Structural Determinants of DNA Binding by a P. falciparum ApiAP2 Transcriptional Regulator

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Putative transcription factors have only recently been identified in the Plasmodium spp., with the major family of regulators comprising the Apicomplexan Apetala2 (AP2) proteins. To better understand the DNA-binding mechanisms of these transcriptional regulators, we characterized the structure and in vitro function of an AP2 DNA-binding domain from a prototypical Apicomplexan AP2 protein, PF14_0633 from Plasmodium falciparum. The X-ray crystal structure of the PF14_0633 AP2 domain bound to DNA reveals a β-sheet fold that binds the DNA major groove through base-specific and backbone contacts; a prominent α-helix supports the β-sheet structure. Substitution of predicted DNA-binding residues with alanine weakened or eliminated DNA binding in solution. In contrast to plant AP2 domains, the PF14_0633 AP2 domain dimerizes upon binding to DNA through a domain-swapping mechanism in which the α-helices of the AP2 domains pack against the β-sheets of the dimer mates. DNA-induced dimerization of PF14_0633 may be important for tethering two distal DNA loci together in the nucleus and/or for inducing functional rearrangements of its domains to facilitate transcriptional regulation. Consistent with a multisite binding mode, at least two copies of the consensus sequence recognized by PF14_0633 are present upstream of a previously identified group of sporozoite-stage genes. Taken together, these findings illustrate how Plasmodium has adapted the AP2 DNA-binding domain for genome-wide transcriptional regulation.

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Keywords: malaria; Plasmodium falciparum; AP2 domain; X-ray crystallography; domain swapping

Introduction

The malaria parasite Plasmodium falciparum exhibits a complex life cycle that involves infection of both a vertebrate human host and an invertebrate mosquito vector. The various host cell types and morphological transitions involved in the development of the parasite suggest the utilization of fine-tuned regulation of gene expression at all stages. However, dissection of transcriptional regulation in P. falciparum and other related apicomplexan parasites has proven difficult because their genomes possess few proteins with detectable homology to established transcription factors in model eukaryotes. To date, only one major family of transcriptional regulators has been identified, the Apicomplexan Apetala2 (ApiAP2) family, which has undergone a lineage-specific expansion in the apicomplexan phylum.²

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Abbreviations used: AP2, Apetala2; ApiAP2, Apicomplexan AP2; ERF, ethylene response factor; dsDNA, double-stranded DNA; SeMet, selenomethionine; EMSA, electrophoretic mobility shift assay; PBM, protein binding microarray; GST, glutathione S-transferase.

ApipAP2 proteins vary in size from hundreds to thousands of amino acids, and each contains at least one copy of a conserved ~60-residue AP2 domain that is weakly homologous to the plant AP2 domain. In the plant Arabidopsis thaliana, AP2 domains are the second most common DNA-binding domain after the Myb domain and are typically found in small proteins on the order of 300 amino acids.³

³Plant AP2 domains mediate specific DNA recognition in all members of the AP2/ethylene response factor (ERF) family, which control development and stress responses in species throughout the plant
 kingdom. Typically in plants, the AP2 domains are found as one- or two-domain arrangements in a given protein, and two-domain AP2 proteins have been shown to contact up to 17 contiguous base pairs of DNA. An NMR solution structure of a single A. thaliana AP2 domain in complex with DNA reveals a mode of DNA binding where three β-strands contact the DNA backbone and specific bases within the major groove. The β-strands are stabilized by a C-terminal α-helix that does not contact the DNA.

Despite having limited homology to plant AP2 domains, Apicomplexan AP2 domains bind DNA with high sequence specificity and can function as regulators of transcription. There are a total of 26 ApiAP2 proteins predicted in the Plasmodium genus, and although a given AP2 DNA-binding domain can be highly conserved across related species, there is sparse homology outside of the DNA-binding domain. Presumably, the ApiAP2 proteins must also contain activation domains and additional protein–protein interaction domains, although there are no additional Pfam domains predicted for any of these proteins.

