Mode of DNA binding with γ-butyrolactone receptor protein CprB from *Streptomyces coelicolor* revealed by site-specific fluorescence dynamics

Anwesha Biswas a, Satya Narayan b, Mamata V. Kallianpur b, G. Krishnamoorthy b,*, Ruchi Anand a, *

a Department of Chemistry, Indian Institute of Technology Bombay, Mumbai, Maharashtra 400076, India

b Department of Chemical Sciences, Tata Institute of Fundamental Research, Mumbai, Maharashtra 400005, India

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**A B S T R A C T**

**Background:** The γ-butyrolactone (GBL) binding transcription factors in *Streptomyces* species are known for their involvement in quorum sensing where they control the expression of various genes initiating secondary metabolic pathways. The structurally characterized member of this family CprB from *Streptomyces coelicolor* had earlier been demonstrated to bind a multitude of sequences containing a specific binding signature. Though structural breakthrough has been obtained for its complex with a consensus DNA sequence there is, however, a dearth of information regarding the overall and site specific dynamics of protein–DNA interaction.

**Methods:** To delineate the effect of CprB on the bound DNA, changes in motional dynamics of the fluorescent probe 2-aminopurine were monitored at three conserved base positions (5th, 12th and 23rd) for two DNA sequences: the consensus and the biologically relevant cognate element, on complex formation.

**Results:** The changes in lifetime and generalized order parameter revealed a similarity in the binding pattern of the protein to both sequences with greater dynamic restriction at the end positions, 5th and 23rd, as compared to the middle 12th position. Also differences within this pattern demonstrated the influence of even small changes in sequence on protein interactions.

**Conclusions:** Here the study of motional dynamics was instrumental in establishing a structural footprint for the cognate DNA sequence and explaining the dynamics for the consensus DNA from structural correspondence. General significance: Motional dynamics can be a powerful tool to efficiently study the mode of DNA binding to proteins that interact differentially with a plethora of DNA sequences, even in the absence of structural breakthrough.

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1. Introduction

The system controlling protein expression in a cell relies on a class of proteins known as transcription factors [1–4]. These proteins recognize their cognate DNA via different DNA binding motifs, like the zinc finger [5,6] and helix-turn-helix (HTH) motifs [7–9]. In most cases binding to DNA induces a conformational change that primes them for either transcription repression or activation [2,3]. Many transcription factors are pleiotropic in nature and bind to a multitude of DNA sequences containing conserved elements [10,11]. This allows them to exert a finer level of control over DNA by allowing gradations of expression and at times they also serve as autoregulators of their own expression [12,13]. In *Streptomyces* the onset of secondary metabolic pathways like morphological differentiation, biofilm formation, including the crucial function of antibiotic production is under the control of transcription factors from the GBL binding family [10,13–16]. GBLs are small diffusible molecules that control the quorum sensing process, which links the cell density to downstream gene regulation in *Streptomyces* genus [17–19].

Structural breakthrough in the GBL binding family of transcriptional factors has been achieved only for the protein CprB from *Streptomyces coelicolor* [15,16]. The structure confirmed the presence of the two domains, one associated with DNA-binding and the other responsible for signal detection through the formation of ligand–protein complex. The DNA binding domain is constituted of a HTH motif and is highly conserved among this family of proteins (Fig. 1a). By virtue of sequence and structural similarity it has been classified into the generic superfamily of tetracycline transcriptional regulator (TetR) family of repressors [20]. The TetR family of proteins consists of several proteins most of which bind to antibiotics and serve as regulators of the efflux pump pathway. Very recently the structure of CprB along with a consensus DNA (CS sequence; Fig. 1b) was solved [13,21]. The structure revealed that CprB binds to this semi-palindromic sequence as a dimer of dimer. This binding event induces a pendulum like motion that results in reorganization of various interactions across the individual dimeric interfaces. Comparison of the X-ray structure revealed that the DNA
bound form of CprB was structurally most similar to the multidrug efflux regulator QacR protein from the antibiotic resistant strain Staphylococcus aureus [22], thereby indicating the possibility of an analogous role in S. coelicolor.

In a recent study, CprB was also shown to bind a host of promoter sequences in S. coelicolor with varying affinity [13]. Most of these sequences influence both GBL and antibiotic production in the organism. Furthermore, based on the consensus sequence the potential cognate DNA of CprB was identified to be a 27 base pair sequence (OPB sequence, Fig. 1b) located upstream of its own gene. The study revealed that CprB, like many other TetR family proteins, apart from controlling the expression of a non-invasive efflux regulator QacR protein from the antibiotic resistant strain Staphylococcus aureus [22], thereby indicating the possibility of an analogous role in S. coelicolor.

