Identification of a Novel and Unique Transcription Factor in the Intraerythrocytic Stage of *Plasmodium falciparum*

Kanako Komaki-Yasuda1*, Mitsuru Okuwaki2, Kyosuke Nagata2, Shin-ichiro Kawazu3, Shigeyuki Kano1

1 Research Institute, National Center for Global Health and Medicine, Shinjuku-ku, Tokyo, Japan, 2 Graduate School of Comprehensive Human Sciences and Institute of Basic Medical Sciences, University of Tsukuba, Ibaraki, Japan, 3 National Research Center for Protozoan Diseases, Obihiro University of Agriculture and Veterinary Medicine, Obihiro, Japan

Abstract

The mechanisms of stage-specific gene regulation in the malaria parasite *Plasmodium falciparum* are largely unclear, with only a small number of specific regulatory transcription factors (AP2 family) having been identified. In particular, the transcription factors that function in the intraerythrocytic stage remain to be elucidated. Previously, as a model case for stage-specific transcription in the *P. falciparum* intraerythrocytic stage, we analyzed the transcriptional regulation of *pf1-cys-prx*, a trophozoite/schizont-specific gene, and suggested that some nuclear factors bind specifically to the cis-element of *pf1-cys-prx* and enhance transcription. In the present study, we purified nuclear factors from parasite nuclear extract by 5 steps of chromatography, and identified a factor termed PREBP. PREBP is not included in the AP2 family, and is a novel protein with four K-homology (KH) domains. The KH domain is known to be found in RNA-binding or single-stranded DNA-binding proteins. PREBP is well conserved in *Plasmodium* species and partially conserved in phylum Apicomplexa. To evaluate the effects of PREBP overexpression, we used a transient overexpression and luciferase assay combined approach. Overexpression of PREBP markedly enhanced luciferase expression under the control of the *pf1-cys-prx* cis-element. These results provide the first evidence of a novel transcription factor that activates the gene expression in the malaria parasite intraerythrocytic stage. These findings enhance our understanding of the evolution of specific transcription machinery in *Plasmodium* and other eukaryotes.

Introduction

*Plasmodium falciparum* is a significant life-threatening parasitic pathogen that causes falciparum malaria in humans [1]. In spite of years of intensive research, no effective vaccine is presently available, and existing antimalarial drugs are becoming less effective because of the rapid emergence of drug-resistant parasites [2]. To develop new strategies for combating *P. falciparum*, a better understanding of the basic biology of this microorganism is required. In particular, the mechanisms of stage-specific gene regulation remain mostly unclear and only a small number of regulatory transcription factors (TFs) have been predicted in the *Plasmodium* genome [3,4].

A bioinformatic analysis of the complete *Plasmodium falciparum* genome revealed that the hypothetical *Apetala2* (AP2) family with plant AP-2-like DNA-binding domains [5] is the gene family that encodes TF candidates in the *Plasmodium* genome [6,7]. Twenty-six AP2-related genes were predicted in the *P. falciparum* genome. In a rodent malaria model, it was revealed that some of the AP2 family proteins indeed regulated specific gene expression in the ookinete stage [8], the sporozoite stage [9] and the liver stage [10]. However, the functions of TF in the intraerythrocytic cell cycle remain to be elucidated at the present time.

The intraerythrocytic cell cycle is important for research because it causes the symptoms of malaria [11]. Studies with microarray techniques revealed that at least 60% of the *P. falciparum* genome is transcriptionally active during intraerythrocytic development [12]. The timing of gene expression appeared to be defined quite rigidly and was related to the intraerythrocytic cell cycle of the parasite [12,13]. As a model of the general transcriptional regulatory mechanisms in *P. falciparum*, we are interested in the mechanism that regulates the transcription of the Pf1-Cys-Prx (PlasmoDB ID, PF3D7_0809200) gene of *P. falciparum* as a model for understanding stage-specific gene expression in the intraerythrocytic stages. Pf1-Cys-Prx is an antioxidant protein and a member of peroxiredoxin family [14]. The expression of *pf1-cys-prx* has been shown to be almost non-existent during the ring stage and markedly elevated during the trophozoite/schizont stage [15]. In a previous study, the authors precisely analyzed the promoter activity of the 5′ region of *pf1-cys-prx*, verified the promoter activities of the 5′ regions of *pf1-cys-prx*, and delineated the 102-bp cis-acting enhancer region in this gene [16]. An electrophoresis mobility shift assay (EMSA) and a DNase I footprinting assay both revealed that the double-stranded 102-bp enhancer sequence of *pf1-cys-prx* was the target of trophozoite/schizont stage-specific DNA-binding proteins in the parasite nucleus. We also observed that the levels of histone acetylation in the 5′ region of *pf1-cys-prx*...
were elevated according to the elevation of transcription. In the pf1-cys-prx, a trophozoite/schizont stage-specific recruitment of PfGCS3 histone acetyltransferase to the putative enhancer region was also observed. These results suggest that *P. falciparum* possesses a sophisticated system of transcriptional regulation during the intraerythrocytic stages that is managed by coordinated interactions of unique cis-elements, trans-acting factors and chromatin modifications.

In the present study, the 102-bp cis-enhancer region was defined as Pex Regulatory Element (PRE), and the hypothetical nuclear factor, which binds specifically to the PRE sequence, was defined as PRE Binding Protein (PREBP). PREBP might be a key molecule for the specific transcriptional regulation of pf1-cys-prx in the intraerythrocytic stages. We attempted to purify and identify PREBP directly from a huge amount of parasite nuclear extract and finally verified a single protein as authentic PREBP, a novel and unique protein in the *Plasmodium* species, with four K-homology domains that has been found in a number of single-stranded DNA or RNA binding proteins [17,18]. PREBP is considered to be a novel transcriptional factor, which functions in the intraerythrocytic stages and which is uniquely evolved in the *Plasmodium* species.