Furthermore, paralogous AP2 domains from Plasmodium spp. share little sequence identity with each other, in contrast to those from Arabidopsis where paralogous AP2/ERF proteins can be functionally grouped on the basis of their primary sequence identity as well as their DNA-binding preferences. This divergence of AP2 domains in sequence identity as well as their DNA-binding actionally grouped on the basis of their primary

Results and Discussion

Structure of the AP2 domain of PF14_0633 specifically bound to dsDNA

To elucidate the mechanism of DNA binding of an Apicomplexan AP2 domain, we determined the 2.2 Å resolution crystal structure of the AP2 domain of PF14_0633 bound to an 8-bp dsDNA comprising its consensus binding sequence TGCATGCA (Fig. 1 and Supplemental Fig. 1). Single-wavelength anomalous dispersion phasing of a crystal of selenomethionine (SeMet)-incorporated AP2 domain bound to DNA produced a high-quality experimental electron density map into which a partial molecular model was built. This model was used as a molecular replacement search model to phase the higher-resolution diffraction data from a native complex, which yielded electron density maps that allowed building of the final model (Fig. 1; Table 1).

The consistent unit contains two molecules of the AP2 domain of PF14_0633, with each monomer bound to an individual dsDNA molecule. The overall structure of the DNA-bound AP2 domain of PF14_0633 retains many of the canonical features of similar DNA-binding domains for which structures have been determined, such as ATERF1 (A. thaliana ethylene response factor 1) and the yeast PI-SceI homing endonuclease. Three anti-parallel β-strands from the AP2 domain of PF14_0633 wrap into the major groove of the bound dsDNA and contain all residues making base-specific contacts. A flexible loop connects these strands with an extended α-helix, which supports the opposing face of the β-sheet. However, our structure demonstrates that PF14_0633 dimerizes through a three-dimensional domain-swapping mechanism in which the α-helix of one protomer is packed against the β-sheet of its dimer mate. Similar dimerization was not previously observed in the NMR solution structure of the related plant AP2 protein ATERF1. The interface derived from domain swapping results in burial of a large surface area (~3140 Å²) of primarily
hydrophobic residues. Dimerization of the PF14_0633 AP2 domain also aligns Cys76 residues of each monomer with one another with sufficient proximity to allow disulfide bond formation (Fig. 1b). We note that the Cys76 residue is conserved in all orthologues of PF14_0633 in Plasmodium spp. but is not conserved in other related apicomplexan species.

Several contacts between the PF14_0633 AP2 domain and its DNA-binding sequence appear to be important to binding specificity (Fig. 1c, summarized in Supplemental Table 1). Residues of particular note are Asn72, Arg74, Arg88, and Ser90, which make base-specific contacts to the consensus binding sequence. Of these, only one residue (Asn72) was previously implicated in DNA binding by computational modeling.2 The two arginine side chains can each contact both strands of DNA (O6 atoms of dG2 and dG6) (Fig. 1c). The side chains of Asn72 and Ser90 contact a central AT base pair, each to one strand. All four of these residues are invariant among apicomplexan orthologues of PF14_0633 with BLAST expect scores <1e–05, including AP2 domains from Plasmodium spp. and Toxoplasma, Theileria, Babesia and Cryptosporidium genera. The AP2 domain in this protein therefore appears to have conserved specific binding to the TGCATGCA consensus DNA over a large evolutionary distance separating these apicomplexan species by conservation of four key DNA contact residues. Indeed, the distantly related Cryptosporidium PF14_0633 orthologue binds specifically to the same TGCATGCA sequence.6 In addition to these base-specific contacts, several phosphodiester backbone contacts are made, most notably His94, which is located in the hinge region between the α-helix and the β-sheet of the AP2 domain. His94, Pro95, and several adjacent residues are all well conserved among the orthologous PF14_0633 AP2 domains. A common mechanistic model for domain swapping implicates strain in Pro-containing hinge regions, which relies on Pro isomerization to help drive conformational changes during dimerization.17,18 The interaction of His94 with the phosphodiester backbone may influence the conformation of the adjacent Pro95 side chain and act as a trigger for domain swapping (discussed further below).

