Single-stranded DNA binding protein from human malarial parasite *Plasmodium falciparum* is encoded in the nucleus and targeted to the apicoplast

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**ABSTRACT**

Apicoplast, an essential organelle of human malaria parasite *Plasmodium falciparum* contains a ~35kb circular genome and is a possible target for therapy. Proteins required for the replication and maintenance of the apicoplast DNA are not clearly known. Here we report the presence of single–stranded DNA binding protein (SSB) in *P falciparum*. PfSSB is targeted to the apicoplast and it binds to apicoplast DNA. A strong ssDNA binding activity specific to SSB was also detected in *P. falciparum* lysate. Both the recombinant and endogenous proteins form tetramers and the homology modelling shows the presence of an oligosaccharide/oligonucleotide-binding fold responsible for ssDNA binding. Additionally, we used SSB as a tool to track the mechanism of delayed death phenomena shown by apicoplast targeted drugs ciprofloxacin and tetracycline. We find that the transport of PfSSB is severely affected during the second life cycle following drug treatment. Moreover, the translation of PfSSB protein and not the transcription of PfSSB seem to be down-regulated specifically during second life cycle although there is no considerable change in protein expression profile between drug-treated and untreated parasites. These results suggest dual control of translocation and translation of apicoplast targeted proteins behind the delayed death phenomena.

**INTRODUCTION**

*Plasmodium falciparum*, the causative agent of most severe form of human malaria belongs to the phylum apicomplexa (1). The apicomplexan parasites possess a non-photosynthetic plastid like organelle called apicoplast. The apicoplast is the site for several biosynthetic pathways (fatty acid, isoprenoid, heme biosynthesis) that are central to the core cellular functioning of the parasite (2). The biosynthetic pathways located within the apicoplast provide novel sites of drug intervention against the malaria disease (3). Moreover, the housekeeping metabolic processes like DNA replication, transcription and translation operative within the apicoplast are prokaryotic in nature and therefore they are good targets of existing antibacterials or novel drugs (4).

The ~35kb, extremely A+T rich (~86%) apicoplast DNA is circular and replicates via a D-loop/bi-directional theta mode in late trophozoite/early schizont stages of intraerythrocytic life cycle (5). The origins of apicoplast DNA replication have been mapped in the inverted repeat (IR) regions (6). The apicoplast genome houses only 68 open reading frames (ORFs) for rRNAs, tRNAs, ribosomes, RNA polymerase, translational elongation factor (EF-Tu), ClpC chaperone and Fe–S cluster protein (SulB) (1,7). None of the ORFs on the apicoplast DNA codes for replication protein. Most of the proteins involved in the apicoplast metabolic pathways are encoded in the nucleus, synthesized in the cytoplasm and subsequently imported into the apicoplast (2). The nuclear encoded proteins are targeted to apicoplast by the apicoplast localization signal that is cleaved off from the protein once inside the apicoplast (8). Recently, a ~220kDa nuclear encoded multifunctional enzyme PfPRex (*P. falciparum* plastidic...
DNA replication/repair enzyme complex) has been shown to localize in the apicoplast. This enzyme has three important activities (helicase, primase and polymerase) associated with DNA replication (9). A histone like protein (HU) exhibiting the DNA condensation property is imported into apicoplast suggesting its role in organization of apicoplast genome (10). We and others have shown that the gyrase subunits present in the parasite are targeted to apicoplast where they might be involved in negative supercoiling of the DNA circle, an essential step for the replication process (11,12). Consistent with the above findings, quinolone (ciprofloxacin) or coumarin (coumermycin, novobiocin) antibiotics target the parasitic gyrase and inhibit the apicoplast DNA replication leading to the parasite death (12,13).

Since apicoplast is of prokaryotic origin, several antibiotics against bacterial replication, transcription and translation processes have been used successfully to block parasitic growth. However, majority of these antibiotics show typical delayed growth phenotype, characterized by defect in parasite growth and decrease in parasitemia only during second life cycle following the addition of these drugs (4). It has been suggested that these drugs affect apicoplast morphology, segregation and most importantly the transport of essential proteins in the apicoplast. Using fusion protein containing apicoplast signal sequence of acyl carrier protein (ACP) and GFP, the effect of these drugs on protein translocation have been studied (14,15). However, no endogenous apicoplast targeted protein involved in housekeeping function has been followed after drug treatment in the above studies. Neither the transcription and translation status of apicoplast targeted proteins was investigated simultaneously in the presence of these drugs.

To gain further insight into the enzymology of \( P. falciparum \) apicoplast DNA replication, we wanted to study another essential key protein, the homologue of bacterial single-stranded DNA binding protein (SSB) (16). The prokaryotic type circular \( P. falciparum \) apicoplast DNA suggests a possible requirement of bacterial type SSB for various DNA metabolic processes in that organelle. The analysis of the parasite genome indeed reveals the presence of a bacterial type \( ssb \) on the chromosome \( V \) of the nuclear DNA (PFE0435c). The N-terminal extension in the primary sequence of the protein is predicted to be a potential apicoplast targeting sequence.

SSBs are known to play essential roles in many aspects of nucleic acid metabolism including DNA replication, recombination and repair. SSBs protect and stabilize the single-stranded DNA (ssDNA) intermediates as well as remove secondary structures in the DNA. Any misstep in DNA replication or a failure to properly recombine or repair DNA can lead to gross aberrations in the genome, signifying the role of SSBs in these processes (17).

In eukaryotes, a heterotrimeric complex called replication protein A (RPA) (18) carries out the ssDNA binding activity in the nucleus. Mitochondrial DNA replication involves the ssDNA binding protein (mtSSB) that is quite distinct from nuclear RPA. mtSSB shows a great degree of sequence homology with bacterial SSB (19). In \( P. falciparum \), however, ssDNA binding activity in the nucleus has been reported earlier corresponding to the large subunit (~55 kDa) of RPA (20).

SSB proteins from different organisms share sequence homology as well as distinct biochemical and structural characteristics. The SSBs have a common oligosaccharide/oligonucleotide-binding fold (OB fold) which binds to ssDNA. The SSBs from all prokaryotic organisms have an acidic C-terminal tail that is essential for DNA replication by mediating protein–protein interactions at the replication fork (17). In solution, SSBs are found in different oligomeric states. They are found as homodimers (Bacteriophages, \( T. thermophilus \), \( T. aquaticus \) and \( Deinococcus radiodurans \)), heterotrimer (eukaryotic RPAs) and homotetramers (mitochondrial and most prokaryotic SSBs) (21).