In the present study we analyze the extent of perturbation at three base positions, 5th, 12th and 23rd, of both the consensus (CS_5, CS_12 and CS_23) and cognate (OPB_5, OPB_12 and OPB_23) sequences upon protein binding (Fig. 1c). This was achieved by substituting 2-AP in place of the conserved adenine at these positions. The fluorescence lifetimes and anisotropy decay parameters of 2-AP were measured for the DNA and DNA in complex with CprB. Together, the information generated from 2-AP dynamics at the three conserved base positions and the available crystal structure were utilized to understand the altered motional dynamics of DNA in the presence of the protein and generate a structural footprint of DNA binding for the cognate OPB sequence.

2. Material and methods

2.1. Purification of proteins

The clone of the native protein CprB in pET28a(+) expression vector was kindly provided to us by Ryo Natsume (Japan Biological Informatics Consortium (JBIC), Tokyo, Japan). The procedures for cloning, expression and purification have been previously reported [40]. The proteins were finally desalted into buffer containing 50 mM sodium phosphate pH 7 and 180 mM NaCl and used for fluorescence studies.

2.2. DNA synthesis

The oligonucleotides with 2-AP modification and their complementary sequences were synthesized in using a MerMade 4 automated synthesizer (BioAutomation, Plano, Texas, USA). All strands were synthesized at 1 μmole scale with appropriate Controlled Pore Glass beads (Proligo Reagents, Hamburg, Germany) used as 3’ solid support. 2-Aminopurine phosphoramidite (Glen Research Corporation, Virginia, USA) was added for the incorporation of 2-AP in a sequence. The DNA sequences synthesized were purified by denaturing PAGE (20%, 7 M urea) using standard protocols. All the DNA sequences with the 2-AP modification have been characterized by MALDI-TOF-MS (Supporting information, Table S1).

The oligonucleotides were quantified by employing spectrophotometer (GE, GeneQuant 1300, WI, USA) at 260 nm. The complementary strands of the modified (2-AP incorporated) DNA were taken in 20% excess, so as to ensure complete annealing of the fluorophore incorporated DNA. The DNA sequences were annealed in the presence of 1 X annealing buffer (5 mM Tris–HCl, pH 7.5, 15 mM NaCl, 0.1 mM EDTA). The mixture was subjected to a temperature of 95 °C for 5 min by placing in a water bath which was then allowed to cool slowly to room temperature. The samples were stored at −20 °C.

2.3. Electrophoretic mobility shift assay

The binding affinity studies of CprB to the CS and OPB sequences were carried out with the help of 5’-end radio labeled oligonucleotides.
Protein concentrations from 6 μM to 90 nM (in buffer 50 mM sodium phosphate pH 7, 180 mM NaCl) obtained by 2-fold serial dilution, were incubated with approximately 1 nM of radiolabeled and annealed DNA. The protocol that was followed for the assay has been reported previously [13].

2.4. Time-resolved fluorescence

Time resolved fluorescence decay and anisotropy decay measurements were performed using a Rhodamine 6G dye laser that generates pulses of 1 ps width. The dye laser was pumped by a passively mode-locked frequency doubled Nd:YAG laser (Vanguard, Spectra Physics). The oligonucleotide samples containing 2-AP (with or without the protein) were excited at 310 nm which was the second harmonic output of angle-tuned KDP crystal. The curves for fluorescence decay were obtained from a time correlated single-photon counting setup which was coupled to a microchannel plate photomultiplier (model 2809U, Hamamatsu Corp.). The instrument response function (IRF) was obtained at 310 nm using a very dilute colloidal suspension of dried non-dairy coffee whitenner. The half-width of the IRF was ~40 ps. The fluorescence emission for the excited samples was collected through a 345 nm cut-off filter followed by a monochromator at 370 nm with a collection bandwidth of 3 nm. The number of counts in the peak channel was ~10,000 for most cases except where the decay was very fast, for which the counts were kept at ~25,000. The fluorescence emission for lifetime measurements was monitored at 54.7° (magic angle) to avoid contribution from anisotropy decay. The emission was collected at directions, parallel and perpendicular to that of the incident polarized light, for the time-resolved anisotropy measurements. For more details see elsewhere [27,28,35,41].