**Results**

**Purification of Cis-Element Binding Protein**

In order to identify PREBP, we set up a huge-scale protein purification scheme using 5 x 10^11 parasite cells obtained from the culture of 33 liters. PREBP was purified from parasite nuclear extract as summarized in Figure 1A and Table S1. In the final DNA affinity chromatography step, the pooled fraction obtained immediately before Mono S chromatography was incubated with magnetic beads, which tethered the 102-bp PRE sequence (Figure 1B). Each fraction obtained from each DNA affinity chromatography step was checked with EMSA and as a result the first eluents of both the first and second purification step showed specific cis-element binding activity (Figure 1C).

The purity of the fractions was assessed by SDS-PAGE followed by silver staining and numerous bands appeared (Figure 1D). Then, to gain insight into the molecular weight of the true binding protein, we performed small-scale protein purification using DNA-coupled magnetic beads. The roughly purified fraction was separated by SDS-PAGE and a series of gel pieces were excised in order of molecular weight (Figure 1E, right panel). Proteins were then eluted from each gel piece and renatured. Each renatured fraction was subjected to EMSA. As a result, DNA-binding activity was detected in the second fraction, corresponding to a molecular weight range of between 55-kDa and 70-kDa (Figure 1E, left panel). In the final fractions of the huge-scale purification, 3 bands were confirmed within the expected molecular weight range (Figure 1F).

**Identification of PREBP by Mass Spectrometry**

The final fraction obtained from DNA-affinity chromatography was subjected to SDS-PAGE and visualized by silver staining specialized for mass spectrometry (Figure S1). The 3 bands were excised from the gel slice, and subjected to mass spectrometry analysis. Database searches with the fragmentation spectra obtained by LC-MS/MS analysis revealed that these bands contained several *P. falciparum* proteins. The 16 proteins that showed the most frequent spectra are indicated in Table S2. To detect true PREBP among these candidate proteins, 16 candidate proteins were first synthesized using a cell-free translation system, and their PRE binding activity was checked by EMSA. However, no recombinant protein showed the specific binding activity (Figure S2, Method S1). Next step, 4 transgenic parasite lines, which expressed protein numbers 1 (PF3D7_0217500), 2 (PF3D7_0617200), 3 (PF3D7_0617200) and 4 (PF3D7_1011800) excessively, were generated (Figure 2A). In the LC/MS/MS analysis, signals corresponding to protein No. 1, 2, 3 and 4 were detected with high frequency and were thus thought to be the 4 most preferable candidates. The expressed recombinant candidate proteins were then roughly purified from the transgenic parasites (Figures 2B and 2C) and their activity was checked by EMSA. Only protein No. 4, which corresponded to PF3D7_1011800, showed specific PRE binding activity and formed a shift-band with similar mobility to that formed with the parasite nuclear extract (Figure 2D). To evaluate the sequence specificity of the interaction, cross-competition assays were performed. The 102-bp sequences of the PRE or other AT-rich DNA fragments derived from the 5’ region of the pf1-cys-prx were added to the EMSA reaction mixtures as competitors. A 90-fold molar excess of unlabeled probe prevented the formation of DNA-protein complex (Figure 3A, lane 4). It is noteworthy that the 270-fold molar excess of AT-rich fragments obtained from the 5’ regions of pf1-cys-prx showed less inhibition of DNA-protein complex formation. Thus, this fragment could not completely prevent a band shift. These results indicated that the interaction between the PRE and the recombinant PF3D_1011800 was sequence-specific. Furthermore, the peptide-antibody against the PF3D_1011800 candidate protein was raised (anti-PREBP pep.) and added into the reaction mixtures of EMSA using the parasite nuclear extract and the 102-bp PRE sequence as a labeled probe, which resulted in a band-shift corresponding to the complex of the probe and authentic PREBP in the nuclear extract. The band-shift disappeared and a super-shift band appeared when the antibody was added into the reaction mix, indicating that the antibody against the PF3D_1011800 protein bound specifically to the PREBP-cis-element complex (Figure 3B). Thus, PF3D7_1011800 was verified as an authentic PREBP.

**Molecular Features of the Identified PREBP, PF3D7_1011800**

The amino acid sequence and characteristic features of the PF3D7_1011800 are summarized in Figure S3 and Figure 4. In PlasmoDB, PF3D7_1011800 is registered as “QF122 antigen” [19]. The deduced molecular mass of PREBP was 132 kDa. The size of PREBP estimated by SDS-PAGE was 60 kDa (Figure 1F). The peptides detected by LC-MS/MS covered the central 60 kDa region of PF3D7_1011800. No peptides corresponding to the N or C terminal regions were detected. Thus, the band of ~60 kDa obtained from the purification steps (Figure 1F) was deduced to be a shortened form due to the processing of protein in the parasite cells, or artificially cut form that can be produced during the purification steps. The characteristic feature of PREBP is that it contains four putative K-homology (KH) domains (Figure 4). The orthologs of PREBP are highly conserved in *Plasmodium* species and those with lower identity have been found in many other Apicomplexan parasites, *Cryptosporidium*, *Theileria* and *Babesia* (Table 1). No ortholog was found in the *Toxoplasma* genome. The PREBP mRNA expression profile during intraerythrocytic development was analyzed by real-time RT-PCR using cDNA obtained from tightly synchronized parasites. PREBP mRNA expression was almost constant during the intraerythrocytic development, although only slight changes of expression were observed (Figure S4, Method S2).
Figure 1. Purification of PREBP. (A) Scheme for purification of PREBP from parasite nuclear extract by 5 chromatography steps. (B) Detailed scheme of the final DNA-affinity chromatography step. (C) EMSA of each elution and washing fraction obtained from DNA-affinity purification. The number of the fraction corresponds to the purification scheme shown in Figure 1B. 0.1 μl of each fraction and the 32P-labeled PRE probe were used for the EMSA. Lanes N and P indicate the negative control (probe only) and positive control (assay with the 0.6 μg of parasite nuclear extract). Positions of the free probe and the shifted band are indicated on the left. (D) SDS-PAGE and silver staining of the PRE affinity-purified fractions. 0.5 μl of each PRE-purified fraction was separated on SDS-PAGE and proteins were visualized by silver staining. Molecular mass markers are indicated in kDa on the left. (E) Estimation of the molecular size of PREBP. A partially purified PREBP fraction was applied to an SDS-PAGE. Gel pieces were cut from the gel according to the 8 grades of molecular weight. The range of molecular weight of each gel piece is shown in the right panel. The proteins were eluted from each gel piece and renatured. The PRE-binding activity of the each renatured fraction was checked by EMSA (left panel). (F) Probable candidate bands observed in the final fractions. The 3 candidate bands were indicated with arrows.