We report here the presence of a prokaryotic SSB homologue in the protozoan parasite \( P. falciparum \) (PiSSB) that is encoded in the nucleus and targeted to the apicoplast. PiSSB is expressed in all the three stages of intraerythrocytic life cycle of parasite and binds specifically to apicoplast DNA. Both recombinant as well as endogenous PiSSB proteins form tetramers in solution like \( E. coli \) SSB protein although it fails to complement \( E. coli \) ssb mutant cells suggesting subtle differences between these proteins possibly due to the differences in the C-terminal region. Consistent with the expression of PiSSB in the parasites, a strong ssDNA binding activity specific to PiSSB was found in the parasite lysate. Further, we show that inhibitors against apicoplast replication and translation machinery modulate PiSSB expression and function significantly, possibly by affecting the translocation of the protein to the apicoplast as well as intervening with the translation of the protein during the second life cycle following drug treatment. These results suggest a mechanism for delayed death phenomena shown by parasites treated with drugs that affect apicoplast housekeeping functions.

**MATERIALS AND METHODS**

**Construction of recombinant expression vector**

The expression vector pET28a (Novagen) with N-terminal His\(_6\)-tag was used to clone \( ssb \). Pfssb was amplified (omitting apicoplast targeting signal) from the genomic DNA by means of polymerase chain reaction using triple master mix PCR enzyme (Eppendorf) and specific primers (P1–P2). Please see Table 1 for all primer sequences. The amplified DNA product was double digested with \( \text{BamHI} \) and \( \text{XhoI} \) restriction enzymes (New England Biolabs), ligated into pET28a expression vector and used to transform DH10b bacterial cells. The recombinant clone was subsequently sequenced and the sequence perfectly matched with the sequence reported in PLASMODB.ORG (PFE0435c).

**Protein purification**

His\(_6\)-PiSSB was purified using Ni–NTA resin (QIAGEN) as per the protocol suggested by the vendor. The details of the protocol are described in the supplementary section (‘Materials and Methods’ section). GST-PiSSB and GST
proteins were purified using glutathione sepharose 4B beads as described in the Supplementary Data ('Materials and Methods' section).

**Immunodepletion assay**

GST-PfSSB (50–60 µg) or GST bound glutathione sepharose 4B beads were incubated overnight with 30 µl crude anti-SSB sera at 4°C. These beads were separated by low-speed centrifugation (3000 r.p.m.) from immunodepleted sera. Each immunodepleted sera (3 µl) were taken out and the rest were further incubated with respective fresh protein bound beads for another 8 h at 4°C. After the second round of immunodepletion process, the sera were separated from the beads as described earlier. Western blot experiments were performed using four (1–4) different PVDF strips having equal amount of parasite lysate. The first strip was treated with untreated anti-SSB sera (1:5000 dilution) as control, second strip was treated with second round GST-depleted anti-sera whereas third and fourth strips were treated with first and second round GST-PfSSB depleted anti-sera with same dilution as control. All the blots were retreated with anti-Pf Actin sera to confirm equal loading.

**Electrophoretic mobility shift assay**

To investigate the ssDNA binding activity of PfSSB, EMSA was performed either using a mixture of M13mp18 circular ssDNA and pUC18 circular dsDNA or radiolabelled 70 base synthetic oligonucleotide as described elsewhere (22,26). The details of the EMSA protocol are described in the Supplementary Data ('Materials and Methods' section).

**Generation of antibodies**

Polyclonal antibodies against purified His<sub>6</sub>-PfSSB were raised in mice using essentially the protocol described previously by Harlow and Lane (23).

**Western blotting and immunoprecipitation**

Either mixed stage or synchronized parasites were released from the host RBCs by saponin lysis and the parasites were resuspended in 50 mM Tris–Cl (pH 7.5) containing protease inhibitor cocktail (Sigma) followed by addition of SDS–PAGE loading buffer and boiling for 5 min at 95°C. The cell lysates were resolved in 12% SDS–PAGE and subsequently used for western blot analysis. The immunoprecipitation was carried out using the protocol described elsewhere (10).

**Parasite culture and drug treatment**

Plasmodium falciparum (3D7) strain was cultured as described earlier (11). Synchronization of parasite culture was obtained by treating the RBC-infected parasites in 5% Sorbitol (W/V) solution in every 48 h at ring stage. Study of effect of different drugs on parasites and the modulation of PfSSB and other proteins was conducted by treatment with ciprofloxacin (5, 10 and 15 µM) or tetracyclin (10 µM) or artemisinin (10, 20 and 30 nM) in parasite culture containing 0.5% parasitemia in 6-well culture plate. Fresh medium with different drugs were replaced in every 24 h. Finally, parasites were harvested in different stages (first or second life cycle). MG132 (100 nM) was added in the ciprofloxacin treated culture during the second life cycle.

**Chromatin immunoprecipitation assay**

Chromatin immunoprecipitation (ChIP) assay was performed essentially following the protocol described by Ram et al. (10) with appropriate modifications. Please see the Supplementary Data ('Materials and Methods' section) for details.

**RT-PCR analysis**

Total RNA from the parasite was isolated by Trizol method (11) RNA was treated with DNase I (Fermentas) to avoid DNA contamination. Synthesis of cDNA was performed by using Superscript II invitrogen

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**Table 1. List of primers**

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Kit as per the manufacture’s instruction. RNA without reverse transcriptase was considered as negative control during cDNA synthesis. PCR was carried out with 4 µL cDNA sample or negative control by using primers for PiGAPDH (P6–P7) or PiSSB (P4–P5). PCR products were further analysed by agarose gel electrophoresis.

Sucrose gradient ultracentrifugation

Sucrose gradient ultracentrifugation assay was done following the protocol described by Choudhury et al. (24). Recombinant PiSSB (150 µg) or 150 µg P. falciparum whole cell extract (in lysis buffer) obtained from 50 µl mixed stage parasite pellet or different marker proteins (~200 µg) were layered on top of 4.9 ml 10–40% (w/v) sucrose step gradient in a buffer containing 25 mM Tris–HCl (pH 7.5), 250 mM NaCl, 2 mM sodium bisulphate and 0.05% Triton X-100. Samples were centrifuged in a Beckman MLS-50 rotor at 35 000 r.p.m. for 18 h at 4°C. Several fractions (~170 µl) were collected for each protein. Fractions of recombinant protein and marker proteins were subjected to 10% SDS–PAGE analysis where as fractions of parasite extract were TCA precipitated and subjected to western blot analysis by anti-PfSSB antibodies. Standard curve was generated using Microsoft excel application program by plotting molecular mass as the function of sedimentation distance for various marker proteins, e.g. ovalbumin (43 kDa), aldolase (158 kDa), catalase (232 kDa), ferritin (440 kDa) and thyroglobulin (669 kDa) (Amasharm Biosciences, USA). Regression analysis using the Microsoft excel application programme yielded the equation y = 2.0155x + 0.108. ’y’ represents the log of the molecular mass in kDa and ‘x’ represents log of fraction number. The peak fraction numbers of recombinant PiSSB (fraction no. 9) and endogenous PiSSB (fraction no. 8) were fitted to the standard curve and relative molecular masses were estimated.