**Fig. 1.** Structure of the AP2 domain of PF14_0633 specifically bound to dsDNA. (a) Secondary structural elements of the AP2 domain of PF14_0633 are depicted above the primary sequence as arrows (β-strands) and a rectangle (α-helix). Residues that make base-specific contacts to dsDNA are underlined. (b) (Top) Ribbon diagram of the crystal structure of a domain-swapped dimer of the AP2 domain of PF14_0633 (one monomer shaded blue, another red) bound to independent dsDNA molecules (shaded green). Disulfide-bonded Cys76 residues are indicated and shaded yellow. (Bottom) Orthogonal view with protein electrostatic surface potentials depicted (blue, electropositive; red, electronegative). (c) DNA contact map depicting direct base-specific contacts between the Asn72, Arg74, Arg88, or Ser90 side chains and the indicated DNA bases.
ture, we generated four variants of the PF14_0633 AP2 domain by substituting alanine for the residues observed to make base-specific DNA contacts (Asn72Ala, Arg74Ala, Arg88Ala, and Ser90Ala). These variants were subjected to electrophoretic mobility shift assays (EMSAs; Fig. 2) and protein binding microarrays19,20 (PBMs; Supplemental Fig. 4) to assess whether these amino acid substitutions altered DNA-binding stability and specificity in solution. All four variants were found to have circular dichroism spectra highly similar to that of the wild-type AP2 domain (Supplemental Fig. 2), indicating that functional differences were not due to differences in protein folding.

We assessed the affinity of the purified wild-type PF14_0633 AP2 domain for its consensus sequence using EMSA (Supplementary Fig. 3). Binding of this domain to two probes (28 and 122 bp in length), each containing a core 28-bp region derived from the upstream region of the pfi0540w gene of the P. falciparum 3D7 genome, as used previously,6 was measured (see Materials and Methods). The $K_{d,app}$ of the wild-type domain for its TGCATGCA target sequence was determined to be 0.5–1.0 μM. Mutation of the central two AT base pairs of the consensus sequence to GC significantly reduced the binding affinity of PF14_0633, confirming that these bases are critical for specific DNA recognition.

The ability of the variant AP2 domains (Asn72Ala, Arg74Ala, Arg88Ala, and Ser90Ala) to bind specifically to dsDNA was evaluated next. The 28-bp dsDNA probe (described above) was incubated with increasing concentrations of the wild-type protein or its variants. A representative gel image for wild-type and Arg74Ala AP2 domains is shown in Fig. 2a, where the wild-type protein forms a specific complex with dsDNA with a $K_{d,app}$ of 0.5 μM (primary shift). A slower-mobility form of the wild-type protein is observed at concentrations of 2.5 μM and above (secondary shift). Quantification of the percentage of shifted probe for each Ala-substituted variant AP2 domain demonstrated that all bind dsDNA very poorly, binding specifically only at concentrations just below those where nonspecific binding to both single-stranded DNA and dsDNA occurs (Fig. 2b). This is consistent with the protein–DNA interface observed in the crystal structure being used in solution. We also assessed the contribution of the AT-hook element (residues 39–50) to DNA binding by extending the expressed consensus sequence to GC significantly reduced the binding affinity of PF14_0633, confirming that these bases are critical for specific DNA recognition.

### Table 1. Data collection, phasing, and refinement statistics of the AP2 domain of PF14_0633 specifically bound to dsDNA

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Values in parentheses are for the highest-resolution shell.