doi:10.1371/journal.pone.0074701.g001
Endogenous PREBP in the Trophozoite/Schizont Nucleus was Expressed as Proteins of Approximately 132 kDa

To detect the stage-specific expression and cellular localization of the endogenous PREBP protein, IP-Western blotting experiments with nuclear extract and cytoplasmic fraction prepared from both ring and trophozoite/schizont parasites were performed (Figure 5). For this purpose, antibody was raised against recombinant partial PREBP protein, which corresponded to amino acid numbers 699–849 (anti-PREBP rec.). As a result, 2 bands of ~130 kDa were detected in the anti-PREBP antibody-precipitant from the nuclear extract of trophozoites/schizont parasites. From the nuclear extract and cytoplasmic fraction of ring stage parasites, only faint bands were detected in the anti-PREBP antibody-precipitant. From the cytoplasmic fraction of trophozoite/schizont parasites, only a sub-band of higher mobility was detected. Thus, most endogenous PREBP protein was localized at the nucleus of trophozoite/schizont parasites. No other band was detected at ~60 kDa in any sample. Thus, it is suggested that endogenous PREBP exits in the parasite nucleus as ~132 kDa proteins, which correspond to the full length deduced from the sequence of ORF registered in PlasmoDB. The ~60 kDa band obtained from purification steps (Figure 1) was deduced to be an artificially cut and shortened form through the purification process.

Figure 2. Preparation of the recombinant candidate proteins using gene overexpression system in P. falciparum. (A) Detailed descriptions for each candidate protein. (B) Schematic diagram of plasmid constructs used for the gene overexpression in P. falciparum. Genes for candidate protein Nos. 1, 2, 3 and 4 were inserted into the expression site of the pHCl vector. For protein purification, the FLAG-tag and His-tag were added to the N and C termini, respectively. The restriction endonuclease recognition sequences shown is X, XhoI. (C) Immunopurification of the recombinant proteins from transgenic parasites. Expressed recombinant proteins were immunopurified with the anti-FLAG antibody and subjected to Western blotting with anti-FLAG antibody. The bands corresponding to the recombinant proteins are indicated with white arrowheads. Sizes are indicated in kDa on the left. (D) EMSA with partially purified recombinant proteins. The same labeled probe was used in Figure 1 in each assay. Lane N is probe only. EMSA was performed with 0.1 μg of each fraction of immunopurified recombinant protein. Lane P is the positive control for the assay with 0.6 μg of nuclear extract derived from the parasite synchronized at the trophozoite/schizont stage. Positions of the free probe and the shifted band are indicated on the left.

doi:10.1371/journal.pone.0074701.g002
PF3D7_1011800 protein showed specific binding activity to the PRE sequence. (A) Cross-competition assays. EMSA was performed with the same labeled probe used in Figure 1. Lane N is probe only. Lane P is the assay with 0.6 μg of nuclear extract from the parasites at tophozoite/schizont stages. Lanes 1–10 represent EMSA with 0.1 μg of immunopurified PF3D7_1011800 fraction (see Figure 2) in the presence or absence of cross-competitors. Lanes 1–5 represent cross-competition with increasing amounts of non-labeled probe. Lanes 6–10 represent another irrelevant 62-bp sequence [16]. The amount of competitor DNA in each reaction mix was indicated at the top of the panel. (B) Super-shift assays. EMSA was performed with the same 32P-labeled probe used in Figure 1. Lane N is probe only. Lane P and Lanes 1–10 represent EMSA with 0.6 μg of nuclear extract from the parasites at tophozoite/schizont stages in the presence or absence of antibodies. The amount of antibody added in each reaction mix is indicated at the top of the panel. Positions of the free probe, shifted band and super-shifted band are indicated on the left.

doi:10.1371/journal.pone.0074701.g003

| Organ                | Gene ID   | Protein length | Similarity (%) | Identity (%) |
|----------------------|-----------|---------------|----------------|--------------|
| P. yoelii            | PY03523   | 979           | 91.7           | 60.7         |
| P. berghei           | PBANKA_121020 | 970           | 90.7           | 61.0         |
| P. chabaudi          | PCHAS_121090 | 975           | 92.3           | 62.3         |
| P. vivax             | PVX_094810 | 985           | 90.5           | 54.6         |
| P. knowlesi          | PKH_081130 | 993           | 88.9           | 54.7         |
| P. cynomoligi        | PCYB_082160 | 964           | 88.5           | 52.8         |
| Cryptosporidium parvum | EAK90228.1  | 818           | 68.8           | 16.6         |
| Theileria perevani   | EAN34110.1 | 864           | 65.8           | 16.2         |
| Babesia bovis        | EDD007257.1 | 850           | 70.6           | 17.1         |

*Gene ID is the PlasmoDB ID for *Plasmodium* genes and Genbank ID for other Apicomplexan genes.

Protein length is number of amino acids.

Similarity and identity are calculated by comparison with the PREBP of *P. falciparum*.

doi:10.1371/journal.pone.0074701.t001
steps. The expression level of pf1-cys-prx in the trophozoite/schizont stages of the PREBP central ~60 kDa overexpressor was quantified by real-time RT-PCR and compared with that of the parental line. No significant difference was observed between the transgenic and parental parasites (Figure S5, Method S2).

Furthermore, to investigate the effect of overexpression of full-length PREBP, a plasmid construct for the excessive expression of full-length PREBP was transfected into the parasite. To obtain stable overexpressor, transgenic parasites with episomal plasmids were selected by pyrimethamine pressure. However, in spite of repetitive trials, no living transgenic parasite was obtained after drug selection (data not shown).