Dynamic light-scattering study of Pf SSB

The particle size measurement was performed by a dynamic light scattering (DLS) technique (mostly at scattering angle = 90° and laser wave length = 660.00 nm) on a digital correlator (Photocor instrument, USA). Scattered light obtained from the sample kept in a quartz cuvette was detected by a photo multiplier tube (Hamamatsu). The signal was converted to an intensity autocorrelation function by a digital correlator. The data analysis was done using CONTIN software provided by Brookhaven instrument. Further details about DLS and data analysis can be found elsewhere (25).

Fluorescence and confocal microscopy

Smears of parasitized RBC of different developmental stages were made and fixed in chilled methanol. Blocking was done by incubating the slide with a solution containing 3% BSA and 0.05% saponin in a humidified chamber for 2 h at 37°C. The slides were washed with 1× PBS and incubated with anti-PiSSB/anti-PfACP antibodies at 1:2000 dilutions for overnight at 4°C. After washing the slides thoroughly with 1× PBS, they were again incubated with 1× PBS containing anti-mouse/anti rabbit secondary antibodies and DAPI (Molecular probe, Invitrogen) for 1 h at room temperature. Finally, the slides were washed and mounted with anti-fed reagent and viewed under 100× oil immersion objective. Confocal images were collected using Laser scanning Olympus microscope or Carl Zeiss Apotome fluorescence microscope and the figures were prepared using Adobe photoshop.

Complementation assay—plasmid bumping assay

For complementation assay, E. coli RDP317 strain (a kind gift from Dr Umesh Varshney, Indian Institute of Science, Bangalore, India) was used. In this strain, the chromosomal ssb gene is replaced by a kanamycin resistance (ssb::kan) marker and it harbours a support plasmid pRPZ150 (ColEl ori, TcR) coding wild-type SSB protein. The complementation assay was performed essentially following the protocol described elsewhere (26) with some modifications as described in the Supplementary Data (‘Materials and Methods’ section).

Metabolic labelling of parasite proteins

Synchronized parasites (~12–14 h post-invasion) were treated with ciprofloxacin (10 µM). Untreated and ciprofloxacin treated trophozoite stage parasites during second life cycle (~68 h post drug treatment) were incubated with 35S-methionine (specific activity = 15 mCi/ml, added to a final concentration of 150 µCi/ml) for 3 h. The parasites were then released by saponin lysis, washed with 1× PBS and subsequently lysed in lysis buffer (30 mM Tris–HCl, pH 8.0, 300 mM NaCl, 1 mM EDTA, 0.7% Triton X, Protease inhibitor cocktail). The lysate was used for immunoprecipitation assay with anti-PiSSB and anti-PfActin antibodies or respective pre-immune sera as described earlier (10). Immunoprecipitated samples were further subjected to analysis by SDS–PAGE and autoradiography.

RESULTS

PiSSB is constitutively expressed in intra-erythrocytic life cycle of P. falciparum and it is targeted to the apicoplast

PLASMODB contains an ORF (PFE0435c) that shows 39% identity and 66% homology with E. coli SSB (Figure 1A). It has a long stretch of 76 amino acid residues at the N-terminus that shows putative apicoplast targeting sequence. PiSSB also contains an extension of 28 amino acid residues at the C-terminus that does not show homology with EcSSB. Overall, the N-terminal ssDNA binding region (excluding the apicoplast targeting sequence) shows more homology than the C-terminal protein–protein interacting domain (excluding the extreme C-terminal 28 residues) that is rich in asparagine residues in contrast to the similar domain of EcSSB that is rich in glycine residues (Figure 1A). Analysis of primary amino acid sequence of PiSSB reveals evolutionary conserved oligosaccharide/oligonucleotide binding domain (from amino acid 80–184) (OBD) implicated in ssDNA binding. The three tryptophan residues...
Figure 1. Amino acid sequence alignment, protein expression profile and immunolocalization of PfSSB. (A) Clustal W amino acid sequence alignment of PfSSB with E. coli SSB (EcSSB). Identical residues are marked with ‘asterisk’ whereas ‘colon’ and ‘end note’ signs are used to indicate strongly and weakly similar residues, respectively. (B) Coomassie-stained gel picture of induction and purification of His6-tagged Pf SSB. Protein markers (M) are indicated on the right. (C) The right panel shows the western blot analysis using pre-immune (PI) or immune (I) sera against PfSSB (lanes 1 and 2) against 3D7 parasite lysate. The left panel shows the coomassie stained gel following transfer of the proteins on the membrane (loading control) (D) Expression profile of PfSSB at the different erythrocytic stages as indicated on top is shown by western blot analysis using polyclonal antibodies against PfSSB. The same blot was reprobed with antibodies against PfActin (middle panel). The bottom panel shows the coomassie stained gel following transfer analysis using polyclonal antibodies against SSB. Arrowhead indicates the position of PfSSB protein. (E) Immunoprecipitation of PfSSB protein from parasite lysate using immune (I) or pre-immune sera (PI) followed by western blot analysis using polyclonal antibodies against SSB. Arrowhead indicates the position of PfSSB protein. (F) Localization of PfSSB by immunofluoroscence analysis using glass slide containing fixed parasites obtained from trophozoite and schizont stages in the presence of anti-PfSSB antibodies. DAPI indicates nuclei whereas merged panel shows the presence of PfSSB with respect to the nuclei. The last panel shows the phase picture of parasite infected RBC. (G) Immuno-colocalization of PfSSB and apicoplast marker acyl carrier protein (ACP). Merged panel shows the co-localization of both the proteins.
and His6-PfSSB protein was purified using Ni–NTA sequence reported in PLASMODB.ORG. BL21 codon genomic DNA from 3D7 strain of \( P. falciparum \) by using specific primers (as shown in Table 1) and cloned ssb was further sequenced completely. The sequence of the expression vector pET28a and the recombinant construct amplified PCR product was subsequently cloned in the between these two proteins.

In order to investigate whether PfSSB is expressed in the parasites, western blot analysis was performed using parasite lysate obtained from mixed stage parasites in the presence of pre-immune and immune sera against PfSSB. A specific band (~30 kDa) was obtained only in the immune sera treated lane and not in the pre-immune sera treated lane (right panel, Figure 1C). The left panel shows commassie stained gel following transfer of the proteins onto the membrane as loading control. To crosscheck the specificity of the anti-PfSSB serum, immunodepletion assay was performed. For this purpose, anti-PfSSB sera were incubated overnight with glutathione sepharose 4B beads containing GST proteins or GST-PfSSB proteins. GST or GST-PfSSB treated sera were further immunodepleted using fresh GST or GST-PfSSB proteins containing beads for another 8h. These different immunodepleted sera and untreated sera were used for western blot analysis against parasite lysate obtained from mixed stage parasites. We find that both untreated and GST protein depleted sera show a strong band corresponding to PfSSB. However, the intensity of the PfSSB band was reduced drastically when GST-PfSSB immuno-depleted sera were used for western blot analysis (Supplementary Figure S1A). The same membranes were treated with anti-PfActin sera resulting in a strong band corresponding to PfActin in all the lanes confirming the loading of equivalent amount of proteins in all the lanes (Supplementary Figure S1A). The immunodepletion results clearly indicate the specificity of the anti-PfSSB sera. Further, the expression pattern of the endogenous protein at the different developmental stages were investigated by western blot analysis using anti-PfSSB antibodies against parasite lysate obtained from synchronized ring, trophozoite and schizont stage parasites. A specific band (~30 kDa) was obtained in all the stages suggesting that PfSSB is expressed during all the stages of the asexual developmental cycle (Figure 1D). The deduced molecular mass of PfSSB is ~24.6 kDa. We find that PfSSB runs anomalously in SDS–PAGE consistent with many other \( Plasmodium \) proteins (10). As a loading control, the same blots was used to track the expression of PfActin at different stages (Figure 1D). To further investigate the authenticity of the band obtained in western blot analysis, immunoprecipitation experiment was performed using parasite lysate and pre-immune and immune sera against PfSSB. Western blot analysis following immunoprecipitation indicates the presence of a specific band in the immune sera precipitated lysate but not in pre-immune sera treated samples (Figure 1E). These results truly reflect the endogenous expression of PfSSB and the specificity of the antibodies.