- Data statistics from one crystal of native or SeMet PF14_0633.
- $R_{sym} = \sum_j |I_j - \langle I \rangle| / SI_j$, where $I_j$ is the intensity measurement for reflection $j$ and $\langle I \rangle$ is the mean intensity for multiply recorded reflections.
portion of PF14_0633 to include residues 38–123. AT-hooks are short auxiliary elements that contact DNA in a non-sequence-specific manner and are commonly found adjacent to specific DNA-binding domains. The addition of the AT-hook motif did not significantly affect the binding affinity of wild-type PF14_0633 to the 28-bp probe (Fig. 2b). These results demonstrate that specific DNA binding was weakened in the variant AP2 domains, consistent with elimination of base-specific contacts provided by each side chain.

We reasoned that alteration of the individual residues that make base-specific contacts to the DNA-binding site might alter the sequence specificity of the PF14_0633 AP2 domain. To address this, we assayed glutathione S-transferase (GST)-tagged forms of these variants of PF14_0633 using PBMs, and sequence-specific binding was determined as previously described. As expected, the wild-type AP2 domain bound the consensus binding sequence TGCATGCA. Surprisingly, two of the four variants (Asn72Ala and Ser90Ala) also bound this same sequence, albeit more weakly (Supplemental Fig. 4). This result implies that these residues are not necessary for DNA-binding specificity. Furthermore, it also suggests that the PBM methodology is far more sensitive for detecting lower-affinity DNA-binding events, which we could not detect by EMSA. In contrast to the Asn and Ser variants, DNA binding was completely abrogated with both Arg variants.

Taken together with the results from the EMSAs comparing wild-type and mutant DNA probes (Fig. 2b; Supplemental Fig. 3), the observation that Asn72Ala and Ser90Ala variants still recognize the consensus binding sequence with high statistical significance indicates that base-specific contacts to the central AT base pairs are not necessary for specific DNA binding. Instead, an appropriate balance of DNA rigidity/flexibility provided by central AT base pairs in the palindromic sequence may be important for specific binding. In contrast, Arg74 and Arg88 are required for sequence-specific DNA binding. These data imply that the identified consensus binding sequence represents the minimal binding sequence for the AP2 domain of PF14_0633, which is largely determined by the base-specific contacts of both Arg74 and Arg88.

**DNA binding stimulates dimerization by domain swapping within the PF14_0633 AP2 domain**

We first addressed these questions by assessing whether a disulfide bond is formed at the dimerization interface upon binding to dsDNA. Under nonreducing conditions, no significant difference in disulfide bond formation was observed between the apo and dsDNA-bound forms of the AP2 domain of PF14_0633 (Fig. 3a, nonreducing gel: lanes 1, 3, and 5). As an alternate approach, we tested whether the homobifunctional cross-linker BM(PEG)3, which can covalently react with free thiols on exposed Cys residues, has a differential effect on the apo and DNA-bound forms of the AP2 domain. As shown in Fig. 3a, the presence of cognate dsDNA (containing the consensus binding sequence of PF14_0633’s AP2 domain, see Materials and Methods for details) stimulated cross-linking of two monomers to form a dimer approximately 50-fold relative to the apo domain (Fig. 3a, reducing gel: lane 4 versus lane 2). This effect is likely due to the proximal localization of the Cys residues by dimerization upon DNA binding. Incubation with non-cognate dsDNA also stimulated cross-linking (Fig. 3a, reducing gel: lane 6), but with a reduced efficiency relative to cognate dsDNA. In agreement with the observation that these Cys residues are buried and/or disulfide bonded upon DNA binding, accessibility of these Cys residues was stimulated 9- to 13-fold by the presence of dsDNA as observed by an iodoacetamide protection/2-nitro-5-thiocyanatobenzoic acid cleavage assay (Supplemental Fig. 5). These data indicate that DNA binding induces dimerization of the PF14_0633 AP2 domain.