Overexpression of PREBP Enhanced the Expression of Reporter Gene Depending on the PRE Sequence in a Transient Luciferase Assay

To evaluate the effects of overexpression of full-length PREBP, we combined transient overexpression and a luciferase assay. The plasmid constructs used for the assays are indicated in Figure 6A. Plasmid p1–10R was the basal construct, which contains a firefly luciferase gene expression cassette controlled under the 0.8-kbp 5' region of the pf1-cys-prx gene. In p1–10R-PREBP-full and p1-10R-PREBP-short, the expression cassette for full-length PREBP and partial, ~60 kDa PREBP, respectively, were also inserted into the basal p1–10R under the control of 5' of calmodulin. Luciferase expression in the p1–10R-PREBP-full transfected cell would be expected to be enhanced if the overexpressed full-length PREBP affected the 5' promoter sequence of pf1-cys-prx gene on the same plasmid. Although the transfection efficiency of the electroporation system for P. falciparum is very low [20], enhanced expression of luciferase could be detected by the luciferase assay system because of its high sensitivity. As negative controls, we prepared p1–10R-Mock, which contained an empty overexpression cassette and p1-10R-No.2, which contained a candidate protein No. 2 (PF3D7_0617200: hypothetical Bfr1p homologue) overexpression cassette. These plasmid constructs were introduced into parasites synchronized in the ring stage by electroporation. After 20–24 hours, luciferase activity was determined at the trophozoite/schizont stages, before the completion of the cell cycle after transformation. Parasites transfected with p1–10R-PREBP-short or p1–10R-PREBP-full showed clear and drastic elevation of the reporter gene in comparison to parasites transfected with the basal plasmid, p1–10R (P<0.01). In contrast, no detectable elevation of reporter activity was observed in parasites transfected with p1–10R-Mock or p1–10R-No.2 (Figure 6B). These results suggest that in the parasites transfected with p1–10R-PREBP-full or p1–10R-PREBP-short, the overexpressed PREBP proteins affected the 5' sequence of pf1-cys-prx and enhanced the expression of the reporter gene (Figure S6: model scheme).

Furthermore, to investigate the dependency of the 102-bp PRE sequence on the enhancing activity of PREBP, a mutant plasmid of p1–10R-PREBP-full, which lacked the 102-bp PRE sequence of the 5' region of pf1-cys-prx gene, was constructed (p1–10R-Dcis-PREBP-full) and transfected into the parasite. As a result, reporter activity of the parasites transfected with the mutant plasmid, p1–10R-Dcis-PREBP-full, decreased to less than 30% of the reporter
activity of the transfectant of the original plasmid, 1–10R-PREBP-full ($P<0.01$, Figure 6B).

**Discussion**

Here, we report for the first time the purification and identification of a specific cis-element binding protein from the
The identified PREBP (PF3D7_1011800) binds the 102-bp PRE sequences with high affinity. In addition, we show that the overexpression of PREBP enhances the expression of the reporter gene under the control of the 5’ sequence of pfl-cys-prx sequence using the transient luciferase assay system and that this enhancing activity is dependent on the 102-bp PRE sequence. This result strongly suggests that the PREBP is a novel transcription factor in *Plasmodium*.

Despite exhaustive research on transcriptomes and proteomes based on the genome database of *Plasmodium* [12,13,21,22], little has been known about the regulation mechanisms of gene expression in the parasite [3,4]. The paucity of the knowledge of the basic biology might reflect the phylogenetic distance of the parasites from other eukaryotes. The phylum Apicomplexa, which is composed of a number of protozoan parasites including *Plasmodium*, belongs to the Alveolate phylogenetic group, the members of which are vastly distant from members of both the Opisthokonta (including most of model eukaryotes from yeasts to humans) and Planta branches [23].

The identified PREBP was registered as “QF122 antigen” in the PlasmoDB database because the partial sequence of this protein was determined in a previous report [19]. The screening of the parasite cDNA expression library with a monoclonal antibody bound to the merozoite surface and Western blotting showed that this monoclonal antibody recognized the 51 kDa protein in the parasite cell lysate. However, no follow-up was reported for this “antigen”. The detected PREBP was not a member of AP2 family, which is the only known specific transcription factor in *Plasmodium* [6,7,8,9,10]. Rather, it was a novel protein with four K-homology (KH) domains. The KH domain was first identified in human heterogeneous nuclear ribonucleoprotein (hnRNP) K [24]. The KH motif consists of approximately 70 amino acids, and is found in a diverse variety of proteins in archaea, bacteria and eukaryota [17,24]. It is known that the KH domain is contained in a number of RNA or single-strand DNA binding proteins that perform wide range of cellular functions [17,18]. Previously, it had been reported that a human protein with a KH domain could function as a transcription factor to activate the expression of the human c-myc gene [25]. This transcription factor (FBP) activates gene expression by binding to the single-stranded cis-element (FUSE: Far UpStream Element) [25,26]. In our previous study, we showed that the authentic PREBP in the nuclear extract could interact with the double-stranded cis-element sequence by DNaseI footprinting assay [16]. Thus, this study is the first report that shows that a protein with a KH domain functions as a transcription factor, which binds to specific double-stranded cis-element sequence. In future studies, it will be informative to elucidate whether PREBP binds to single-stranded-DNA and RNA in addition to the double-stranded PRE sequence. With the exception of *Plasmodium* species and some Apicomplexan species, no PREBP orthologs have been found in any other organisms. These findings indicate that this factor evolved uniquely in the *Plasmodium* species.

The mRNA expression of PREBP was not trophozoite/schizont stage-specific during the intraerythrocytic stages. However, the expression of PREBP protein was increased at the trophozoite/ schizont stages, suggesting that the expression of PREBP protein is regulated by post-transcriptional mechanisms. In addition, the immunoprecipitation-Western blotting analyses represented two bands of approximately 130 kDa as the endogenous PREBP and 60 kDa PREBP also enhanced the expression of the reporter gene under the control of the 5’ sequence of pfl-cys-prx 5’, and the expression of many other genes in the parasite cell and have a lethal effect on the parasite physiology. We tested the effects of overexpression of full-length PREBP using the transient luciferase assay system and the results clearly showed that overexpression of full-length PREBP enhances the expression of the reporter gene under the control of the 5’ sequence of pfl-cys-prx. Unexpectedly, partial ~60 kDa PREBP also enhanced the expression of the reporter gene under the promoter of pfl-cys-prx, despite the lack of detection of any significant affection in the transgenic parasites that stably overexpressed partial ~60 kDa PREBP. In the transient reporter assay system, the introduced plasmid DNA is considered to not be a formed chromatin structure. Thus, our results might suggest that full-length PREBP controls the expression of the endogenous gene in the chromatin structure. Partial ~60 kDa PREBP might bind the naked DNA sequence of PRE, but may not be able to access the cis-element of the endogenous gene because of its chromatin structure. The N and C terminal regions of PREBP protein might associate with other chromatin remodeling factors such as histone acetyl transferases and reconstitute the cis-element as an accessible form.