We further investigated the subcellular localization of PfSSB during asexual developmental cycle by immunofluorescence assay using antibodies against PfSSB. Distinct subnuclear spot corresponding to apicoplast was observed in late ring/trophozoite stage parasites with undivided nucleus (Figure 1F, first row, top panel) whereas multiple subnuclear spots were observed in multinucleated schizont stage parasites (Figure 1F, bottom two rows, top panel). These results are consistent with the multiplication of apicoplast organelle with parasite growth. To further confirm whether PfSSB is localized in the apicoplast organelle of the parasite, immuno-colocalization experiment was performed using antibodies against \( P. falciparum \) ACP (a marker for apicoplast) and PfSSB (11). We find that PfACP is localized in the sub-nuclear apicoplast compartment. Interestingly, PfSSB signal was completely merged with the PfACP signal suggesting that PfSSB is truly localized in the apicoplast (Figure 1F, the bottom panel).

Oligomeric status PfSSB in vitro and in vivo and homology modelling

EcSSB forms homo-tetramer in solution (28). It will be interesting to see the oligomeric status of recombinant PfSSB and endogenous PfSSB from parasite lysate in solution. For this purpose, both recombinant PfSSB and \( Plasmodium \) lysate from mixed stage parasites were subjected to sucrose density gradient centrifugation as described in the ‘Materials and Methods’ section. Different molecular mass markers were also used for centrifugation under the same experimental conditions. Several fractions were collected for each molecular mass marker and peak fraction was identified (shown with arrow heads on top, Figure 2A). Subsequently, the peak fraction of recombinant PfSSB was identified by SDS–PAGE and coomassie staining of the gel containing different fractions following centrifugation (Figure 2A, top panel). The peak fraction of endogenous PfSSB was identified by western blot analysis of each fraction using anti-PfSSB antibodies (Figure 2A, bottom panel). A standard curve was plotted using the log fraction number (peak fraction) against molecular mass of the standard proteins (Figure 2B). This helped us to estimate the molecular mass of recombinant PfSSB (~104 kDa) and native PfSSB (~92 kDa). The apparent
difference in the calculated molecular masses between the recombinant and endogenous proteins could be due to the presence of six histidine residues and some extra amino acid residues in recombinant PfSSB contributed by the expression vector.

Comparison of the calculated molecular mass of monomeric form of recombinant PISSB (25.2 kDa) and endogenous PfSSB (24.6 kDa) with the estimated molecular mass obtained from sucrose gradient experiments revealed that both the recombinant PISSB and endogenous PfSSB would be in tetrameric form.

Further, DLS experiment was performed to evaluate whether tetrameric form of PISSB were present predominantly in the solution. For this purpose, SSB protein from gastric pathogen Helicobacter pylori was taken as positive control. The molecular mass of HpSSB is ~25 kDa (close to ~His6-PfSSB, ~25.2 kDa) and it has been shown recently that HpSSB exists as tetramer in solution (26). DLS allows us to study the average particle size (hydrodynamic diameter) and its distribution in solution. The average hydrodynamic diameter of Pf SSB was found to be 152.1 ± 2.65 nm which is quite close to the hydrodynamic diameter 139.1 ± 4.12 nm of HpSSB (Figure 2C). Thus, it can be inferred that majority of PISSB remains as tetramer as HpSSB in solution. These results also reconfirm sucrose density gradient ultracentrifugation results.

Consistent with the tetrameric form of PISSB as obtained using various methods described earlier, we performed the homology modelling of the protein using EcSSB structure as template (29). Monomer model of putative Oligo nucleotide Binding domain of PISSB (80–193 aa) is modelled using Modeller 9v7 software with the Oligo nucleotide binding domain of EcSSB (ssDNA bound subunit A1001–1112 of PDBID:1EYG) as a template since it has significant sequence homology.
Modelled monomer of PfSSB was superimposed using Dali pairwise structure alignment program (http://ekhidna.biocenter.helsinki.fi/dali/start) onto individual monomer of tetramer (A–D subunits in PDBID:1EYG) of EcSSB to generate the tetramer of PfSSB with root mean square deviation (RMSD) of 1.8, 1.6, 2.0 and 1.8 Å, respectively. The model revealed three conserved tryptophan residues (Trp-117, Trp-131 and Trp-166) which are essential for ssDNA interaction. His 55 residue in EcSSB structure where it has been implicated for oligomerization of SSB tetramer is also conserved in PfSSB (His132) (29). The homology model of PfSSB homotetrameric oligo binding domain is shown in Figure 3 where different monomers (chains/subunits) (A–D) are shown in different colours using Pymol software. A homology model was also generated using tetrameric form of PfSSB along with two stretches of ssDNA (chains A and B) that shows the interaction of ssDNA with the oligonucleotide binding domain of PfSSB (Supplementary Figure S1B).

**Complementation of E. coli ssb mutant cells with wild-type and deletion mutant of PfSSB**

In order to investigate whether PfSSB is a true homologue of SSB in vivo, we performed genetic complementation experiments using an E. coli ssb mutant strain with the help of plasmid bumping experiment using EcSSB and H. pylori SSB as positive control (26). The details of this method are described in the ‘Materials and Methods’ section and the results are shown in Table 2. We found that both EcSSB and HpSSB could complement this mutant strain whereas PfSSB could not complement under the same experimental conditions. Further, a construct deleting the unique extreme C-terminal 28 residues present in PfSSB also cannot complement the E. coli ssb mutant strain suggesting that PfSSB is inherently different from EcSSB. It has been reported earlier that the extreme 10 amino acid residues of EcSSB are essential for its in vivo function (30). Poor homology at the extreme C-terminal regions of these two proteins may contribute to the inability of PfSSB to complement E. coli ssb mutant strain.

