myc fusion expressed by either var or var2csa promoters and selected using increasing levels of blasticidin. hpi – hours post invasion; NC: Negative control (untransfected); PC: positive control. Anti-Aldolase antibody was used as a loading control. (G) Immunofluorescence microscopy of BSD expressing parasites under the control of a var (upper panel) or var2csa promoter (lower panel). BSD was labeled with an anti-BSD antibody (Red). Nuclei were stained with DAPI (blue). Scale bars: 1 μm. (Total of 120 pVBbIDh expressing parasites all showed diffused cytoplasmic localization; Total of 45 pV2BbIDh expressing parasites all showed either no signal or faint foci around the nucleus).
Figure 2. mRNA and protein levels expressed by the var2csa promoter are determined by the uORF. (A) Schematic of plasmids. Left: pVB+uORF, bsd is expressed with a typical var promoter in which the var2csa uORF was inserted into the 5' UTR. Right: pV2BΔuORF, bsd is expressed with var2csa promoter in which the uORF was deleted. (B and D) Steady-state transcript levels of bsd and western blot analyses of BSD expression from a typical var promoter which contains the uORF of the var2csa promoter (pVB+uORF) selected with increasing blasticidin concentrations. ns: not significant (P > 0.5: two-tailed, unpaired, Student's t-test). (C and E) Steady-state transcript levels of bsd and western blot analyses of BSD expression, from an uORF deleted var2csa promoter (pV2BΔuORF) grown in the presence of increasing blasticidin concentrations. Significance is marked with ** (P < 0.01: two-tailed, unpaired, Student's t-test). mRNA levels are presented as relative copy number to the housekeeping gene arginine-tRNA ligase (PF3D7_1218600). Aldolase was used as a loading control. (F) Immunofluorescence microscopy of BSD expressing parasites. Upper panel: expression of BSD by an uORF-deleted var2csa promoter (pV2BΔuORF). Lower panel: expression of BSD from a var promoter which contains the uORF of the var2csa promoter (pVB+uORF). BSD was labeled with an anti-BSD antibody (red). Nuclei were stained with DAPI (blue). Scale bars: 2 μm (39/47 pV2BΔuORF expressing parasites showed diffused cytoplasmic localization; total of 39 pVB+ORF expressing parasites were counted and showed no signal).
blot and IFA (Figure 2B, D and F, lower panel). These results are reciprocal to the original transfections with intact var and var2csa promoters, indicating that the uORF found in var2csa 5’ UTR sequence is responsible for the differences in both RNA and protein phenotypes.

Struck by the observation that the parasites expressing BSD by the intact var2csa promoters (in pV2Bh, pV2B-myc and pV2BbIDh) displayed resistance to blasticidin despite virtually undetectable levels of BSD (Figure 1), we reasoned that it is likely that in these circumstances, the parasite expresses very small, undetectable amounts of the BSD that are sufficient to confer resistance. One way to achieve resistance in spite of very low levels of deaminase is if the deaminase is expressed at the location where blasticidin S blocks translation. This might create a subcellular niche that enables enough protein translation for the parasite to survive. We applied Stochastic Optical Reconstruction Microscopy (STORM), which is a technique sensitive enough to detect photons from single molecules, and were able to detect the deaminase expressed by var2csa in small complexes localized mainly to the ER (Figure 3A–C), while the protein expressed by a typical var promoter was distributed throughout the parasite cytoplasm (Figure 3D). Further, we decided to validate that the protein expressed by the var2csa promoter enters the ER-Golgi system by fusion of an ER retention signal to the expressed protein. We hypothesized that such a signal would retain more of the deaminase expressed by var2csa in the ER and would help the parasite to resist increasing blasticidin concentrations. We further predicted that addition of this signal would have little effect on cytoplasmically located BSD expressed from a typical var promoter. The four amino acid motif KDEL, which has been previously shown to function as an ER retention signal that retrieves proteins from the Golgi back to the ER by COPI vesicles (41) was fused at the C’ terminus of the gfp-bsd fusion protein expressed either by var2csa or var promoters. We then transfected parasites with these expression vectors as well as with versions of the plasmids that did not contain the KDEL signal for each of the promoter types (Figure 3E). Parasites were first selected using WR99210 for stable episomal transfection and then with 2 μg/ml blasticidin for activation of either var2csa or var promoters. After the different parasite populations carrying each of the plasmids were established, we increased blasticidin concentration to 6 μg/ml and followed their growth over time. We found that cultures of parasites expressing bsd by a typical var promoter grew at a normal growth rate regardless whether the bsd was fused with the KDEL signal or not, as expected for a cytosolic protein. However, in those parasites expressing the var2csa promoter there was a significant improvement in growth rate of parasites in which the bsd was fused to the KDEL motif compared to those that did not contain it (Figure 3F), even though the protein is still not detected by Western blot (Supplementary Figure S4). Very similar results were obtained when we increased blasticidin-S concentration further from 6 to 10 μg/ml, indicating that only for the var2csa promoters, the fusion of the ER retention signal fosters the parasite’s ability to resist increasing drug concentrations (Figure 3G). Altogether, these data suggest that protein expressed by var2csa promoters find their way to the ER-Golgi trafficking machinery.