Our previous study showed that the level of histone acetylation of the promoter region of pfl-cys-prx gene was elevated at the trophozoite/schizont stages according to the activation of gene expression, and that the PGHCN5 protein, the histone acetyltransferase of *P. falciparum* [29], was recruited into the PRE sequence [16]. It has been suggested that the mechanisms of histone modification and epigenetic regulation for transcription are as well conserved in the parasite as they are in other eukaryotes [30], and that the degree of histone acetylation reflects the activity of transcription [16,31]. The PREBP protein might interact with the PGHCN5 and regulate the chromatin structure for the effective activation of gene expression.

In the transient reporter assay, the transfectant with the plasmid p1–10R-Acis-PREBP-full showed a significant decrease in firefly luciferase activity, indicating that the enhancer activity of PREBP depends on the PRE sequence. However, this level of decrease was not as low as that in the transfectant of p1–10R-Mock and p1–10R-No.2. This some elevation of expression might be the effect of non-specific binding of PREBP to the promoter sequence in the plasmid, which does not form a chromatin structure. Our previous report described the measurement of the promoter activity of the pfl-cys-prx 5’ sequence with and without the PRE element [16]. The plasmids used in the previous study did not express PREBP. The promoter without PRE showed about 70% of promoter activity of that with the PRE element. Thus, it is suggested that the PRE element was more effective with an excess of PREBP.

It is also possible that PREBP regulates the numbers of genes other than pfl-cys-prx. In mammals, the expression of I-Cys-PxR is induced by oxidative stresses through the function of the antioxidant response element in the promoter [32]. Oxidative stress, however, could not enhance the expression of the pfl-cys-prx gene (Komaki-Yasuda K, unpublished data) in the intraerythrocytic stage. Thus, its expression is rigidly regulated in accordance
with the progression of the parasite cell cycle in the intraerythrocytic stages [12,13,15]. The expression of other genes regulated by PREBP might be elevated at the same time-point in the trophozoite/schizont stages. In future studies, it would be interesting to use ChIP-seq methods to discover the genes that are regulated by PREBP binding [33]. The structural analyses would elucidate the means by which PREBP recognizes and specifically binds to the sequence of cis-element in the AT-rich non-coding DNA sequence in *P. falciparum*. It would also be of great interest to detect the co-factors that function with PREBP in the process of transcriptional activation.

It is known that an enormous variety of specific transcription factors is present across different eukaryotic lineages [4,34]. Thus, only a small number of specific transcription factors have been identified in *Plasmodium* spp., and across the range of protist species [4]. The discovery of PREBP showed the possibility that unknown specific transcription factors with unique DNA binding domain had evolved in the protists. As a model case of the general transcriptional regulatory mechanisms in *P. falciparum*, further analysis of the transcriptional activation of *pf1-cys-prx* will yield profound insights into the gene activation mechanisms present during the intraerythrocytic growth of the parasite. Further detailed characterization of novel components of the transcriptional machinery in malaria parasites may help lead to the black box of evolution of specific transcription machinery in the protists, and may pave the way for the identification of unique drug targets for alternative malaria chemotherapies.

Materials and Methods

Parasite Culture

The FCR-3 strain of *P. falciparum* was cultured in human O erythrocytes and RPMI medium (Life Technologies, Carlsbad, CA) supplemented with 10% human serum using the modified method of Trager and Jensen [35]. For the preparation of nuclear extracts, parasites were cultured in RPMI medium supplemented with 0.5% Albulamx (Life Technologies). Growth synchronization was achieved with 5% D-sorbitol [36].

Parasite Nuclear Extracts and Cytoplasmic Fractions

Parasite nuclear extracts were prepared with modifications according to the previous study [37]. In brief, parasites were released from red blood cells by treatment with PBS containing 0.05% saponin and washed twice in PBS. The parasite pellet was resuspended in ice-cold lysis buffer (20 mM Hepes, pH 7.8, 10 mM KCl, 1 mM EDTA, 1 mM DTT, 1 mM PMSF and 0.65% Nonidet P-40) and incubated for 5 min on ice. Nuclei were pelleted at 2,500 *g* for 5 min, and the supernatants were saved as cytoplasmic fractions. The nuclear pellet was washed twice with lysis buffer before resuspension in one pellet volume of extraction buffer (20 mM Hepes, pH 7.8, 800 mM KCl, 1 mM EDTA, 1 mM DTT and 1 mM PMSF) containing Complete Protease Inhibitor Cocktail (Roche Applied Science). After vigorous shaking at 4°C for 30 min, the extract was cleared by centrifugation at 13,000 *g* for 30 min. The supernatant (nuclear proteins) was diluted with an equal volume of dilution buffer (20 mM Hepes, pH 7.8, 1 mM EDTA, 1 mM DTT and 30% glycerol), then immediately frozen in liquid nitrogen and stored at −80°C.

**Electrophoretic Mobility Shift Assay (EMSA)**

To generate the labeled probe for EMSA, the 102-bp putative enhancer sequence (PRE; region between 490–591 bp upstream of the ATG translational start site of *pf1-cys-prx*) in the 5′ region of *pf1-cys-prx* was labeled with [γ-32P]ATP with T4 polynucleotide kinase as described previously [16]. For EMSA reactions, each protein sample was incubated with 10 fmol of labeled probe in EMSA buffer (20 mM Hepes, pH 7.8, 60 mM KCl, 0.5 mM EDTA, 2 mM DTT, 0.1% Triton X-100 and 10% glycerol) in a 20 µl reaction volume for 20 min at 25°C. To decrease nonspecific binding, 0–0.025 μg/µl of poly-dI-dC was added to the reaction mix. To stabilize the DNA-protein complex, 0–0.0125 μg/µl of poly-L-lysine was also added to the reaction mix. Binding reactions were analyzed by 6% polyacrylamide gel electrophoresis in 0.25 × TBE buffer at 4°C. For the super-shift assay, 0.9–27.0 μg of antibodies was added into the reaction mix of EMSA and incubated overnight at 4°C before electrophoresis.