**In vitro and in vivo ssDNA binding activity of PfSSB**

During DNA replication, recombination and repair, SSB protein stabilizes the ssDNA intermediates. ssDNA binding activity of purified recombinant PfSSB was monitored using agarose gel mobility shift assay in the presence of a mixture of M13 ssDNA and pUC18 dsDNA. The results indicated that the mobility of single-stranded M13 DNA was retarded significantly with increasing amount of PfSSB protein compared to pUC18 DNA whose mobility did not change at all (Figure 4A). Further, gel mobility shift assay was performed using

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<td>pTRC+PfssbΔC28</td>
<td>40</td>
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**Figure 3.** Homology modelling of PfSSB. Pf SSB structure was modelled using the crystal structure of oligonucleotide binding domain of E. coli SSB as template. The positions of the conserved amino acid residues (Trp-117, Trp-131, Trp-167 and His-132) are shown in the figure. Four monomers of SSB that form the tetramer are shown in different colours (chains A–D). The model also shows two ssDNA molecules (A and B) interacting with the tetramer model.
Figure 4. In vitro and in vivo DNA binding property of Pf SSB. (A) PfSSB binds specifically to ssDNA. Different amount of Pf SSB was incubated with the mixture of M13 mp18 ssDNA and pUC18 dsDNA followed by separation of the protein-DNA complex by agarose gel electrophoresis. ssDNA was progressively retarded (asterisk) with increasing concentration of PfSSB whereas mobility of dsDNA was not affected. (B) Electrophoretic mobility shift assay using short radiolabeled single-stranded oligonucleotide probe with increasing amount of Pf SSB. The arrowhead indicates the protein bound DNA. (C) Gel retardation analysis of single stranded DNA binding activity from P. falciparum cell extract. Labelled ssDNA probe was incubated with P. falciparum cell extract in the absence or presence of anti-PfSSB or anti-PCNA antibodies or pre-immune sera against SSB followed by PAGE analysis of the protein–DNA complexes (shown by arrowhead). Supershift of the DNA–protein complex was observed only in the presence of immune sera against PfSSB (lane 4) but not in the presence of pre-immune sera (lane 3) or non-immune sera (anti-PCNA) (lane 5). (D) Single-stranded DNA binding activity of cytoplasmic or nuclear extract or immunodepleted samples from each fraction in the presence of preimmune and immune sera against PfSSB. A single shifted band of labelled probe was observed in cytoplasmic fraction (lane 2) whereas two such bands were observed in nuclear extract treated samples (lane 3). No band was observed in the presence of immunodepleted cytoplasmic extract (lane 5) whereas the bottom band was not visible in the presence of immunodepleted nuclear extract treated sample (lane 7). Pre-immune depleted samples did not affect the pattern of bands (lanes 4 and 6) found in extract only lanes (lanes 2 and 3). (E) The upper part of the left panel shows the western blot analysis of immune and pre-immune depleted Plasmodium cytoplasmic extract whereas the lower part represents the coomassie stained gel as loading control following transfer of the proteins on the membrane. Similarly, the upper part of the right panel shows the western blot analysis of the immunodepleted nuclear extract (immune and pre-immune treated) and the lower part of the same panel represents the coomassie stained gel as loading control. The arrowhead indicates the position of the PfSSB protein. (F) Specific binding of PfSSB to apicoplast DNA in ChIP assay. PCR amplification of ori region of apicoplast (api.) DNA was found only in the immunoprecipitated samples using immune sera (I) against PfSSB but not with the pre-immune (PI) sera whereas nuclear (nuc.) ssb gene could not be amplified using ssb gene specific primers (set 1). Further, PCR amplification of immunoprecipitated DNA along with input (inp.) genomic DNA were performed using specific primer sets from nuclear orc5, apicoplast ori and IRA region (set 2) or nuclear sir2 and apicoplast tufA regions (set 3). Specific products were found only in the immunoprecipitated samples using immune sera and primer sets from apicoplast DNA. No significant amplification was found using primer sets from nuclear DNA. PCR products were obtained in all the control input lanes.
radioactive ssDNA probe and purified PfSSB. We find similar results as above where increasing amount of PfSSB results major shift of the ssDNA probe (Figure 4B). We also investigated whether *P. falciparum* extract would contain ssDNA binding activity contributed by PfSSB. For this purpose, radiolabelled ssDNA probe (70 mer) was incubated with *P. falciparum* parasite extract followed by PAGE analysis of the reaction mixture. We find that *P. falciparum* extract cause a considerable shift of the ssDNA probe (lane 2, Figure 4C). To identify whether the binding is specific for PfSSB, the reaction mixture was incubated further in the presence of undiluted anti-PfSSB or pre-immune sera as described in the ‘Materials and Methods’ section. We find, while pre-immune sera do not affect the ssDNA binding activity, immune sera causes a supershift suggesting that the binding is specific for PfSSB (lanes 3 and 4, Figure 4C). Non-immune PfPCNA antibodies also do not affect the ssDNA binding activity further confirming the presence of specific PfSSB like ssDNA binding activity in the *P. falciparum* extract.

Specific interaction of the high molecular mass containing tetrameric form of PfSSB with high titer antibodies against PfSSB may form huge complex causing supershift of the DNA–protein complex that may not enter the polyacrylamide gel properly as shown above. The supershift experiments were repeated with some modifications like using different concentrations of diluted PfSSB antibodies (1–5 µl of 1:1000 diluted anti-PfSSB sera in 1× PBS) and resolving the DNA–protein complex for a longer time during PAGE analysis. We find that pre-immune sera against PfSSB do not show any supershifted band even at the highest concentration whereas immune sera show specific supershifted bands at all concentrations. However, with increasing concentration of immune sera, the intensity of the original DNA–protein complex is reduced significantly with the appearance of a higher molecular mass containing population that does not enter the gel properly apart from the supershifted band that entered the gel (Supplementary Figure S2). The complex that does not enter the gel may be the result of the accumulation of huge complex containing ssDNA–Protein-Ab at the higher concentration of antibodies.

