Identification and bioinformatics characterization of translation initiation complex eIF4F components and poly(A)-binding protein from *Plasmodium falciparum*

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Protein synthesis in eukaryotes initiates with binding of the multisubunit translation initiation complex eIF4F. This complex contains eIF4E, eIF4A and eIF4G. eIF4E directly interacts with the cap structure, eIF4A is an RNA helicase and eIF4G acts as a scaffold for the complex. eIF4G contains the binding sites for both the subunits i.e., eIF4A and eIF4E and it also interacts with poly(A)-binding protein (PABP). In present study we have identified and characterized the main components of the eIF4F complex i.e., eIF4E, eIF4A and eIF4G and PABP from *Plasmodium falciparum*. Molecular modeling of PfeIF4E, PfeIF4G and PfPABP confirms that they contain all the characteristic conserved structural features. We have annotated some of the genes of *P. falciparum* and as a result these studies demonstrate that the components of translation initiation complex are highly conserved. Therefore these studies will contribute to understand the basic biology and components of translation complex in *P. falciparum*.

Generally eukaryotic mRNAs contain a 5’ cap (m(7)GpppN) where N is any nucleotide and a 3’ poly(A) tail (50–300 nucleotides), both of which are essential for efficient mRNA translation and are specifically recognized by the proteins. Eukaryotic translation initiation is a multistage process, which involves several sequential steps performed by a multisubunit protein complex termed as eukaryotic translation initiation factor 4F (eIF4F).1,2 eIF4F is composed of multiple translation factors, which assemble at the monomethylated cap present on the 5’-end of the mRNA to promote the recruitment of the ribosome. eIF4F mainly consists of three subunits, eIF4A (eukaryotic translation initiation factor 4A), an ATP-dependent RNA helicase and a member of the DEAD-box protein family (DEAD corresponds to Asp-Glu-Ala-Asp), eIF4E (eukaryotic translation initiation factor 4E) which binds to the 5’ cap structure of the mRNA and eIF4G (eukaryotic translation initiation factor 4G) that acts as a scaffold for the complex with binding sites for both eIF4A and eIF4E. eIF4G is also responsible for the circularization of the mRNA via interaction with poly(A) binding protein (PABP).3 The 5’ cap and the 3’ poly(A) tail of eukaryotic mRNAs cooperate to synergistically increase translation.

The functions of the components of eIF4F complex have been well established by studies in various systems. eIF4A is believed to be involved in the unwinding of the inhibitory secondary structures present in the 5’ untranslated region of the mRNA and its maintenance in a single stranded form to facilitate the binding and scanning of the ribosomes to the initiator AUG codon.4,5 eIF4E mediates cap identification during initiation and is absolutely necessary for cap-dependent translation.6 Most organisms contain a single gene, which encodes eIF4E but some organisms contain more than one eIF4E gene, which belong to different families.7 eIF4G is an adapter protein with a modular structure, which is required to mediate the ribosome recruitment via the cap structure and poly(A) tail.1 Mammals possess two functional isoforms of eIF4G eIF4GI and eIF4GII.2 The binding sites for numerous interaction partners of eIF4G have been determined by mutation and deletion analysis and it has been suggested that eIF4G provides multiple contact points for interaction with other components of the initiation complex.2 It has been shown that mammalian eIF4G contains two interaction domains for eIF4A, a C-terminal domain and a central domain and the interaction of eIF4A with the middle region of eIF4GI is necessary for translation, whereas the interaction of eIF4A with the C-terminal region plays a modulatory role.8,9 PABPs usually bind poly(A) using one or more globular domains containing RNA-recognition motifs (RRMs) and therefore help in stabilization of mRNAs. PABPs are involved in interaction of the poly(A) tail with the translation initiation complex and simultaneous binding of eIF4E and PABP to eIF4G is essential for efficient translation.2,3,10 The highly conserved PABP polypeptides are found only in eukaryotes; single-celled eukaryotes each contains 1–2 PABP genes, while humans have five and Arabidopsis has up to eight PABP genes.3,11 Several molecules of PABP cover the entire length of the poly(A) tail of the mRNA and this interaction occurs via RRMs of PABP.3,11