Gels were dried, and the result was visualized using a BAS-2000 system (Fuji Film, Tokyo, Japan).

Purification of the Specific DNA-Binding Protein

Throughout all of the purification steps, the activities of PREBP were checked by EMSA with the labeled PRE sequence. The frozen stock of nuclear extract (110 ml, obtained from 5 × 1011 parasite cells synchronized at trophozoite/schizont stage) was thawed and dialyzed against buffer 1 (60 mM KCl, 20 mM Hepes (pH 7.8), 0.5 mM EDTA, 2 mM DTT, 0.1% Triton X-100, 10% glycerol, 1 mM PMSF). The dialyzed nuclear extract was loaded onto HiPrep Q column (GE Healthcare, Buckinghamshire, UK) with an HPLC system (AKTA Prime, GE healthcare). The column was washed with the same buffer and the bound materials were eluted with 200 ml of an increasing (60–1000 mM) KCl gradient. Activity was eluted at ~0.3 M KCl. Active fractions were pooled, diluted with KCl-free buffer 1 until they reached 60 mM KCl concentration, and loaded onto the HiTrap CM column (GE Healthcare) using the AKTA prime system. After the column was washed with buffer 1, the bound protein was eluted with stepwise increasing Na2SO4 (50 ml of buffer 2 (0.3 M Na2SO4, 20 mM Hepes (pH 7.8), 0.5 mM EDTA, 2 mM DTT, 0.1% Triton X-100, 10% glycerol, 1 mM PMSF), 50 ml of buffer 2 with 0.6 M Na2SO4 and 50 ml of buffer 2 with 1.2 M Na2SO4). Active fractions (with 0.3 M Na2SO4) were pooled and buffer 2 was added with 3 M Na2SO4 to reach 1.2 M Na2SO4 concentration and then loaded onto a RESOURCE PHE column (GE Healthcare) using the AKTA Prime system. The column was washed with buffer 2 containing 1.2 M Na2SO4 and the bound protein was eluted with 20 ml of a decreasing (from 1.2 to 0.1 M) Na2SO4 gradient and 10 ml of buffer 2 containing 0.1 M Na2SO4. The active fractions (with ~0.1 M Na2SO4) were pooled and dialyzed against buffer 1, then loaded onto a Mono S column (GE Healthcare) using the SMART system (GE Healthcare). The bound proteins were eluted with 3 ml of a linear gradient from 60 to 1000 M KCl. Active fractions (with ~0.3 M KCl) were pooled, diluted to 60 mM KCl concentration and applied to the DNA affinity purification.

As a matrix for affinity purification, the biotinylated PRE DNA sequence was tethered to magnetic streptavidin-coated Dynabeads M280 (Life Technologies) according to the supplier’s instructions. The dialyzed fractions were supplied with 25 μg/ml of poly(dI-dC) and 2.5 μg/ml of poly-L-lysine, then incubated with 0.5 mg of Dynabeads (100 pmol oligo/mg beads) on a rotating wheel for 3 h. Dynabeads were collected using a magnetic stand and washed twice with 1 ml of wash buffer 1. Bound proteins were eluted from the beads with 10 μl of elution buffer 1 (20 mM Hepes, pH 7.8, 500 mM KCl, 1 mM EDTA, 1 mM DTT, 1 mM PMSF, 0.01% Triton X-100, 5% glycerol) twice and 10 μl of elution buffer 2 (elution buffer 1 containing 1.0 M KCl). The flow through fraction was pooled and applied onto the DNA-tethered Dynabeads, and the same washing and elution process was repeated twice.
Estimation of the Molecular Size of PREBP

Two hundred μl of parasite nuclear extract was dialyzed against buffer 1 (60 mM KCl, 20 mM Heps (pH 7.8), 0.5 mM EDTA, 2 mM DTT, 0.1% Triton X-100, 10% glycerol, 1 mM PMSE), added to 25 μg/ml of poly(dI-dC) and 2.5 μg/ml of poly-L-lysine, and treated with 0.2 mg of the DNA-affinity magnetic beads tethered to the 102-bp PRE sequence. The bound protein was eluted with 8 μl of the buffer 1 with 0.5 M of KCl, and applied to 5–20% SDS-PAGE. The gel pieces were cut from the gel according to the 8 grades of molecular weights, ranging 10–100 kDa. The proteins were eluted from each gel piece by treatment with 200 μl of elution buffer (50 mM Tris pH 7.9, 150 mM NaCl, 0.1 mM EDTA, 0.1% SDS, 0.05 mg/ml BSA), precipitated with 80% of acetone, and the precipitated proteins were dissolved in 100 μl of the buffer (50 mM Hepe7.9, 10 mM DTT, 6 M Guanidine HCl). The proteins were then renatured by removing the guanidine HCl by dialysis against buffer 1. To concentrate the DNA binding protein, each renaturated fraction was supplied with 25 μg/ml of poly(dI-dC) and 2.5 μg/ml of poly-L-lysine, then again treated with 0.02 mg of the DNA-affinity magnetic beads tethered to the 102-bp PRE sequence. The bound protein was eluted with 2 μl of the buffer 1 with 0.5 M of KCl. The PRE-binding activity of each fraction was checked by EMSA.

Mass Spectrometry and Data Analysis

Seven μl of the final fraction obtained from the purification steps was subjected to 5–20% SDS-PAGE and stained with silver stain system specialized for mass spectrometry (SilverQuest, Life Technologies). The 3 candidate bands were excised as gel slices, digested with trypsin, and analyzed by nanoliquid chromatography and mass spectrometry on an LTQ-Orbitrap XL tandem mass spectrometer (Thermo Fisher Scientific, San Jose, CA). Data were searched against the Mascot concatenated forward-and-reversed v3.46 International Protein Index database (Matrix Science, London, UK) and collated into nonredundant lists using Scaffold software (Proteome Software, Portland, OR).