To confirm whether the ssDNA binding activity present in *P. falciparum* parasite extract is due to the presence of PfSSB, sub-cellular fractionation of the parasite extract was performed to obtain cytoplasmic extract and nuclear extract as described earlier (31). Each fraction was immunodepleted using immune and pre-immune sera against PfSSB. Immunodepleted cytoplasmic and nuclear extract were used for gel retardation assay using labelled ssDNA probe. We find that cytoplasmic extract yields one shifted band whereas nuclear extract results in two shifted bands. An extra band was obtained on top of the band corresponding to the band obtained from the cytoplasmic extract (lanes 2 and 3, Figure 4D). Immunodepleted cytoplasmic extract using PfSSB immune sera resulted in complete loss of DNA binding activity (lane 5). Interestingly, immunodepleted nuclear extract resulted in the loss of band corresponding to the band obtained in cytoplasmic extract. However, the upper band obtained in nuclear extract was not affected following immunodepletion suggesting the shift represented by upper band would not be due to the presence of PfSSB (lanes 6 and 7, Figure 4D). This ssDNA binding activity could be due to the presence of specific nuclear protein present in the nuclear extract. This could be due to the presence of RPA in the nuclear fraction as it has been reported earlier (20). Immunodepletion of cytoplasmic extract was further confirmed by western blot analysis of pre-immune and immune sera depleted cytoplasmic extract using anti-PfSSB antibodies (Figure 4E, left panel). The results indicate the presence of a strong band corresponding to PfSSB in the pre-immune sera depleted sample compared to the very weak band present in the immune-sera depleted sample. The similar loading in both the lanes is indicated by the coomassie stained gel following transfer of the proteins onto the membrane. Immunodepletion of nuclear extract using pre-immune sera and PfSSB antibodies followed by western blot analysis indicate the presence of PfSSB in the nuclear extract possibly as a contamination of nuclear fraction with cyttoplasmic proteins (Figure 4E, right panel) that may have resulted the appearance of lower band in the gel shift assay using nuclear fraction as shown earlier. These results indicate that the specific immunodepletion of PfSSB from the cytoplasmic extract may have caused the inability of the immunodepleted extract to bind to ssDNA.

We further investigated whether PfSSB binds specifically to apicoplast DNA *in vivo*. For this purpose, ChIP experiments were performed using pre-immune or immune sera against PfSSB. Recently a similar approach has worked nicely to show that PfGyrB binds specifically to its target apicoplast DNA with no affinity towards nuclear DNA (22). ChIP results indicate that PfSSB binds strongly to the apicoplast ori region compared to the nuclear ssb gene as control (Figure 4F, set 1). ChIP experiments were repeated several times followed by PCR using primer sets from different nuclear loci and apicoplast loci. We find that the intensity of the PCR product using primer set from apicoplast DNA is always higher many fold than that obtained from nuclear DNA in different sets of experiments (Figure 4F, sets 2 and 3). These results indeed suggest that PfSSB binds specifically to apicoplast DNA as compared to nuclear DNA.

**Modulation of PfSSB expression and activity in the presence of inhibitors against apicoplast machinery**

*Plasmodium* contains both the subunits of prokaryote type gyrase (a and b) that are nuclear encoded but targeted to the apicoplast (11). Gyrase is required for the maintenance of the apicoplast DNA as positive supercoils are generated ahead of the replication fork that should be relieved intermittently for the faithful replication of apicoplast DNA. It has been shown earlier that ciprofloxacin, a gyrase A specific inhibitor affects the apicoplast DNA replication specifically without affecting nuclear DNA replication (32). Interestingly, ciprofloxacin shows typical delayed death phenotype of the parasites where parasitemia drops significantly during the second life cycle of the parasites.
following the addition of the drug without affecting the parasites during the first life cycle (13). We were interested to study whether ciprofloxacin would modulate PfSSB function since it is also a key molecule for apicoplast DNA replication.

We first investigated the effect of ciprofloxacin on parasite growth during first and second life cycle of parasite development. Although ciprofloxacin caused ~40% decline in parasitemia at 15 μM concentration, it did not cause any significant effect at lower drug concentration (5/10 μM) at the end of the first life cycle (~48 h). However, at the end of the second life cycle (~92 h), there was a dramatic drop in parasitemia at 5/10 μM drug concentration that resembled the delayed death effect of ciprofloxacin as described earlier (Figure 5A) (13).

It has been reported earlier that drugs that affect apicoplast housekeeping functions (e.g. tetracycline), inhibit the translocation of the proteins to the apicoplast (during second life cycle) using signal sequence of ACP fused to GFP as marker (14,15). No endogenous apicoplast targeted protein has been tracked in the above studies. We were interested to investigate whether treatment of the parasites with ciprofloxacin would modulate PfSSB translocation and function. Subsequently, we performed western blot analysis using ciprofloxacin (10 μM) treated parasite lysate obtained from first life cycle (~34 h post-drug treatment) and second life cycle (~82 h post-drug treatment) by using antibodies raised against PfSSB. Antibodies against PfActin and PIPCNA were used as control for western blot experiments. We find that during the first life cycle, addition of ciprofloxacin does not change the expression of PfSSB protein compared to the untreated parasites (Figure 5B). However, during second life cycle, the expression of PfSSB is found to be reduced in ciprofloxacin treated parasites compared to the untreated parasites. In contrary, the expression level of control protein PfActin and nuclear replication protein PIPCNA did not

![Figure 5](https://academic.oup.com/nar/article-abstract/38/20/7037/1303712)
change significantly. Moreover, a higher molecular weight band is observed in drug treated samples that is absent in the untreated samples (Figure 5B). This may be the unprocessed form of PfSSB before the cleavage of signal peptide and transit peptide required for the faithful translocation of apicoplast proteins. Apicoplast specific drugs often result in accumulation of the immature form of the apicoplast targeted proteins during second life cycle (15). However, the decrease in overall PfSSB protein level during second life cycle following ciprofloxacin treatment is very striking.

The previous experiment did not allow us to investigate the timing of the effect of ciprofloxacin on PfSSB protein level during second life cycle. For this purpose, ciprofloxacin treated or untreated parasite pellets were obtained from the trophozoite (~68 h post-drug treatment) and schizont stages (~82 h post-drug treatment) during the second life cycle followed by analysis of the PfSSB protein level by western blot analysis using anti-PfSSB antibodies (Figure 5C, left panel). Parasite lysate from the first life cycle was also taken as control. We find that the expression level of PfSSB protein is reduced even at the trophozoite stage that continues until schizont stage during second life cycle whereas the PfActin level does not change at all. A faint band corresponding to unprocessed form of PfSSB was also visible at the latter stage.

We further show that the decrease in overall PfSSB protein level and the appearance of the mature form of PfSSB following ciprofloxacin treatment is specific for apicoplast targeted drugs. Use of increasing amount of artemisinin that cause considerable parasitic death resulting in huge decrease in parasitemia does not affect the level of PfSSB at all. Neither the mature form of PfSSB could be detected even at the highest concentration of artemisinin used (30 nM) (Figure 5C, right panel).

We are interested to find out whether modulation of PfSSB expression and maturation is ciprofloxacin-specific or it is a manifestation of apicoplast targeted drugs to modulate apicoplast targeted proteins in general. For this purpose, we performed western blot analysis using ciprofloxacin and tetracycline (protein translation inhibitor that also affects apicoplast) treated parasite extract in the presence of PfSSB and PfHGU (histone like protein) antibodies. Western blot analysis indicates that the treatment of ciprofloxacin and tetracycline affects the expression and maturation of both PfSSB and PfHGU protein to the similar extent (Figure 5D). However, these drugs did not have any effect on control protein PfActin.

Further, to investigate whether ciprofloxacin truly affects the translocation of apicoplast targeted proteins like PfSSB during second life cycle, we have performed immunofluorescence assay using untreated and drug treated parasites in the presence of PfSSB antibodies. We find specific apicoplast localized PfSSB signal in the untreated parasites whereas the apicoplast localization of PfSSB is severely affected in the drug-treated parasites under the same experimental conditions (Supplementary Figure S3). These results strongly suggest that ciprofloxacin inhibits the translocation of SSB during second life cycle.