Malaria is a major parasitic infection caused by the protozoan parasites of the genus Plasmodium and *Plasmodium falciparum* causes
the most virulent form of malaria. In order to understand the basic biology and specially the translation process in this protozoan parasite we have bioinformatically characterized various components of the eIF4F complex from *P. falciparum*. In previous studies we have reported the detailed characterization of an eIF4A homologue designated as PfH45 from *P. falciparum*. In the present manuscript we present characterization of the other principal components of the eIF4F complex such as eIF4E, eIF4G and PABP from *P. falciparum*. Our studies reveal that *P. falciparum* contains only one gene each encoding eIF4E, eIF4G and PABP. The bioinformatics characterization of these proteins reveals that PfIF4E, PfIF4G and PfPABP all contain the characteristics features. PfPABP also contains the characteristic RRMs. A three-dimensional homology model of the middle domain of PfIF4G shows that there is sufficient conservation to human eIF4G to make the models virtually super-imposable. The studies reported here show that despite the insignificant homology of PfIF4G at the sequence level its structure is highly conserved. These studies will make an important contribution towards understanding the process of translation in the parasite.

**Sequence Analysis and Computer Based Prediction of Structure of eIF4E Homologue from *P. falciparum***

The search of the *P. falciparum* genome using eIF4E as query revealed that only one homologue of eIF4E is present in the genome. The 684 base-pair open reading frame of *P. falciparum* eIF4E (PfeIF4E) was analyzed. The deduced amino acid sequence of PfeIF4E revealed a protein consisting of 227 amino acids with a predicted molecular mass of approximately 27 kDa and it is a basic protein with a calculated isoelectric point of ~8.4. The sequence analysis confirmed the presence of characteristic motifs of this family including the conserved tryptophan residues. The 'PlasmoDB' (http://www.plasmodb.org) entry number for this gene is PFC0635c and it contains no introns. A multiple alignment of amino-acid sequence homology search using NCBI database revealed that PfeIF4E aligned contiguously and showed highest homology with its counterparts from *Plasmodium berghei* (~92%) and ~19–40% homology with eIF4E from other sources (Fig. 1A and B).

The 3 dimensional structures of the yeast eIF4E determined by NMR and mouse and human eIF4E proteins determined by X-ray crystallography have been reported. The murine protein has a cupped hand shape and contains eight-stranded anti-parallel β-sheets, backed by three α-helices on its convex side. A thorough examination of the alignment of PfeIF4E revealed that the cap-binding pocket is highly conserved. A comparison of the amino acids of human eIF4E and PfeIF4E is shown in Figure 2A. The interactions mediated by the amino acids required for specific binding to the 7-methyl-GDP are taken over by the amino acids Trp37, Asp102, Trp115, Glu116, Lys170, Arg180, His125 and Trp184 in PfE27,16 (Fig. 2A, marked by red asterisk). Similarly the alignment showed that the amino acids His37, Pro38, Val69, Trp73, Leu128 and Leu135 of mammalian eIF4E, which are responsible for binding to eIF4G, are replaced by the amino acids Leu31, Leu32, Val67, Trp71, Asn145 and Leu149 in PfeIF4E (Fig. 2A, marked by blue asterisk). For the location of the binding pocket on PfeIF4E, the complete protein sequence was submitted to the JIGsaw program (http://www.bmm.icnet.uk/servers/3d jig saw/). A highly similar and superimposable model based on the crystal structure of human eIF4E was obtained [Fig. 2B (i)]. In these models the amino acids responsible for the binding of eIF4G have been marked using the VMD software (www.ks.uiuc.edu) and are shown in yellow color [Fig. 2B (ii) and (iii)]. These results further show that the binding pocket is present on the similar planes in PfeIF4E and human eIF4E.