Since one of the fluorescence lifetimes was ~40 ps (see Results and discussion) the decay kinetics were also recorded by using a Streak Camera set-up (Model SC-10, Optronics, Kehl, Germany). Half width of the IRF was ~5 ps. In the Streak Camera experiments only one ultrafast lifetime was obtained (τ1). This is may be have been due to the instrument response time of Streak Camera being longer than that of the down-conversion set-up. The samples consisted of 10 μM oligonucleotides, with and without the protein at 40 μM concentration. The complex was incubated at room temperature for 10 min before data collection. The buffer consisted of 50 mM Phosphate and 180 mM NaCl, pH 7.

2.5. Data analysis

The analysis of the data of fluorescence decay for the lifetime measurements were performed using FluoFit Pro Version 4.4 (PicoQuant) by a non-linear least-squares iterative reconvolution method based on the Levenberg–Marquardt algorithm [42] and expressed as a sum of exponentials with the equation:

\[ I(t) = \sum \alpha_i \exp(-t/\tau_i) \]  

where \( \alpha_i \) represents the amplitude of the ith component associated with fluorescence lifetime \( \tau_i \) such that \( \sum \alpha_i = 1 \). \( \Sigma \alpha_i \tau_i \) gives us the mean lifetime \( \tau_m \) of the system which gives knowledge about the average fluorescence yield of the system. The support plane analysis was performed to determine the confidence intervals for each parameter [43].

2.5.1. Analysis of fluorescence anisotropy decay kinetics

The time-resolved anisotropy decay curves were derived from the experimentally obtained \( I_\parallel(t) \) and \( I_\perp(t) \) with the equation:

\[ r(t) = \frac{I_\parallel(t) - G(\lambda)I_\perp(t)}{I_\parallel(t) + 2G(\lambda)I_\perp(t)} \]  

where \( r(t) \) is the time dependent anisotropy, \( I_\parallel(t) \) and \( I_\perp(t) \) are the fluorescence intensity collected with the emission polarizer at 0° (parallel) and 90° (perpendicular), with respect to excitation polarizer, respectively, and \( G(\lambda) \) is the geometry factor at the wavelength of emission \( \lambda \). The value of \( G(\lambda) \) for the optics for measuring emission was calculated independently using 50 μM solution of 2-AP.

\[ I_\parallel(t) = I(t)[1 + 2r(t)]/3 \] \[ I_\perp(t) = I(t)[1 - r(t)]/3 \]

The fluorescence anisotropy decays were fitted using FluoFit into a bi-exponential model as follows:

\[ r(t) = r_0[β_1 \exp(-t/τ_1) + β_2 \exp(-t/τ_2)] \]  

where \( r_0 \) is the anisotropy in the absence of any rotational diffusion and \( β_i \) is the amplitude associated with the ith rotational correlation times \( τ_i \) such that \( \sum β_i = 1 \). This model assumes that the sample contains a population having uniform motional dynamics properties with each species associated with two rotational correlation times [35]. In general, fluorescence anisotropy decay kinetics can be analyzed by one of the following two models [44]: (A) Multiple populations of the fluorophore (revealed by multiple exponentials in fluorescence intensity decay) have the same motional dynamics and (B) Multiple populations of fluorophore with each population having their own motional dynamics. Model B has been used in situations where more than one population is encountered [45–47]. In such situations anisotropy decay kinetics has complex shapes such as ‘flip-and-rise’ which has been fitted by model B. In contrast, multiple exponentials arising from microstates of macromolecules have been assumed to have similar motional dynamics (Model A) since multiple exponentials can have their origin in several ways including excited state reactions etc. Hence in the present work we have analyzed our results based on model A. Further justification for the use of this model is given later.

The \( r_0 \) value was estimated by a separate experiment with a sample in 70% glycerol, which was 0.31. The goodness of fits was assessed from the reduced \( χ^2 \) values (1.2–1.7) as well as from the randomness of the residuals. The support plane analysis was again performed to determine the confidence intervals for each parameter. For more details see elsewhere [27,28,35,41]. The results of the anisotropy decay analysis have been analyzed in terms of the change in \( S^2 \) for the probe, before and after the binding of protein, where \( S \) is the generalized order parameter [48]. \( S \) provides a measure of the restriction in the local motion of the probe and is given by the following equation [25,49]:

\[ S^2 = β_2/(β_1 + β_2) \]  

2.6. Maximum entropy method of analysis

The distribution of 2-AP conformers with different lifetimes, in the duplex DNA as well as in DNA–protein complex was also analyzed by the Maximum Entropy Method (MEM) of analysis [50,51]. It is a model free approach which initially assumes the fluorescence decay as a distribution of discrete lifetime values that are equally spaced in the \( \log(τ) \) space within a range which is determined by the nature of the particular fluorophore. A range of 10 ps to 20 ns was given for 2-AP. The method initially gives equal amplitude to all the lifetime values and with each iteration the distribution changes towards obtaining minimization of \( χ^2 \) and maximization of the Shannon–Jaynes entropy function given by:

\[ S = \sum \alpha_i \log \alpha_i \]  

where \( \alpha_i \) represents the amplitude of the ith lifetime.
3. Results and discussion