Size-exclusion chromatography of the wild-type AP2 domain in the absence or presence of cognate or non-cognate DNA was used as an independent means of testing whether DNA binding stimulates dimerization in solution (Fig. 3b). The wild-type AP2 domain eluted at a volume consistent with the molecular size of a monomer (8.8 kDa) in the absence of DNA. In the presence of cognate (8 bp, 4.8 kDa) or non-cognate (17 bp, 10.6 kDa) dsDNA, the AP2 domain eluted at two distinct peaks each, at volumes consistent with an AP2 dimer bound to one or two dsDNA molecules. The presence of DNA in these elution peaks was confirmed by a decrease in the A280/A260 ratio from 1.56 (apo) to 0.56–0.59 (DNA-bound), as well as by the large increases in peak UV absorbance due to the relatively large extinction coefficients of dsDNA. Samples from both peaks of the cognate dsDNA-bound AP2 complexes were compared to the apo form via reducing and nonreducing 20% SDS-PAGE (Fig. 3b, inset). The faster-eluting species (marked as “⁎”, elution peak at 60.0 ml, 7% of total) contained a disulfide-bonded dimer, whereas the slower-eluting species (marked as “⁎⁎”, elution peak at 67.5 ml, 93% of total) was composed of a non-covalently bonded dimer (Fig. 3b, inset). Taken together, these experiments indicate that DNA binding stimulates the dimerization of the wild-type AP2 domain in solution.

In order to “capture” the domain-swapped form of the PF14_0633 dimer in solution, we engineered a triple substitution variant AP2 domain for fur-
The cross-linking studies. This protein variant (Cys76Ala, Ala86Cys, Asn118Cys) lacks the single native Cys residue in the dimerization interface but introduces one Cys residue on the domain-swapped α-helix and another in a spatially proximal loop between β-strands (illustrated in Fig. 3c). The introduced Cys residues allow the assessment of domain swapping between dimer mates whereby the β-sheet of one monomer can cross-link to the α-helix donated by the other monomer. As observed above with the wild-type PF14_0633 AP2 domain, the presence of dsDNA containing the consensus binding sequence stimulated cross-linking of the triple point variant (Fig. 3d, reducing gel: lane 4 versus lane 2). Again, a reduced level of dimerization is seen in the presence of nonspecific DNA (Fig. 3a, 563Plasmodium falciparum ApiAP2 Structure
We hypothesize that DNA binding to the domain swapping also occurs in solution conditions. These observations imply that reducing gel: lane 6). These observations imply that domain swapping also occurs in solution conditions. We hypothesize that DNA binding to the β-sheet may stabilize this portion of the domain, thus allowing the α-helix to adopt the extended conformation and form a domain-swapped dimer.

In light of these data, we propose a model in which the quaternary structure of the AP2 domain is influenced by specific DNA binding (Fig. 4). The DNA trigger could act by stabilizing the dimeric state in a preexisting monomer–dimer equilibrium of the AP2 domain, or, alternatively, DNA binding by an AP2 monomer could induce a conformational change that attracts a second monomer to bind. The AP2 dimer can be stabilized by several factors, including hinge region effects and stabilizing interactions made by the dimer interface or by ligand binding. In the case of the AP2 domain of PF14_0633, both of these factors could be important. First, Pro95 in the hinge region is in a trans conformation in the dimer crystal structure. If the closed monomeric form requires Pro95 to be in a cis conformation, isomerization could both relieve strain and provide energy to produce a conformational change. Notably, other residues in the hinge make backbone contacts with the DNA that may help drive DNA-binding stabilization of the dimeric form of the AP2 domain. Additionally, the structural and biochemical evidence presented herein indicates that the formation of a disulfide bond at the dimerization interface (Cys76) could also increase the stability of the dimer form. Lastly, observations from our three solution-based experiments confirm that binding dsDNA specifically stimulates the formation and/or stabilization of the domain-swapped dimer, as observed in the crystallographic structural model. The influence of these structural features indicates that the composition of the AP2 domain of PF14_0633 is well suited for the formation of a bona fide domain-swapped dimer, which may be relevant to its role in transcriptional regulation.