The above results prompted us to investigate whether parasite extract obtained from ciprofloxacin treated parasites would yield ssDNA binding activity. This would also help us to confirm whether the ssDNA binding activity from the parasite extract is truly dependent on PfSSB. Gel retardation assay was performed using parasite extract obtained from ciprofloxacin treated and untreated control parasites. We find a strong shift of the radiolabelled probe in the presence of untreated parasite extract compared to the free probe. Interestingly, ciprofloxacin treated parasite extract fails to show any prominent shifted band suggesting that the inability of ssDNA binding may be attributed to the poor presence of the mature form of PfSSB in ciprofloxacin treated samples (Figure 5E, right panel). Western blot analysis using extract obtained from ciprofloxacin treated and untreated parasites in the presence of PfSSB antibodies further confirm the above results (Figure 5E, left panel).

**Synthesis of SSB protein is affected during second life cycle following ciprofloxacin treatment**

We find that apicoplast targeted drug, ciprofloxacin affects the translocation of the proteins to the apicoplast as evidenced by the accumulation of the unprocessed form of the protein in the presence of the drug. It is also interesting that there is an overall decrease (~4-fold) in PfSSB protein level compared to control protein PfActin following drug treatment (Figure 6A). Transcription of ssb is up-regulated in *E. coli* cells in the presence of ciprofloxacin (33). It is possible that the decrease in SSB protein level following ciprofloxacin treatment during second life cycle is caused by the reduced transcription of *Pfssb* in the nucleus. To address this issue, we performed semi-quantitative RT-PCR analysis using cDNA prepared from ciprofloxacin treated and untreated samples. We find that there is no significant difference in the transcript level of *ssb* in the presence or absence of ciprofloxacin (Figure 6B). *PfGAPDH* was used as control that also did not change its expression as *ssb*. These results indicate that down-regulation of SSB protein in the ciprofloxacin treated parasites may not be due to the transcriptional defect. This is consistent with the earlier published results where the microarray analysis of tetracyclin treated parasites do not show change in transcription level of nuclear genes (including genes required for nuclear encoded apicoplast-targeted proteins) compared to the untreated parasites (14). The other possibility is that SSB protein was degraded with time leading to the decrease in protein level. We used MG132, a proteosomal degradation pathway inhibitor during the second life cycle following ciprofloxacin treatment. Recently, we have shown that MG132 can stabilize PfORC1 by inhibiting degradation of PfORC1 at the late schizont stage (34). Western blot analysis using parasite lysate obtained from 100 nM MG132 treated and untreated samples (during second cycle following ciprofloxacin treatment) reveals that MG132 does not change the SSB protein level (Figure 6C) suggesting that the decrease in SSB protein level would not be due to the degradation of the protein. MG132 was active in the parasites as addition of similar
concentration of drug could stabilize PfORC1 from degradation at the late schizont stage as shown by immunofluorescence and western blot analysis (Supplementary Figure S4A and B).

These results suggest that the decrease in the SSB protein level during the second life cycle following ciprofloxacin treatment might be attributed to the translational defect. To address this issue further, ciprofloxacin-treated or untreated parasites were incubated in the presence of $^{35}$S-methionine during second life cycle (trophozoite stage, $\sim$68 h post-drug treatment) for 3 h. The parasite lysate were obtained from both control and drug-treated parasites followed by SDS–PAGE analysis and autoradiography to visualize the labelled proteins. The results indicate that there is no gross change in the metabolic labelling of the proteins during second life cycle in the presence or absence of the drug (Figure 6D, left panel). The above lysate were used to immunoprecipitate newly synthesized SSB protein during second life cycle using pre-immune and immune sera against SSB. We find that SSB can be immunoprecipitated specifically using immune sera but not pre-immune sera during second life cycle from the untreated sample only (Figure 6D, right panel). Interestingly, PfActin can be immunoprecipitated from both the untreated and drug-treated sample to the similar extent using specific antibodies against PfActin under the similar experimental conditions (Figure 6D, middle panel). We also performed immunoprecipitation experiments using pre-immune and immune sera against PfHU under the same experimental conditions. We find that the translation of newly synthesized PfHU protein during second life cycle...
following drug treatment is also significantly reduced compared to the untreated samples (Figure 6D, bottom panel). These results suggest although there is no significant change in protein expression profile during trophozoite stage of the second life cycle between drug-treated and untreated samples, there is a specific down-regulation of translation of nuclear encoded apicoplast targeted proteins like SSB and HU during second life cycle following ciprofloxacin treatment. It is important to note that all apicoplast targeted proteins may not be affected to the similar extent at a particular time since there may be additional mechanisms to control their expression profile that need to be explored further.

**DISCUSSION**

The replication of apicoplast DNA circle begins in trophozoite stage and continues into schizogony (5). Very little is known about the process and enzymology involved in the replication of apicoplast genome. So far, few enzymes including gyrase and PfPRex have been reported that could be the members of active replication/recombination machinery within the apicoplast. This study is an attempt to identify and functionally characterize another key DNA replication protein, the ssDNA binding protein in *P. falciparum*. To our knowledge, the identification of SSB in *P. falciparum* is the first report of the presence of bacterial type SSB in any eukaryotic parasite. In *E. coli* and other prokaryotes, in addition to DNA replication the role of SSB has been extended to DNA recombination and repair events. PISSB may be involved in these processes to maintain the apicoplast genome.

It is known that SSB protein binds to circular supercoiled DNA containing ssDNA stretches that have been induced by topological stress generated by different cellular processes like DNA replication, transcription and recombination (35). Our data suggest that PISSB is exclusively targeted to the apicoplast and it binds to the apicoplast DNA and not to the nuclear DNA. Together, these data confirm apicoplast specific role of PISSB. The apicoplast DNA replication occurs in the late trophozoite-early schizont stages. However, PISSB is constitutively expressed and localized into the apicoplast during all the three stages of intraerythrocytic developmental cycle of *P. falciparum*. This may explain the possible involvement of PISSB in different DNA transactions other than DNA replication during the non-replicative phases of intra-erythrocytic parasitic life cycle (ring, early trophozoite).

Earlier, ssDNA binding activity from *P. falciparum* had been assigned to large subunit of RPA. The ssDNA binding activity exhibited by RPA has been shown to be present in both nuclear and cytoplasmic fractions. However, specificity of the ssDNA binding activity could not be tested due to the non-availability of specific antibodies against RPA. In contrast, we show that the ssDNA binding activity from cytoplasmic fraction is largely contributed by PISSB. The loss of ssDNA binding activity in immunodepleted cytosolic fraction using specific anti-PISSB antibodies strongly substantiates our claim.