3.1. DNA binding studies of CprB with CS and OPB DNA

The first sequence identified to bind CprB was the CS sequence [21] and subsequently this sequence was used to find the OPB sequence [13]. While CS sequence had been selectively pulled out from a random pool of chemically synthesized DNA sequences, the OPB sequence appears in the S. coelicolor genome and is therefore more biologically significant. The overall binding strength of CprB to the two sequences was estimated from electrophoretic mobility shift assays (Fig. 2). For the CS and OPB sequences an average of three data points was used to estimate the dissociation constant.

CprB exhibited similar levels of affinity for both the sequences (Kd for CS 1.26 ± 0.63 μM; Kd for OPB, 1.53 ± 0.96 μM). This could be attributed to the fact that both the sequences exhibit 52% sequence similarity and hence the binding to the two sequences CS and OPB is possibly arising from certain specific interactions among the conserved bases. However, this is not always the case as in an earlier study CprB had been shown to exhibit gradation in binding affinity with sequences of the TetR family of proteins to which the protein bears similarity in the DNA binding motif [13]. The assays performed here only give the overall binding affinity of the DNA–protein system and does not shed any light on the mode of binding or site specific dynamics which is an important aspect that needs investigation. To confirm whether the substitution of adenine by 2-AP had caused a change in the affinity of the protein towards the sequences, EMSA was also performed with all the modified DNA sequences (Fig. S1). The modified DNA sequences exhibited affinities similar to the original sequence.

3.2. Fluorescence intensity decay kinetics of 2-AP in OPB and CS DNA sequences

Free 2-AP has a quantum yield of 0.68 in an aqueous solution (pH 7) [52] and a single lifetime of ~11 ns [53]. However, when the fluorophore is incorporated in an oligonucleotide sequence, a very significant reduction in its quantum yield is often observed as a result of various interactions of 2-AP with its neighboring bases [23,24]. The degree of quenching of 2-AP is highly susceptible to perturbations in the local environment of the fluorophore in the sequence [23,24,53,54]. Moreover, 2-AP when present within a DNA displays a multi-exponential decay representative of a heterogeneous population of fluorophores with time constants ranging from ~10 ps to ~10 ns [23,24,35,41]. The population heterogeneity has been explained in the existence of an array of microstates, with different degrees of DNA stacking.

The shortest lifetime component that is representative of base-pair formation in a double stranded DNA, is also related to the stronger stacking interactions of 2-AP with the neighboring bases. However, there could be other models such as excited state electron transfer to nearby guanines [23,24,54–56] resulting in multiple exponentials.

Fluorescence decay kinetics for all the DNA sequences were best represented as a sum of four components with distinct lifetimes (1). The decay was also analyzed by the MEM that presents an impartial representation of the four lifetimes.

As seen in Fig. 3, four distinguishable peaks corresponding to the lifetimes obtained from the discrete analysis (Table 1) indicate the validity of four lifetimes inferred from MEM. Decay parameters listed in Table 1 show that the shortest lifetime component (τ1) is the major component in all the cases. Both the CS and OPB duplex DNA exhibit a very short value of mean lifetime τm due to the large amplitude (α1) of the shortest lifetime species. Compared to the τm of single stranded DNA (Supporting information, Table S2), 2-AP in duplex DNA shows significant decrease in the value of τm except in the case of CS_23. The decrease is due to (i) the enhancement of stacking interaction brought about in the duplex DNA due to base pairing as compared to the less ordered single stranded DNA [23,24,55,56] and/or (ii) electron transfer to bases (mainly guanine) either adjacent to 2-AP or in the opposite strand [55]. Significantly less reduction, observed on going from single stranded to double stranded CS_23 (which does not have any guanine either adjacent or in the opposite strand), is a clear indicator that the very short lifetime in other duplex DNA is primarily due to electron transfer quenching by nearby guanine. Addition of CprB caused significant changes in the fluorescence decay kinetics of 2-AP in all the DNA sequences. The main changes are the increase in the value of τ1, decrease in the value of α1 and increase in the amplitude (α4) of the longest lifetime component (τ4) that result in the significant increase in the value of τm in all the cases (Table 1). The increase in the mean lifetime was a combined result of an overall increase in all the individual lifetimes as well as a shift of the population of fluorophores. For example, a comparison across the OPB DNA shows an increase in mean lifetime by 5 and 3.6 times at the 5th and 23rd positions as compared to a slightly smaller increase by 2.5 at the 12th position. This increase in the mean lifetime can be due to local perturbation caused at specific positions due to binding by CprB.