Many eukaryotic specific transcription factors form requisite dimers or higher-order multimers to assemble into their functional conformation. Some, such as Oct-1, Pit-1, and Ets-1, become competent for transcriptional regulation by differential homodimerization induced by the sequence and architecture of their DNA-binding sites. Notably, the human FOXP2 transcription factor forms a domain-swapped dimer, with known disease-associated mutations affecting residues of the domain-swapping interface. Similarly to the AP2 domain of PF14_0633, the orientation of the DNA-binding region of the domain-swapped FOXP2 dimer would only allow the binding of two distal DNA-binding sites, thus looping out intervening DNA. The precedence of such a structural explanation of transcriptional regulation by FOXP2 thus lends credence to a similar mechanism for the function of PF14_0633.

**Multiple PF14_0633 AP2-binding sites are found upstream of a set of sporozoite stage genes**

Previous work by Young et al. identified a statistically enriched DNA motif (PIM24.1) located upstream of 30 genes expressed in the mosquito sporozoite stage of the *P. falciparum* life cycle. This motif is highly similar to the consensus TGCATGCA motif recognized by PF14_0633, suggesting that the sporozoite-expressed ApiAP2 protein PF14_0633 may be the transcription factor responsible for the transcriptional regulation of these genes. Our structural model implies that binding to multiple TGCATGCA binding sites may facilitate the function of PF14_0633 as a transcriptional regulator by inducing protein dimerization. To test this possibility, we used the PBM-derived position weight matrix for the TGCATGCA consensus sequence to scan the 2-kb upstream regions of these 30 sporozoite genes for secondary occurrences of the TGCATGCA sequence (see Materials and Methods). Of the 30 sporozoite genes, 27 contained two or more matches to the motif (Supplemental Table 2). Furthermore, 24 of these upstream regions contained pairs of sites that are spaced sufficiently distant (>100 bp apart) to permit dimer-induced DNA looping (Supplemental Table 2).

The occurrence of multiple DNA-binding sites upstream of this subset of well-established sporozoite genes suggests that dimer binding may be important for the function of PF14_0633 in transcriptional regulation. Co-occurrence of motifs was also reported for the ookinete-specific *Plasmodium*...
transcriptional regulator AP2-O.\textsuperscript{7} One model is that the positioning of multiple binding sites in a promoter mediates a higher-affinity interaction of PF\textsubscript{14}\_0633 with DNA, facilitated by dimer formation. It remains to be tested whether higher-order structures are essential for function \textit{in vivo}. Structural rotations of the C-terminal \(\alpha\)-helix, upon dimer formation, may expose anticipated additional domains of PF\textsubscript{14}\_0633 that activate/repress transcription or modulate interaction with other auxiliary proteins required for regulation. Activation and repression domains that mediate transcriptional regulation have been defined only for a few examples of AP2/ERF domain proteins in the plant kingdom,\textsuperscript{26} and to date, no such characterization exists for any ApiAP2 protein. However, the wide range of predicted sizes of ApiAP2 proteins (200–4000 amino acids in \textit{Plasmodium} spp.) presumably speaks to the likely diversity of sequence space available to serve as necessary functional activation and repression domains.

We anticipate that the AP2 DNA-binding domain structure presented herein will serve as a model for assessing the binding interaction of other ApiAP2 proteins. PF\textsubscript{14}\_0633 orthologues from two highly diverged Apicomplexans, \textit{Cryptosporidium parvum} and \textit{P. falciparum}, have conserved their DNA-binding preference for the TGCATGCA sequence,\textsuperscript{6} the most over-represented DNA sequence in all sequenced Apicomplexa.\textsuperscript{27} Notably, the four key DNA-binding residues identified in the PF\textsubscript{14}\_0633 structure are conserved in the \textit{C. parvum} orthologue cdg2\_3490. This conservation suggests that insights gained on transcription factor domains in one apicomplexan species will be broadly applicable to understanding how related Apicomplexan AP2 domains contact DNA and regulate gene expression. Although it could be possible to model the structure of other AP2 domains upon that of PF\textsubscript{14}\_0633, the prediction of DNA-binding specificity and which residues make base-specific DNA contacts would likely not be similarly predictable. Important differences that should be taken into consideration are that many AP2 domains do not contain conserved cysteine residues that are present at the dimerization interface; neither do they possess appropriately placed proline residues that might facilitate the domain swapping identified here. Additional structural and biochemical characterization of these domains will be needed to elucidate these functional details. In addition, Apicomplexan AP2 domains possess affinities for a much more diverse set of DNA sequences than in plants where an overwhelming majority of AP2/ERF proteins bind the canonical GCC box (A/GCCGCC).\textsuperscript{9,11} This difference from plant AP2 domains demonstrates the need for structural and biochemical studies tailored towards Apicomplexan-derived proteins. Such studies are of great importance given that the ApiAP2 proteins are key regulators of parasitic developmental progression, and the absence of any homologous proteins in their mammalian hosts makes these proteins notable candidate drug targets.