There are two ssDNA binding activities in nuclear fraction and only one of them is contributed by PISSB. We show that upon immunodepletion of PISSB from the nuclear extract, the ssDNA binding activity contributed by PISSB is completely lost whereas the other activity remains unaffected. The unaffected ssDNA activity in the nuclear extract may probably be due to RPA as reported earlier (20). However, we cannot rule out the ssDNA binding activity by PISSB in the nuclear extract by cross-contamination of cytoplasmic PISSB during the cell fractionation because our ChIP experiment using specific anti-PISSB antibodies yield robust amplification of apicoplast DNA only and no significant amplification of nuclear DNA.

Generally, SSB modulates the functions of other replication/recombination/repair proteins by interacting with them via acidic C-terminus region. It has been reported that the last 10 amino acid residues of SSB are highly conserved among different bacteria and they are essential for *in vivo* protein–protein interaction (30). The C-terminus of PISSB shows little sequence conservation and there is an extension of 28 amino acid residues. Poor sequence conservation at the C-terminus of PISSB may explain its failure to complement *E. coli* ssb function *in vivo*. The identification of the proteins interacting with PISSB C-terminus would be an essential step towards understanding the role of PISSB during different developmental stages of the parasite erythrocytic life cycle, particularly during the ring stage where DNA replication does not take place.

PISSB forms tetramer in solution similar to EcSSB. His-55 and the region between amino acid residues 89–105 have been suggested to have a role in tetramerization of EcSSB. In PISSB, His-132 is equivalent to His-55 of EcSSB and the region between 165 and 194 shows a significant homology with the region 89–105 of EcSSB. These similarities may explain the tetrameric nature of the protein both *in vitro* and *in vivo* as shown here. Our ongoing crystallization studies of PISSB may provide a clear insight into oligomerization interfaces of the protein.

Ciprofloxacin is a quinolone antibiotic that principally targets DNA gyrase and results in the breakage of the *P. falciparum* apicoplast DNA without affecting the nuclear DNA during second life cycle at an IC$_{50}$ of $\sim$7 µM. Surprisingly, our results indicate that blocking apicoplast replication with ciprofloxacin results in the drastic reduction in the levels of PISSB and its relocation to the apicoplast as evidenced by the accumulation of PISSB with the signal peptide sequence. However, ciprofloxacin treatment does not change the levels of cytoplasmic actin and nuclear replication protein PCNA1. The level of PISSB is not affected when the parasites are treated with commonly used non-apicoplast targeted anti-malarial drug artemisinin, suggesting that the drug stress does not lead to the decrease in PISSB levels. However, treatment of the parasites with tetracycline that affects apicoplast translation also modulates PISSB expression and maturation like ciprofloxacin suggesting...
that these drugs may follow general mechanism that interfere with the translocation of the proteins to the apicoplast and the protein level. The similar fate of HU, another protein targeted to the apicoplast, following ciprofloxacin and tetracyclin treatment further confirms the generalized mechanism of apicoplast targeted drugs. Simultaneous treatment of parasites with ciprofloxacin and proteosome blocker MG132 clearly indicates that protein is not degraded in the presence of ciprofloxacin during the second cycle. Moreover, the inhibition of apicoplast DNA replication does not affect the level of the ssb transcripts produced in the nucleus housing ssb gene on ChrV as shown by the semi-quantitative RT-PCR analysis. Incubation of the parasites with 35S-methionine to track the newly synthesized proteins during second life cycle suggests that translation of SSB and HU transcripts are specifically affected during second life cycle following drug treatment although both untreated and drug-treated parasites show comparable protein expression profile and there is no change in expression level of control PfActin protein. Like PfSSB, the expression level of newly synthesized PfHU protein during the second life cycle following drug treatment was severely affected. It seems that apicoplast targeted drug ciprofloxacin may selectively inhibit the translation of apicoplast targeted proteins during second life cycle without affecting the translation of other proteins. It remains to be seen further, how this differential regulation of translation is maintained in the parasites. This work shed some light to understand the mechanism behind the delayed death phenomena showed by some drugs like ciprofloxacin and tetracycline that affect apicoplast function. The parasites undergo normal division and segregation of organelles during the first life cycle. However, severe defects in apicoplast growth, morphology and cytokinesis take place in the second life cycle. Consistent with the above results, we find that the translocation of apicoplast targeted proteins like SSB is not affected during the first life cycle. It is possible that the accumulation of the drugs in the four membrane bound apicoplast is a slow process and the parasites can bypass the first life cycle since the drug may be limiting and it may not reach the critical concentration that may affect the apicoplast function immediately. There is also a constant translocation of fresh proteins from the cytoplasm to the apicoplast during this stage. However, during the second life cycle, the parasites may acquire the critical concentration of the drugs that may affect the apicoplast function, growth and morphology, which in turn affect the protein translocation in the apicoplast, leading to the accumulation of the unprocessed form of the proteins in the cytoplasm. This, in turn may be sensed by the cytoplasmic translational machinery involved in synthesis of apicoplast targeted proteins, to selectively reduce their synthesis. Latter, the complete inhibition of apicoplast function and growth may affect overall growth of the parasite leading to the parasitic death since apicoplast is the indispensable organelle of the parasite involved in various important pathways like fatty acid synthesis and haem biosynthesis. Therefore, dual mechanism consisting of translocation and translation defect of apicoplast targeted proteins during second life cycle may contribute to the delayed death phenomena shown by some apicoplast targeted drugs like ciprofloxacin and tetracyclin (Figure 7). Although ciprofloxacin is a specific inhibitor of gyrase, it remains to be seen further whether ciprofloxacin has a direct effect on the translational machinery during second life cycle following drug treatment.

To understand the fundamental process of apicoplast DNA replication completely, it is essential to identify all the key enzymes and study their role for the replication of DNA.

Figure 7. Model showing the effect of ciprofloxacin on parasites. In the absence of apicoplast targeted drug like ciprofloxacin, nuclear encoded apicoplast targeted (NEAT) genes are transcribed in the nucleus (N) and transported into the cytoplasm (C) where they are translated as unprocessed protein that are translocated to the apicoplast (A) and gets processed. During second life cycle following drug treatment, the apicoplast achieve critical concentration of the drug that affects apicoplast function and morphology that inhibits translocation of NEAT proteins into the apicoplast which in turn may down-regulate the translation of these proteins. Therefore, a dual mechanism of inhibition of translocation of NEAT proteins and modulation of translation level of these proteins may contribute to the delayed death effect exerted by such drugs like ciprofloxacin. The solid and broken circles in apicoplast indicate the apicoplast genome in the absence and presence of drug.
35 kb DNA circle. Our results provide conclusive evidence that apicoplast imports a bacterial homologue of nuclear encoded ssDNA binding protein that could be required in ssDNA intermediate stabilization during the process of DNA replication. The apicoplast localized PfSSB can be used as a tool to identify other components of the replication machinery as its homologue in bacteria has been shown to interact and modulate the activities of different members of replication proteins during the fork progression.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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REFERENCES