3.3. Motional dynamics from fluorescence anisotropy decay

Unlike fluorescence lifetime which is modulated by a variety of causes, fluorescence anisotropy kinetics is dominantly controlled by motional dynamics of the fluorophore and hence could be a robust read-out of protein binding to DNA. Fluorescence anisotropy decays of 2-AP in the DNA and DNA–protein complexes represented in Fig. 4 show position dependent motional dynamics of 2-AP in both the free OPB and CS sequences.

The decays were fitted satisfactorily to a sum of two exponentials. The shorter correlation time (ϕ1), represents the local motion of the 2-AP with respect to the DNA backbone while the longer one (ϕ2) represents the combined effect of both segmental motion within the DNA strand as well as the or the overall tumbling motion of whole strand. Most importantly, from the analysis of the anisotropy decay, the spatial
restrictions felt by the probe within the DNA can be comprehended from the value of $S^2$ as seen in Table 2.

Several observations could be made on the position dependence of motional dynamics of 2-AP in the two DNA sequences studied: (i) local motion represented by $\phi_1$, is significantly slower in CS DNA when compared to OPB DNA and (ii) 2-AP at the CS_23 position has significantly slower dynamics when compared to other positions in the CS DNA. Position-dependence of motional dynamics of 2-AP in duplex DNA and especially the unusual slow dynamics of 2-AP in between two thy-DNA. Position-dependence of motional dynamics of 2-AP in duplex DNA and especially the unusual slow dynamics of 2-AP in between two thy-DNA. Position-dependence of motional dynamics of 2-AP in duplex DNA and especially the unusual slow dynamics of 2-AP in between two thy-DNA. Position-dependence of motional dynamics of 2-AP in duplex DNA and especially the unusual slow dynamics of 2-AP in between two thy-DNA. Position-dependence of motional dynamics of 2-AP in duplex DNA and especially the unusual slow dynamics of 2-AP in between two thy-DNA. Position-dependence of motional dynamics of 2-AP in duplex DNA and especially the unusual slow dynamics of 2-AP in between two thy-DNA.

Comparison of the CprB binding induced increase in the value of $S^2$ indicated that the level of restriction imposed by the protein varies across both the DNA sequences (Fig. 5). It was observed that in the CS series of DNA, there was a large increase in $S^2$ by approximately 0.55 for the 2-AP at the 5th position and also an increase of 0.43 for the 23rd position. The 12th position showed the minimum increase of 0.32. It is evident from the figure that the protein poses greater restrictions to the movement of the 2-AP, at either end of the oligomer than in the center. For the OPB DNA series as well there is a clear indication of restriction imposed by complex formation to the movement of 2-AP.

The value of $S^2$ increases by 0.35, 0.23 and 0.40 for OPB_5, OPB_12 and OPB_23 respectively. Comparison of the results of anisotropy decay analysis between the two sequences reveals a remarkable similarity in the manner the bases at the equivalent positions in the DNA are restricted by the protein (Fig. 5). Even though the amount of restriction at the equivalent positions varies, there is high resemblance in the trend by which the restriction is imposed at the positions. The dynamics in both the DNA sequences seem to be largely affected by protein binding towards the end positions and much less in the central position. However, it was observed that the CS sequence exhibited an overall larger restriction in dynamics on complex formation than the OPB sequence. The reason could be attributed to the difference in the inherent nature of the two sequences. The OPB sequence is comparatively more GC rich than the CS sequence and probably this is the reason it exhibits higher degree of base stacking even in the absence of the protein. Protein binding therefore seems to induce a greater degree of structure in the CS sequence as is evident from the greater overall restriction observed for this sequence as compared to the OPB sequence.

In order to validate the observations from the site-specific studies performed with the CprB-binding sequences, a control sequence was also analyzed. The control sequence consisted of the probe incorporated in a non-interacting position created by a poly-A extension at the 3' end of the OPB sequence (Control sequence, C3: 5'-AGGCAGGCAGGCGGCACGGT CTTTGAAGTTAAAAA*A-3'; A* represents 2-AP). The successful binding of the overall sequence was evident from the increase in value of the longer correlation time $\phi_2$, which indicates slower global motion of the probe due to increase in its bulkiness. As observed from the generalized order parameter, the effect on the restriction to local motion of the probe in the control sequence was however negligible ($\Delta S^2$ = 0.03, Table 2, Supporting information Fig. S2). This reveals the specific nature of the CprB binding preferentially to the signature sequence and confirms that the changes in $\Delta S^2$ seen earlier are due to specific interaction of the probe with CprB.