### Materials and Methods

#### Purification of the PF\textsubscript{14}\_0633 AP2 domain and variants

The PF\textsubscript{14}\_0633 AP2 domain (residues 63–123) or its variants were expressed as a GST-fusion protein as previously described.\textsuperscript{6} Cells were lysed by sonication in GST lysis buffer (50 mM Tris, pH 8.0, 150 mM NaCl, and 10% glycerol) containing 1 mM PMSF and 1 mM benzamidine. Proteins were purified from the soluble fraction by binding to glutathione Sepharose 4B resin (GE Healthcare) and eluting with GST lysis buffer containing 20 mM reduced l-glutathione (Acros). The GST epitope tag was removed by thrombin cleavage. Proteins were dialyzed exhaustively to remove glutathione and were further purified by cation-exchange and size-exclusion chromatography, which yielded >95% pure samples in monomeric form. SeMet-incorporated PF\textsubscript{14}\_0633 was expressed in \textit{Escherichia coli} grown in M9 minimal media containing SeMet (Acros) and purified as above, except that 2 mM fresh dithiothreitol (DTT) was included in buffers. Protein concentrations were determined by measurement of the absorbance at 280 nm.

#### Crystallization and structure determination of DNA-bound PF\textsubscript{14}\_0633 AP2 domain

PF\textsubscript{14}\_0633 AP2 domain (15 mg/mL in GST lysis buffer) was mixed at a 1:1.2 molar ratio (protein:dsDNA) with a palindromic dsDNA (5’-TGCATGCA-3’, annealed in 10 mM Tris, pH 8.0, and 50 mM NaCl). Equivalent volumes of the complex and mother liquor (10 mM CoCl\textsubscript{2}, 100 mM sodium acetate, pH 5.1, 0.5–1.25 M L,6-hexanediol, and 10–20% v/v ethylene glycol) were mixed, and crystals of the complex formed in 1–3 days by hanging drop vapor diffusion. Crystals were cryoprotected in mother liquor adjusted to 25% ethylene glycol prior to being flash-frozen in liquid nitrogen. Diffraction data were collected at the Advanced Photon Source at Argonne National Laboratory (Argonne, IL) at the LS-CAT 21-ID-D beamline (Table 1). Data were indexed and scaled with HKL2000,\textsuperscript{28} and an experimental electron density map of the SeMet-incorporated AP2 domain of PF\textsubscript{14}\_0633 bound to dsDNA was generated by single-wavelength anomalous dispersion phasing using SOLVE and solvent flattening using DM.\textsuperscript{29} A partial model of the complex was generated and then used as a molecular replacement search model to determine the structure of the native complex using Phaser.\textsuperscript{30} The native structure model was improved by rounds of manual building using Coot\textsuperscript{31} and refinement with REFMAC.\textsuperscript{32}