**Sequence Analysis and Computer Based Prediction of Structure of eIF4G Homologue from *P. falciparum***

On simple text-based data mining of PlasmoDB with the text term ‘MIF4G’ five genes are reported. These are PF11_0086 (hypothetical protein of 3334 amino acid in PlasmoDB), PF11265w (hypothetical protein of 1754 amino acid in PlasmoDB), PFL1855w (putative cell cycle control protein of 967 amino acid in PlasmoDB), PF14_0113 (hypothetical protein of 943 amino acid in PlasmoDB) and MAL13P1.63 (asparagine-rich protein of 1192 amino acid in PlasmoDB). The detailed analysis of all of these proteins revealed that these contain MIF4G domain but all the others except MAL13P1.63 did not reveal any detectable homology to known proteins from other organisms. The detailed sequence analysis of MAL13P1.63 (~139 kDa protein) confirmed the presence of characteristic motifs of eIF4G. A multiple alignment of amino-acid sequence homology search using NCBI database revealed that PfeIF4G aligned contiguously and showed highest homology with its counterparts from *Plasmodium yoelii* (~49%) and ~6–13% homology with eIF4G from other sources (Fig. 3A and B).

Since PfeIF4G contained insignificant homology at amino acid level therefore to check the homology at structural level, the complete protein sequence of PfeIF4G was submitted to the 3D-JIGSAW program and a three-dimensional model was obtained, which was based on the X-ray structure of the template human eIF4G. On careful analysis it was observed that this model was built by using only the phylogenetically conserved region i.e., amino acid 888 to amino acid 1125 in PfeIF4G (Fig. 4B (i)). It is interesting to note that this domain is composed of ten α helices arranged as five anti-parallel α helical pairs or HEAT repeats (named for Huntington, Elongation factor 3, A subunit of protein phosphatase 2A [PP2A], and Target of rapamycin). Similarly using the complete protein sequence of human eIF4G (AF104913) a three dimensional model was obtained [Fig. 4B (ii)]. This model spanned from amino acid 712 to 944 of human eIF4G. The alignment of amino acid 888 to 1125 of PfeIF4G and amino acid 712 to 944 of human eIF4G showed only ~22% similarity and 27% identity (Fig. 4A). But it is interesting to note that the two structures of these regions are highly similar and largely super-imposable [Fig. 4B (iii)].

**Sequence Analysis and Computer Based Prediction of Structure of PABP Homologue from *P. falciparum***

On simple text-based data mining of PlasmoDB with the text term ‘poly A binding protein’ two genes are reported. These are MAL13P1.303 (putative polyadenylate binding protein of 414 amino acids) and PFL1170w (putative polyadenylate binding protein of 875 amino acids). MAL13P1.303 is small in size to be a bonafide PABP. The detailed sequence analysis of PFL1170w revealed a protein with a predicted molecular mass of ~97 kDa and it is a basic protein with a calculated isoelectric point of ~8.9. A multiple alignment of amino-acid sequence homology search using NCBI database revealed that PFL1170w (PfPABP) aligned contiguously and showed highest
Figure 1. (A) Comparison of amino acid sequences of *P. falciparum* eIF-4E protein (PlasmoDB No. PFC0635c; GenBank accession No. EF043517) with other eIF-4E from *Plasmodium berghei* (XP_674865), *Theileria annulata* (CAI74251), *Cryptosporidium parvum* (CAD98650), *Pisum sativum* (ABG35118), *Saccharomyces cerevisiae* (NP_014502) and *Homo sapiens* (P06730). The accession numbers of the aligned sequences are written in brackets. (B) Phylogenetic guide tree using Clustal W program based on the multiple alignment. A value of 0.1 corresponds to a difference of 10% between two sequences.
homology with its counterparts from *Plasmodium berghei* (~71%) and ~29–37% homology with PABP from other sources (Fig. 5). It was observed that similar to other PABPs, PfPABP also contains four RRM3s, which span from amino acid 16–94; 104–176; 193–261 and 450–527 respectively (Fig. 6A). It contains a few regions, which are Asn rich (amino acid 295–396; 617–799), Thr rich (amino acid 317–362) and Gln rich (amino acid 557–761) (Fig. 6A). The presence of homorepeats is a characteristic feature of proteins from *P. falciparum* and the most common is asparagines followed by lysines and glutamine.25,26 These inserted residues encode non-globular domains of unidentified function, which are extruded from the core of the protein and probably form surface exposed structure that consequently in most of the proteins have no effect on the functional folding of the protein.25,26