Quenching of fluorescence of 2-AP by acrylamide could also provide a site-specific map of binding of CprB to the DNA sequences. Bimolecular quenching constant $k_q$ estimated from Stern-Volmer plots associated with quenching by acrylamide [44] were found to be in the range of $10^{10}$ M$^{-1}$ s$^{-1}$ for all the positions in both the DNA and in the range of $10^9$ M$^{-1}$ s$^{-1}$ for the CprB–DNA complexes (data not shown). This decrease is insensitive to any particular position along the length of the DNA sequence and only provides a global signature of protein binding. Therefore, as mentioned earlier, we believe fluorescence anisotropy as a more reliable technique to study site specific changes.

3.4. Structural correspondence of CprB–CS complex to the observed alteration in base dynamics

In order to get an in-depth understanding of how the motional dynamics at a specific position corresponds to the local structure, we analyzed the relationship between the changes in base motional restriction upon protein binding at each modified position with the interactions observed in the X-ray structure.

From the structure of CprB with a truncated CS sequence, it was observed that the adenine at the 5th position interacts more extensively with the protein than the adenine moieties present at the 12th and 23rd position (Fig. 6a). The region around the 5th base is well anchored into the major groove of the DNA making extensive interactions with residues residing in the recognition helix α3. The phosphate backbone
Table 1
Time resolved fluorescence decay parameters for 2-AP in duplex DNA and in DNA–protein complex with confidence intervals for all parameters calculated using the support plane method.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Fluorescence lifetime ($\tau_i$ ns), amplitude ($\alpha_i$)</th>
<th>Mean lifetime $\tau_m$ (ns)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$\tau_1$ $\alpha_1$ $\tau_2$ $\alpha_2$ $\tau_3$ $\alpha_3$ $\tau_4$ $\alpha_4$</td>
<td></td>
</tr>
<tr>
<td>CS_5</td>
<td>0.021 (−0.019 + 0.002) 0.98 (−0.05 + 0.04) 0.22 (−0.04 + 0.05) 0.012 (−0.001 + 0.026) 1.09 (−0.16 + 0.55) 0.005 (−0.001 + 0.002) 5.32 (−0.11 + 1.77) 0.003 (−0.001 + 0.001) 0.042</td>
<td></td>
</tr>
<tr>
<td>CS_5_CprB</td>
<td>0.029 (−0.028 + 0.004) 0.93 (−0.16 + 0.70) 0.29 (−0.18 + 0.19) 0.027 (−0.009 + 0.040) 1.93 (−0.60 + 1.89) 0.017 (−0.005 + 0.011) 6.16 (−0.53 + 9.43) 0.028 (−0.011 + 0.005) 0.24</td>
<td></td>
</tr>
<tr>
<td>CS_12</td>
<td>0.020 (−0.019 + 0.010) 0.96 (−0.60 + 0.24) 0.21 (−0.05 + 0.082) 0.022 (−0.004 + 0.006) 0.90 (−0.38 + 0.38) 0.009 (−0.002 + 0.006) 5.26 (−0.099 + 2.45) 0.004 (−0.001 + 0.001) 0.053</td>
<td></td>
</tr>
<tr>
<td>CS_12_CprB</td>