These data, together with the results of the reciprocal placement of the var2csa-uORF at a var 5’ UTR, suggest that the uORF could be responsible for diverting the protein expressed by the var2csa promoter to be associated with the ER. To test this assumption, and surpass the translational repressive effect of the uORF, we created a plasmid in which the uORF sequence is fused with a bsd-gfp reporter, which is expressed by a typical var promoter. We compared the cellular localization of the fused reporter with the localization of the non-fused reporter expressed by the same var promoter. Strikingly, while the protein which was not fused to the uORF is cytosolic (Figure 4A), the fusion of the uORF sequence significantly changes its subcellular location. The protein can be visualized around the nucleus as distinct foci associated with the ER (Figure 4B). A closer look on the cellular localization of the fused protein at the ultrastructural level by immuno-Electron Microscopy indicated that the fused protein is primarily associated with membranes surrounding the nuclear envelope (Supplementary Figure S5A–C) corresponding with ER location. To further validate these findings, we performed cellular fractionation followed by western blot analysis and found that the protein fused to the uORF sequence is targeted to the membrane fraction while the one expressed without the uORF-fusion is cytosolic (Figure 4D). This set of experiments shows that the var2csa-uORF contains a signal for expression at a specific cellular localization associated with the ER.

The signal for specific cellular localization could be at the RNA transcript of the uORF or the peptide itself. To differentiate between these two mechanisms, we made a similar construct to the one described above, in which the uORF sequence is fused with a bsd-gfp reporter expressed by a typical var promoter, only in the current plasmid the uORF-coding sequence was replaced with a different DNA sequence, encoding for the same polypeptide (14). We found that bsd-gfp reporter fused to the uORF polypeptide encoded by the alternative sequence localized to a distinct spot similar to the fusion with the endogenous uORF sequence (Figure 4C), indicating that the cellular localization signal is determined by the polypeptide rather than the RNA transcript. In order to demonstrate that the uORF itself is translated into a polypeptide, we constructed a plasmid in which the gfp-bsd fusion protein is expressed by var2csa promoter where the uORF is fused with a Myc epitope tag that allows detection of the polypeptide (Figure 5A). We first confirmed that the uORF functions as a translational repressor and that GFP isn’t detectable (data not shown). Then we showed that the polypeptide encoded by the var2csa uORF is expressed and concentrates mainly around the nucleus at ring and trophozoite stages but is hardly expressed in schizonts (Figure 5B and C). This data shows that in its natural context in the 5’ of var2csa the uORF is translated into a polypeptide.