The complete protein sequence of PfPABP was submitted to the 3D-JIGSAW program and a three-dimensional model was obtained, which was based on the X-ray structure of the template human PABP. These two structures were not similar and not super-imposable (data not shown). The amino acid alignment of full-length human and PfPABP revealed a similarity of ~14% and an identity of ~33% and it further showed that PfPABP contains an insertion of ~150 amino acids in between RRM3 and 4 (Fig. 6A). It is interesting to note that only the modeled structure of PfPABP from amino acids 1–180 (RRM1 and 2) superimposes with that of human PABP [Fig. 6B (i)] whereas the structures of PfPABP from amino acids 1–264 (RRM1-3) and from amino acids 1–531 (RRM1-4) are not superimposable with human PABP [Fig. 6B (ii) and (iii) respectively].

Our unpublished observations indicate that the middle and C-terminal domain of PfIF4G is structurally conserved and is able to interact with PfH45, PfIF4E and PfPABP but there was no detectable interaction between the PfIF4G-N fragment and PfIF4E or PfH45 or PfPABP. These interactions are shown schematically in Figure 7A and B shows the overall initiation complex of *P. falciparum*.

Protein synthesis is a complex process in eukaryotes and it requires numerous different macromolecules. The critical step of translation initiation is a coordinated action of a number of translation initiation factors. Most important of these is the heterotrimeric elf4F complex, which is composed of elf4A, elf4E and the very large protein elf4G, which mediates the interactions between elf4F and other translation factors. Previously we have reported the char-
eIF4F components and PABP from *Plasmodium falciparum*

Previous studies have shown that the HEAT repeat proteins participate in a wide variety of cellular processes, which are dependent on accumulating large multiprotein complexes.23,24 The overall similarity between human eIF4G and PfIF4G is not very high (only ~12%) because it is well established that the HEAT repeat proteins do not share any absolutely conserved amino acids.23 It has been reported that although *Leishmania major* contains five eIF4G homologues but most of them show similarity to the human sequence only at the level of central HEAT domain.28 The comparison of human and *P. falciparum* eIF4G suggests that specifically the middle region of PfIF4G might be involved in the interaction with PfH45 as reported for human eIF4G and eIF4A.24 Due to the presence of HEAT domain, the conserved middle domain of PfIF4G most likely serves as a central assembly platform for its translation initiation machinery. It has been reported previously that mammalian eIF4A characterization of an eIF4A homologue (PfH45) from *P. falciparum* and have shown that PfH45 is localized in the cytoplasm and is essential for parasite survival.13 In the present study we have isolated and characterized the other components i.e., eIF4E and eIF4G of eIF4F complex and PABP from *P. falciparum* using the bioinformatics approach. Previously it has been reported that PfIF4E binds to mRNAs in *P. falciparum*.27

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Figure 3A. For figure legend, see page 253.
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has been reported previously that the amino acid sequences of the PABP-binding site in eIF4G are not conserved between yeast and human. The interaction between PABP and eIF4G is conserved in many species and this interaction can be imparted to RRM 1 and 2 of PABP. Previously it has been reported that the interaction of eIF4G with PABP requires RRM1 and 2 in animal and yeast systems but in wheat PABP RRM1 was sufficient to interact with eIF4G. The structural modeling of PfPABP suggests that the RRM1 and 2 are structurally conserved and PfPABP is also able to interact with the middle and C-terminal domain of Pf eIF4G.

Overall in this study we have reported the bioinformatics characterization of important components of translation initiation complex.

Figure 3A (continued). For figure legend, see page 253.

binds to the central region of eIF4G via its HEAT domain and it also binds to its C-terminal domain. Similarly in yeast it has been shown that eIF4G and eIF4A physically associate and this association is required for translation and is essential for yeast cell viability. The binding site for eIF4E in human eIF4G has been reported previously and the alignment of the human and Pf eIF4G shows that this site in Pf eIF4G is between amino acid 464 to 473 and these amino acids are present in the middle domain of Pf eIF4G.