<td>0.026 (−0.004 + 0.006) 0.93 (−0.32 + 0.08) 0.56 (−0.08 + 0.09) 0.036 (−0.006 + 0.006) 3.72 (−2.73 + 0.87) 0.028 (−0.006 + 0.004) 9.90 (−4.15 + 19.2) 0.010 (−0.006 + 0.009) 0.24</td>
<td></td>
</tr>
<tr>
<td>CS_23</td>
<td>0.057 (−0.029 + 0.028) 0.39 (−0.39 + 0.23) 0.34 (−0.09 + 0.06) 0.49 (−0.18 + 0.04) 0.65 (−0.17 + 0.47) 0.12 (−0.11 + 0.01) 4.93 (−1.10 + 13.3) 0.004 (−0.002 + 0.001) 0.29</td>
<td></td>
</tr>
<tr>
<td>CS_23_CprB</td>
<td>0.047 (−0.021 + 0.032) 0.48 (−0.24 + 0.24) 0.33 (−0.05 + 0.06) 0.44 (−0.09 + 0.05) 0.96 (−0.24 + 1.19) 0.058 (−0.041 + 0.048) 5.44 (−0.45 + 1.38) 0.020 (−0.006 + 0.002) 0.33</td>
<td></td>
</tr>
<tr>
<td>OPB_5</td>
<td>0.024 (−0.022 + 0.004) 0.93 (−0.19 + 0.65) 0.31 (−0.13 + 0.21) 0.033 (−0.007 + 0.016) 1.77 (−0.24 + 0.58) 0.028 (0.008 + 0.012) 6.90 (−0.71 + 1.54) 0.010 (−0.003 + 0.005) 0.15</td>
<td></td>
</tr>
<tr>
<td>OPB_5_CprB</td>
<td>0.031 (−0.010 + 0.007) 0.77 (−0.34 + 0.16) 0.32 (−0.16 + 0.23) 0.064 (−0.016 + 0.037) 2.28 (−0.48 + 0.52) 0.087 (−0.019 + 0.017) 6.50 (−0.64 + 0.80) 0.079 (−0.023 + 0.017) 0.76</td>
<td></td>
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<tr>
<td>OPB_12</td>
<td>0.023 (−0.009 + 0.004) 0.98 (−0.37 + 0.39) 0.35 (−0.31 + 5.87) 0.005 (−0.005 + 0.036) 1.37 (−1.25 + 4.52) 0.008 (−0.007 + 0.003) 5.40 (−1.15 + 16.6) 0.005 (−0.004 + 0.002) 0.62</td>
<td></td>
</tr>
<tr>
<td>OPB_12_CprB</td>
<td>0.026 (−0.005 + 0.005) 0.96 (−0.31 + 0.51) 0.33 (−0.14 + 0.23) 0.015 (−0.004 + 0.008) 2.15 (−0.58 + 1.02) 0.041 (−0.004 + 0.005) 6.18 (−0.69 + 2.59) 0.015 (−0.006 + 0.003) 0.15</td>
<td></td>
</tr>
<tr>
<td>OPB_23</td>
<td>0.037 (−0.091 + 0.006) 0.84 (−0.21 + 0.51) 0.61 (−0.50 + 0.05) 0.079 (−0.049 + 0.064) 1.95 (−0.10 + 1.22) 0.051 (−0.011 + 0.051) 5.74 (−0.60 + 1.71) 0.028 (−0.016 + 0.016) 0.34</td>
<td></td>
</tr>
<tr>
<td>OPB_23_CprB</td>
<td>0.044 (−0.041 + 0.031) 0.62 (−0.30 + 0.31) 0.46 (−0.44 + 0.25) 0.10 (−0.05 + 0.12) 2.09 (−0.39 + 0.92) 0.13 (−0.05 + 0.08) 6.20 (−0.38 + 1.10) 0.140 (−0.03 + 0.03) 1.21</td>
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</table>
of the adjacent bases anchors this adenine group such that it makes several (Fig. 6b) hydrophobic interactions with conserved residues F48, A45 and G44 both with the purine ring as well as the deoxyribose moiety. There is also a hydrogen bonding interaction between the backbone phosphate group (joining the adenine to the neighboring thymine) and T42. Based on structural data it can be concluded that the cumulative effect of these interactions should result in restricted motion upon DNA binding at the 5th position. The large dynamic restriction observed on the 2-AP substituted at this position corroborates these conclusions.