#### Electrophoretic mobility shift assay

Oligonucleotides containing a 28-bp fragment of the genomic sequence from \textit{P. falciparum} located upstream of \textit{pf}0540\_wt that contains the PF\textsubscript{14}\_0633 AP2 consensus binding sequence (GCA\textsubscript{7}TGC) were synthesized by the University of Wisconsin Biotechnology Center to create a DNA substrate (“wt”). A mutant substrate with the central AT base pairs of the consensus binding site changed to GC (GCGGCC, “mut”) was also produced. Both substrates were inserted into the Stul site of the pCR-Blunt vector (Invitrogen) to confirm their sequence identity and to produce 122-bp DNA substrates by restriction with HindIII and XhoI and polyacrylamide gel electrophoretic
purification (probe sequences listed in Supplementary Table 3). One picomole of each duplex DNA probe was end-labeled by T4 Polynucleotide Kinase (New England Biolabs) with \(^{32}\)P-ATP. Ten femtomoles of end-labeled probe was incubated with 0.1–50 \(\mu\)M concentrations of the PF14_0633 AP2 domain (or a variant) in 20 \(\mu\)l GST lysis buffer supplemented with 0.1 mg/ml bovine serum albumin and 5 \(\mu\)g/ml poly(dl-dC) (Sigma) for 30 min at room temperature. Samples were electrophoresed through a 12% polyacrylamide gel and exposed to a phosphor screen (Molecular Dynamics). Signals were visualized by a Storm 860 PhosphorImager, and band intensities were quantified by ImageQuant (Molecular Dynamics).

**Protein binding microarrays**

PBM experiments using the PF14_0633 AP2 variant domains were performed as described previously.\(^33\) A minimum of two replicate experiments were performed for each variant.

**Cysteine cross-linking**

The PF14_0633 AP2 domain (or triple variant: Cys76Ala, Ala86Cys, Asn118Cys) was treated with 2 mM DTT to reduce disulfide bonds and subsequently dialyzed exhaustively against GST lysis buffer (pH 7.4) to remove the DTT. Individual proteins (60 \(\mu\)l) were incubated in the absence of dsDNA or in the presence of 75 \(\mu\)M cognate dsDNA (5’-TGATGCAGCATG-3’) or non-cognate dsDNA (5’-TGATGCAGCATG-3’) for 30 min at 4 \(^\circ\)C. Dimethylformamide (0.5 \(\mu\)l) or 20 mM BM(PEG)\(_3\) (a 17.8-Å homobifunctional maleimide cross-linker, Thermo Scientific) dissolved in dimethylformamide was added to samples for 30 min at 4 \(^\circ\)C. Samples were electrophoresed on reducing or nonreducing 20% SDS-PAGE gels and visualized by Coomassie Blue. Band intensities were analyzed by ImageQuant.

Calibrated size-exclusion chromatography

PF14_0633 AP2 domain (400 \(\mu\)M) was incubated with 500 \(\mu\)M cognate dsDNA, with non-cognate dsDNA (sequences described above), or without DNA for 1–2 h in GST lysis buffer at 4 \(^\circ\)C. Samples were applied to a GE Healthcare S-100 HR column, and elution volumes were determined by detection of UV absorbance peaks. Average molecular weights of each peak were determined by comparing the elution volumes to calibrated standard markers (Bio-Rad Gel Filtration Standard) run just prior to and following experimental runs. Peak fractions were also analyzed by reducing and nonreducing SDS-PAGE as above.

Biological identification of multiple binding sites with transcriptional upstream regions of *P. falciparum*

ScanACE\(^34\) and the PBM-derived PF14_0633 wild-type position weight matrix were used to scan the 2-kb upstream regions of the *P. falciparum* sporozoite-expressed genes (listed in Ref. 24). The 2-kb upstream regions of the 30 sporozoite genes were extracted from the *Plasmodium* genome resource PlasmoDB.org v5.5.\(^{35}\) Scoring cutoffs were set to capture two standard deviations below the mean score in the alignment file.

**Accession numbers**

The X-ray crystallographic structure and structure factors have been deposited in the Protein Data Bank and have been assigned the identifier 3IGM.

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**Supplementary Data**

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.jmb.2009.11.004

**References**