Antibodies

Rabbit polyclonal antiserum was raised against a cocktail of two synthetic peptides (CGMKDDIENLKELIE and CSGNKNNDIDKA, corresponding to the amino acid numbers 699–849) was expressed as a glutathione S-transferase (GST) fusion protein with pGEX-6P-1 vector system (GE Healthcare) and Escherichia coli BL21 strain. The GST-tag on the fusion protein was removed by PreScission protease treatment and GST-glutathione affinity chromatography (GE Healthcare). The recombinant protein was further purified with gel filtration chromatography with a HiPrep Sephacryl S-100 column (GE Healthcare) and used for rabbit immunization. The antibodies were affinity-purified with the HiTrap NHS-activated HP column (GE Healthcare) and used for rabbit immunization. The antibodies against recombinant PREBP protein (the antigen) according to the manufacturer’s instructions. The purified antibody was defined as anti-PREBP pep. In addition, Rabbit polyclonal antiserum was raised against the partial sequence) and affinity-purified as described previously [16]. The antibodies were defined as anti-PREBP rec. The antibody against antibody was defined as anti-PREBP rec. The antibody against FLAG (Sigma-Aldrich, St. Louis, MO) was commercially supplied.

Construction of Plasmids

For construction of plasmid vectors for the overexpression of the candidate proteins, each gene was first amplified by PCR with the parasite cDNA as a template. The sequence of each primer used for the PCR is indicated in Table S3. FLAG-1F and His-1R were used for protein No. 1; FLAG-2F and His-2R were used for protein No. 2; FLAG-3F and His-3R were used for protein No. 3; and FLAG-4F and His-4R were used for protein No. 4. These PCR products contain the sequences for the FLAG-tag and His-tag at the 5’ and 3’ ends, respectively. For insertion of these fragments into the pHC1 expression vector (MRA-4, Malaria Research and Reference Reagent Resource [MR4] Center [30]), the Xhol sites were added at both the 5’ and 3’ ends by reamplification of these fragments by PCR using the primers FLAG-Xhol and His-Xhol. These fragments were then inserted into the Xhol site of the pHC1 and the constructed plasmids were termed pHC1-No.1, pHC1-No.2, pHC1-No.3 and pHC1-No.4. The direction of the inserted fragments was checked by sequencing analysis.

For the overexpression of full-length PREBP, the complete ORF of PREBP gene was amplified using the primers, and PREBP-F and PREBP-R with the parasite cDNA as a template. The amplified fragment contained FLAG-tag at the 5’ end, HA tag at the 3’ end and the Xhol sites at both the 5’ and 3’ ends. The fragments were inserted into the Xhol site of pHC1. Furthermore, for the following operation of the plasmid, the HindIII site in the ORF was disrupted by a Site-Directed Mutagenesis System (Promega), avoiding the non-synonymous substitution of amino acid, with the primers AhindIII-F and AhindIII-R. This plasmid was termed pHC1-PREBP-full. The direction of the inserted fragments was checked by sequencing analysis.

For the transient luciferase assays, a basal plasmid, p1–10R was first constructed. The luciferase expression cassette was amplified by PCR with the primers 1-cys5’-F and 1-cys3’-R. The plc-Fluc plasmid [16] was used as a template. This fragment contains 0.8-kbp of the 5’ region of pf1-cys-prx, 1.7-kb of firefly luciferase gene and 0.9-kb of 3’ region of pf1-cys-prx. This fragment was digested by SfiI and Kpnl and subcloned into the same restriction site of pUC19.

To obtain the p1–10R-PREBP-short, p1–10R-PREBP-full, p1–10R-Mock and p1–10R-No.2, the expression cassettes were excised from pHC1-No.4 (for p1–10R-PREBP-short), pHC1-PREBP-full (for p1–10R-PREBP-full), pHC1 (for p1-10R-Mock) and pHC1-No.2 (for p1–10R-No.2) by HindIII and inserted into the same restriction site of p1–10R. The direction of the inserted fragments was checked by sequencing analysis.

To obtain the p1-10RAPRE-PREBP-full plasmid, the same procedure for preparation of p1-10RAPRE-PREBP-full was repeated, differing only in that plc-Fluc-Δcαs [16], which lacks the 102-bp PREBP sequence, was used as the template for the first PCR.

Stable Transgene Expression in P. falciparum and Immuno-Affinity Purification of the Expressed Protein

The plasmids for overexpression were transformed into the parasite by electroporation at 0.310 kV and 975 μF in a 0.2 cm gap cuvette with a Gene Pulser II (Bio Rad, Hercules, CA) as described in a previous study [39]. Drug selection with 25 mg/ml of pyrimethamine was started 48 h after transfection. Parasites that stably expressed the transformed gene, were then selected. For immuno-affinity purification with anti-FLAG antibody, ~1.5 x 10^9 of transgenic parasite-infected erythrocytes were lysed with phosphate-buffered saline (PBS) containing 0.05% saponin. The parasite pellet was washed several times with PBS and diluted with 1000 μl of homogenizing solution (20 mM Tris-HCl (pH 7.5),
Each transfection cuvette contained 100 μg of plasmid expressing Renilla luciferase, as a control for efficiency normalization. The plasmids were transfected into parasites with a Gene Pulser II (Bio-Rad). After electroporation, the parasites were incubated twice in PBS, and suspended in 50 mM Tris-Cl (pH 8.0), 250 mM NaCl, 10% Glycerol, 0.8% Triton X-100 and then the bound proteins were eluted with the 30 μl of elution buffer (wash buffer with 0.5 mg/ml of the FLAG peptide). The eluents served as immunoprecipitated proteins.