It is interesting to note that *P. falciparum* contains only one PABP, which is localized in the cytoplasm and as reported earlier evolutionarily PIPABP is closer to PABP from *L. major*. Multiple PABPs, both nuclear and cytoplasmic have been reported in human. It has been reported previously that the amino acid sequences of the PABP-binding site in eIF4G are not conserved between yeast and human. The interaction between PABP and eIF4G is conserved in many species and this interaction can be imparted to RRM 1 and 2 of PABP. Previously it has been reported that the interaction of eIF4G with PABP requires RRM1 and 2 in animal and yeast systems but in wheat PABP RRM1 was sufficient to interact with eIF4G. The structural modeling of PfPABP suggests that the RRM1 and 2 are structurally conserved and PfPABP is also able to interact with the middle and C-terminal domain of Pf eIF4G.

Overall in this study we have reported the bioinformatics characterization of important components of translation initiation complex.
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**Figure 3A (continued).** For figure legend, see page 253.

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**Table 1:**

| Species       | Accession Number | Protein | Description |
|---------------|------------------|---------|-------------|
| *P. falciparum* | 561              | VNNNFTS | VNNNFTS     |
| *P. yoelii*    | 463              | TGGMADT | TGGMADT     |
| *S. pombe*     | 647              | BAAAAAAAAAA | BAAAAAAAAAA |
| *O. sativa*    | 398              | LIGELKX | LIGELKX     |
| *H. sapiens*   | 713              | AOSKTVQ | AOSKTVQ     |

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**Figure 3A (continued).** For figure legend, see page 253.
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Figure 3. (A) Comparison of amino acid sequences of *P. falciparum* eIF-4G protein (PlasmoDB No. MAL13P1.63) with other eIF4G from *Plasmodium yoelii* (XP_727991), *Schizosaccharomyces pombe* (Q10475), *Oryza sativa* (AAO72569) and *Homo sapiens* (AF104913). The accession numbers of the aligned sequences are written in brackets. (B) Phylogenetic guide tree using Clustal W program based on the multiple alignment. A value of 0.1 corresponds to a difference of 10% between two sequences.
Figure 4. (A) Comparison of amino acid sequence (from amino acid 888–1125) of *P. falciparum* eIF4G protein (PlasmoDB no. MAL13P1.63) with eIF4G (from amino acid 712–944) of *H. sapiens* (GenBank accession number AF104913). (B) Three-dimensional models for (i) human eIF4G and (ii) *P. falciparum* eIF4G were created as described in text. The structures have been displayed using the same program as described in legend to Figure 1B. This model has been built by using only the phylogenetically conserved middle domain i.e., amino acid 888 to amino acid 1125 in Pf eIF4G. A super-imposed image is shown in (iii).
Figure 5A. For figure legend see page 258.
Figure 5A (continued). For figure legend see page 258.
Figure 5A (continued). For figure legend see page 258.
Figure 5. (A) Comparison of amino acid sequences of *P. falciparum* PABP protein (PlasmoDB No. PFL1170w; GenBank accession No. EF116593) with other PABP from *Plasmodium berghei* (XP_677383), *Theileria annulata* (CAI73150), *Cryptosporidium parvum* (CAD98589), *Schizosaccharomyces pombe* (AAA35320), *Oryza sativa* (XP_481529) and *Homo sapiens* (CAI12298). The accession numbers of the aligned sequences are written in brackets. (B) Phylogenetic guide tree using Clustal W program based on the multiple alignment. A value of 0.1 corresponds to a difference of 10% between two sequences.
Figure 6. (A) Comparison of amino acid sequence of *P. falciparum* (PlasmoDB no. PFL1170w, GenBank accession number EF116593) and human PABP (GenBank accession number CAI 12298). The RNA recognition motifs (RRM) 1–2, 3 and 4 have been highlighted with green, blue and orange color respectively. (b) Three-dimensional models for human PABP and various fragments of PfPABP were created as described in text. The structures have been displayed using the same program as described in legend to Figure 1B. (B) (i)–(iii) Show the modeled structure of human (in yellow) and PfPABP (in green from amino acid 1–180), PfPABP (in blue from amino acid 1–264) and PfPABP (in orange from amino acid 1–531) respectively.
Figure 7. (A) Schematic diagram showing the interaction of various components (eIF4E, eIF4G and PABP) with the N-terminal, Middle and C-terminal domains of eIF4G. (B) Schematic diagram of translation initiation complex in *P. falciparum*. The name and PlasmoDB number of each component is written.