DISCUSSION

Upstream open reading frames (uORFs) are important regulators of eukaryotic gene expression. uORFs may impact expression of a downstream protein coding region by triggering nonsense-mediated decay (NMD) or by controlling protein translation (42). Computational analysis of the P.
Figure 3. The protein expressed by the var2csa promoter is associated with the ER. (A) Super-resolution STORM imaging of parasites expressing BSD-GFP fusion by a var2csa promoter (Green; pV2-GB) that were co-stained with the ER marker Pf39 (red). (B) High magnification of the region indicated in
The *P. falciparum* genome had identified a large number of possible uORFs. However, since translation start and stop codons are AT-rich, and the *P. falciparum* genome is considered one of the most AT-rich among eukaryotes, composed of ~80% AT, it is possible that the number of functional uORFs in *P. falciparum* is much lower. Nonetheless, recent ribosome footprint analysis provides experimental support to the prediction that uORFs are quite abundant in *P. falciparum* 5′ UTRs (43). To the best of our knowledge, the only uORF in *Plasmodium*, to which a defined regulatory role had been demonstrated, is the uORF of var2csa that regulates expression of the dORF at the level of translation re-initiation (13,14). Here, we demonstrate that when forcing activation of the var2csa promoter using a drug selectable marker, the parasites express very low levels of protein while transcribing high levels of mRNA. It is possible that under these experimental conditions the strong translational repression effect of the uORF is maintained and that there was no switch that enhanced the efficiency of translation re-initiation of the dORF. Nonetheless, parasites expressed enough protein that enabled them to overcome drug pressure. The sensitivity of super-resolution STORM imaging, which is based on the ability to visualize single fluorescent molecules, allowed us to detect these low levels of the protein expressed by var2csa promoters and to associate it with the ER. Our data clearly indicate that a regulatory element found at the 5′ UTR of var2csa influenced the localization of the protein encoded by the dORF.

Blasticidin S is a fungal toxin that kills most prokaryotic and eukaryotic cells by inhibiting mRNA-directed translation on ribosomes (44). The specific cellular location of the deaminase expressed by the var2csa promoter, at a site associated with the ER (contrary to the cytosolic localization of the deaminase expressed by a var promoter) which is a primary translation site, might explain how the parasites overcome drug pressure with expression of only very small amounts of the deaminase. One possible explanation is that blasticidin enter the parasite through a retrograde transport, similar to shiga, cholera and ricin toxins, which enter the cell by retrograde transport through endosomes to the Golgi and ER (45). If blasticidin gets into the parasite in a similar pathway, localizing the deaminase to the ER, directly where blasticidin must pass, would greatly increase its efficiency in detoxifying the molecule. Nonetheless, the exact mechanism by which the parasite overcomes blasticidin pressure at very low levels of deaminase expression requires further investigation.

Sequence analysis of the 5′ UTRs of the entire var gene family had indicated they could be classified into several subtypes named UpsA, UpsB and UpsC, while the 5′ UTR of var2csa shows very little similarity to other var subtypes and was classified as UpsE (46). Interestingly, the UpsE promoter could not be completely silenced by promoter titration, and was not responsive to selection by increasing the mRNA levels as observed with other var promoters. This suggests that while var2csa is part of the var gene family for mutually exclusive expression and antigenic switches, it has additional unique regulatory mechanisms. The only regulatory element characterized which is unique to the 5′ UTRs of var2csa is the uORF, and therefore we reasoned that it might be responsible for the different phenotypes of the two promoters. Remarkably, we were able to switch between these two phenotypes by removing the uORF from the var2csa promoter and inserting it into a typical var promoter, indicating that the presence of the uORF at the 5′ UTR determines the fate of the protein expressed by these promoters. The fusion of the uORF sequence with reporter genes overcame the translational repressive effect of the uORF and enabled us to determine that it contains a signal that controls cellular localization. This fusion showed that not only does the uORF sequence change the protein location from the cytoplasm to the ER, but it also diverts it from being a soluble protein to a membrane-associated protein. Interestingly, the uORF does not seem to have a defined hydrophobic region. Nevertheless, both reporter genes used in our experiments contained a predicted hydrophobic domain which could contribute to its association with the ER.