Immunoprecipitations and Western Blotting

Immunoprecipitations (IP) and Western blotting assays with anti-PREBP peptide antibody were performed with the Rabbit IgG TrueBlot system (eBioscience, SanDiego CA). For the IP experiments, 15 μg of nuclear extract or cytoplasmic fraction from synchronized parasites was diluted with 1 ml of IP buffer (20 mM Tris-Cl (pH 7.5), 150 mM NaCl, 1% NP-40, 1 mM PMSF with Protease Inhibitor (Complete Mini, Roche Applied Science, Basel, Switzerland) and lysed by sonication. The parasite extracts were analyzed with a Dual-Luciferase Reporter Assay System (Promega) and Turner luminometer according to the manufacturer’s instructions. The firefly luciferase activity was normalized to that of Renilla luciferase, which was derived from the pHCl-Rhuc transfecant.

Statistical Analysis

Differences were evaluated using Student’s t-test. P<0.01 was considered to be statistically significant.

Accession Numbers

PlasmidDB (www.plasmidDB.org) accession numbers for genes and proteins discussed in this publication are: Pf3D7_0602900; PREBP (P. falciparum); Pf3D7_1011800; PREBP (P.berghei); PBANKA_121020; PREBP (P. yoelii); PY05323; PREBP (P. vivax); PVX_094810; PREBP (P. knowlesi); PKH_061130; PREBP (P. cynomolgi); PCYB_062160. Genbank (www.ncbi.nlm.nih.gov/genbank/) accession numbers for genes and proteins discussed in this publication are: PREBP (Cryptosporidium parvum): EAK90228.1; PREBP (Theileria parva): EAN34110.1; PREBP (Babesia bovis): EDO72571.1.

Supporting Information

Figure S1 Excision of gel slices corresponding to candidate proteins. Seven μl of final fractions obtained from purification steps were loaded onto the 5–20% SDS-PAGE. The gel was stained with silver-stain kit (SilverQuest, Invitrogen) specialized for mass spectrometry. Three candidate bands (indicated with arrows) were excised as gel pieces and subjected to trypsin digestion and LC-MS/MS analysis. Sizes are indicated in kDa on the right. (TIF)

Figure S2 Preparation of the recombinant candidate proteins using the gene cell-free translation system. A) Immunopurification of the recombinant proteins from cell-free translation reaction mix. The expressed recombinant proteins were eluted from the reaction mixes through immunoprecipitation with the anti-FLAG antibody. The precipitated proteins were eluted with FLAG peptides and 5 μl (~1 μg) of eluents were subjected to 5–20% SDS-PAGE and blotted onto PVDF membrane. Blotted proteins were detected with the anti-FLAG antibody. The bands corresponding to the recombinant proteins are indicated with white arrowheads. Protein Nos. 1, 2, 3, 4, 5, 7, 9, 10, 11, 13, 14, 15 and 16 clearly showed the band of the expected molecular weight. Protein No. 12 could not be prepared because the DNA sequence for the gene of protein No. 12 was not amplified by the PCR reaction. Sizes are indicated in kDa at the right of each panel. See Table S2 for detailed descriptions for each protein. B) EMSA with the immunopurified recombinant proteins. The same 32P-labeled probe was used in Figure 2 in each assay. Lane N is probe only. EMSA was performed with 2.0 μl (~0.05 μg) of the fraction of each roughly purified recombinant protein. The reaction mix for EMSA did not contain any non-specific competitor DNA. Lane P is the positive control of the assay with 0.3 μg of nuclear extract derived from the parasite synchronized at the trophozoite/schizont stage. Positions of the free probe and shifted band corresponding to the PREBP-PRE complex are indicated on the left. No recombinant protein showed any shift-band of comparable size to that of the positive control. (TIF)
cells during the erythrocytic stage.

The deduced molecular weight of PF3D7_1011800 is 132 kDa. The peptides detected in mass spectrometry are indicated with gray shading. The predicted K-homology (KH) domains are indicated with open boxes. The start and end points of the partial ~60 kDa recombinant protein, which was expressed in the transgenic parasite, are indicated by arrows.

Figure S4 Patterns of expression of PREBP mRNAs in *P. falciparum* cells during the erythrocytic stage. Parasite cultures were tightly synchronized, and parasite-infected erythrocytes were harvested at the indicated times for total RNA extraction and cDNA synthesis. Images of various stages of parasite growth at indicated times are shown in the lower panel. Time 0 corresponds to the final synchronization by D-sorbitol treatment. Real-time quantitative RT-PCR was performed with cDNA templates, and the values recorded were normalized to the amount of 18 S rRNA in each sample measured in the same RT-PCR run. Patterns of expression of *pf1-cys-prx* mRNAs which were measured using the same cDNA sample are indicated as a sub-panel within the main panel.

Figure S5 Expression level of *pf1-cys-prx* in parasites that overexpress the central ~60 kDa region of PREBP. The parental parasite and the pHCl-No.4 transfectant parasite cultures were synchronized at the trophozoite/schizont stage, and parasite-infected erythrocytes were harvested for total RNA extraction and cDNA synthesis. Real-time quantitative RT-PCR was performed with the cDNA templates and specific primers for *pf1-cys-prx*, and the values recorded were normalized to the amount of 18 S rRNA in each sample measured in the same RT-PCR run. The relative amounts of PREBP mRNA were also measured by RT-PCR using same cDNA samples and indicated by the small panel on the right. Data are shown as the means of three independent assays. Error bars represent standard deviations.

Figure S6 Model scheme for PREBP interaction with PRE sequence and activation of the transcription of *pf1-cys-prx*.

Table S1 Summary of the purification of PREBP from the parasite nuclear extract.

Table S2 List of detected *P. falciparum* proteins by LC-MS/MS.

Table S3 List of primers used for the plasmid constructions.

Method S1 Preparation of recombinant proteins using the cell free translation system.

Method S2 Quantitative real-time PCR.

Acknowledgments

We are grateful to Dr. T. Yasuda of the National Institute of Radiological Sciences (Chiba, Japan) and Dr. K. Yano of National Center for Globol Health and Medicine (Tokyo, Japan) for helpful and critical suggestions and comments on this work.

Author Contributions

Conceived and designed the experiments: KKY MO KN SIK SK. Performed the experiments: KKY. Analyzed the data: KKY MO KN SIK. Contributed reagents/materials/analysis tools: KKY MO KN SIK. Wrote the paper: KKY SK.

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