Most of the proteins that are trafficked from the parasite to the red blood cell surface contain a signal peptide and a transmembrane domain that directs them to the ER. In addition, they have a PEXEL/HT motif positioned ~20 amino acids downstream of the signal peptide (47,48), which is cleaved by plasmepsin V (49,50) and allows them to translocate through a translocon called PTEX, which enables them to enter the parasitophorous vacuole membrane (PVM) (51). However, PfEMP1 does not have a defined signal peptide, its transmembrane domain is located internally, near the C-terminus of the protein (at the 3′ end of exon 1), and its PEXEL-like motif isn’t cleaved (52). Therefore, PfEMP1 is classified as PNEP (PEXEL-negative exported protein). Using a PfEMP1 mini-gene system, Melcher et al.,
Figure 4. The var2csa-uORF directs protein expression to a specific cellular localization associated with the ER. (A and B) Immunofluorescence microscopy of parasites expressing the constructs pV-GB and puORF-GFP (schematically shown above each panel). BSD was labeled with an anti-GFP antibody (green) and the ER with a specific antibody against Pf39 (red). Nuclei were stained with DAPI (blue). Scale bars: 2 μm (total of 158 GFP positive, pV-GB expressing parasites all showed diffused cytoplasmic localization; 37/41 GFP positive, puORF-GFP expressing parasites showed distinct focus associated with the ER). (C) In vivo imaging of parasites expressing the construct pDuORF-GFP, which is similar to the puORF-GFP plasmid only that the uORF sequence was replaced with an alternative sequence that encodes for the same peptide. Nuclei were stained with Hoechst (blue). Scale bars: 2 μm (total of 56 GFP positive, pDuORF-GFP expressing parasites all showed distinct focus at the periphery of the nucleus). (D) Western blot analyses of cellular fractionation of parasites expressing V-GB and uORF-GFP separated into soluble supernatant (S) and pellet (P) fractions. Aldolase was used as a marker of a parasite's cytoplasmic protein.
have demonstrated that the information for PfEMP1 export to the RBC surface is found within its semi-conserved head structure (N terminal segment—NTS) in combination with the predicted transmembrane domain and its cytoplasmic tail (53). This would explain why an unrelated protein that contains no PfEMP1 coding segments expressed from a var promoter will be localized to the parasite cytosol. Our data indicate that in addition to the export information found in PfEMP1 coding regions, the uORF of var2csa contains a unique signal that associates the expressed downstream protein with the ER. Moreover, sequence analysis of ~400 var genes from different parasite isolates showed that PfEMP1 NTS is conserved among all var genes and could be classified into two subtypes (54). However, var2csa is exempt from this sequence conservation and its NTS shows no sequence similarity to the NTS of other var genes, which could be explained by the presence of a cellular localization signal at the uORF of var2csa.

The mechanisms by which the uORF of var2csa influences the cellular localization of the dORF encoded protein are yet to be discovered. One could hypothesize that a specific mRNA splicing event could fuse the uORF to the dORF and create one ORF similar to the one we have created with the reporter gene that associates it with the ER. However, we and others (13) were unable to verify that such a splicing event of the pre-mRNA exists in the native uORF context in the 5′ UTR. Another reasonable hypothesis is that the peptide encoded by the uORF could play a role in targeting the protein encoded by the dORF to its destination. Similar phenomena had been observed in a polyprotein system which were encoded by the same mRNA but separated by the 2A skip peptide (55). When the upstream protein was fused with an N terminal signal sequence the downstream protein, lacking any signal sequence was also translocated into the ER despite the discontinuity from the peptide backbone of the upstream protein. Possible inter-
pretation of these results is that the second protein, lacking any signal, ‘slipstreamed’ through the ribosome–translocon complex formed by the first protein containing the localization signal. uORFs are probably the best studied subclass of regulatory small open reading frames (sORFs) found in eukaryotic genomes. Over the past few years it has become evident that many of the sORFs could encode small proteins involved in regulation of protein–protein interactions, enzymatic activity, signaling and cell–cell communication (reviewed in [56]). It was also demonstrated that the *Drosophila* polar granule component (*pgc*), which is a short protein of 71 amino acids, could regulate recruitment and localization of the kinase P-TEFb to transcription sites (57,58). While the main regulatory mechanism by which uORFs function as repressive genetic elements is by the act of translation per se, one cannot preclude that the short proteins they encode also have a regulatory function. Very little is known about the expression of the putative short protein encoded by the var2csa uORF. Nonetheless, our data strongly supports the presence of a cellular localization signal that determines the fate of the dORF, however, how it functions and regulates these processes requires further investigation.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR online.

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