Similarly, from the analysis of the structure it was predicted that the 12th position should have the least restriction as this region of the DNA is relatively far from the protein. It was observed that the adenine base at the 12th position (Fig. 6c) showed hydrophobic interactions of its purine ring with the long aliphatic side chain of the K43 of α3.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Rotational correlation time ($\phi_1$, ns), amplitude ($\beta_1$)</th>
<th>$\phi_2$</th>
<th>$\beta_2$</th>
<th>$S^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>CS_5</td>
<td>0.44 (−0.06 + 0.06) 0.76 (−0.05 + 0.04)</td>
<td>12.4</td>
<td>0.24 (−0.04 + 0.05)</td>
<td>0.24</td>
</tr>
<tr>
<td>CS_5_CprB</td>
<td>0.26 (−0.12 + 0.31) 0.21 (−0.05 + 0.05)</td>
<td>&gt;20.0</td>
<td>0.79 (−0.02 + 0.02)</td>
<td>0.79</td>
</tr>
<tr>
<td>CS_12</td>
<td>0.61 (-0.09 + 0.10) 0.64 (−0.04 + 0.04)</td>
<td>&gt;20.0</td>
<td>0.36 (−0.04 + 0.04)</td>
<td>0.36</td>
</tr>
<tr>
<td>CS_12_CprB</td>
<td>0.13 (−0.03 + 0.03) 0.32 (−0.03 + 0.03)</td>
<td>&gt;20.0</td>
<td>0.68 (−0.01 + 0.01)</td>
<td>0.68</td>
</tr>
<tr>
<td>CS_23</td>
<td>2.15 (−0.06 + 0.13) 0.79 (−0.01 + 0.01)</td>
<td>&gt;20.0</td>
<td>0.21 (−0.01 + 0.01)</td>
<td>0.21</td>
</tr>
<tr>
<td>CS_23_CprB</td>
<td>1.05 (−0.20 + 0.25) 0.36 (−0.04 + 0.04)</td>
<td>&gt;20.0</td>
<td>0.64 (−0.04 + 0.04)</td>
<td>0.64</td>
</tr>
<tr>
<td>OPB_5</td>
<td>0.17 (−0.02 + 0.02) 0.71 (−0.03 + 0.03)</td>
<td>15.9</td>
<td>0.29 (−0.01 + 0.02)</td>
<td>0.29</td>
</tr>
<tr>
<td>OPB_5_CprB</td>
<td>0.06 (−0.02 + 0.02) 0.36 (−0.05 + 0.06)</td>
<td>&gt;20.0</td>
<td>0.64 (−0.01 + 0.01)</td>
<td>0.64</td>
</tr>
<tr>
<td>OPB_12</td>
<td>0.21 (−0.03 + 0.03) 0.67 (−0.04 + 0.04)</td>
<td>&gt;20.0</td>
<td>0.33 (−0.02 + 0.02)</td>
<td>0.33</td>
</tr>
<tr>
<td>OPB_12_CprB</td>
<td>0.080 (−0.01 + 0.02) 0.44 (−0.03 + 0.03)</td>
<td>&gt;20.0</td>
<td>0.56 (−0.01 + 0.01)</td>
<td>0.56</td>
</tr>
<tr>
<td>OPB_23</td>
<td>0.15 (−0.02 + 0.02) 0.75 (−0.02 + 0.02)</td>
<td>10.9</td>
<td>0.25 (−0.01 + 0.01)</td>
<td>0.25</td>
</tr>
<tr>
<td>OPB_23_CprB</td>
<td>0.07 (−0.01 + 0.02) 0.35 (−0.04 + 0.04)</td>
<td>&gt;20.0</td>
<td>0.65 (−0.01 + 0.01)</td>
<td>0.65</td>
</tr>
<tr>
<td>CS_5_F48A</td>
<td>0.10 (−0.06 + 0.07) 0.25 (−0.04 + 0.04)</td>
<td>&gt;20.0</td>
<td>0.75 (−0.01 + 0.01)</td>
<td>0.75</td>
</tr>
<tr>
<td>CS_12_K43A</td>
<td>0.11 (−0.03 + 0.04) 0.37 (−0.04 + 0.04)</td>
<td>&gt;20.0</td>
<td>0.63 (−0.01 + 0.01)</td>
<td>0.63</td>
</tr>
<tr>
<td>Ct</td>
<td>0.36 (−0.05 + 0.05) 0.47 (−0.02 + 0.02)</td>
<td>10.1</td>
<td>0.53 (−0.02 + 0.02)</td>
<td>0.53</td>
</tr>
<tr>
<td>Ct_CprB</td>
<td>0.32 (−0.05 + 0.06) 0.44 (−0.03 + 0.03)</td>
<td>&gt;20.0</td>
<td>0.56 (−0.02 + 0.02)</td>
<td>0.56</td>
</tr>
</tbody>
</table>

* The lower limit in the estimation of $\phi_2$ is caused by the finite lifetime window available for monitoring the decay of fluorescence anisotropy. Ct is a control sequence where 2-AP is far away from the protein binding site.
noted that among the three positions studied, the change in the value of $S^2$ is least for the 12th position thereby implying the lowest restriction at this position upon protein binding. This is not surprising as the 12th position (as apparent from the structure) lies in the region of the DNA which acts like a bridge between the recognition elements and exhibits least interaction with the protein. The 23rd base position however shows a similarity to the 5th position in terms of the change in $S^2$. This was expected as this position also lies in the region of the recognition sequence, which induced fits into the helix-turn-helix motif of CprB (Fig. 6d). The overall dynamic restriction profile of the DNA–protein system consisting of greater restriction at the end positions as compared to the central position therefore corresponded well with the environment of the bases as observed from the structure. Hence, this analysis highlights the fact that the motional dynamics along the length of the DNA, possess an underlying structural footprint. This form of analysis can be employed to predict the structural orientation of the DNA with respect to the protein. Furthermore, the correspondence between the site-specific changes in the generalized order parameter and the known structural features of the CS-DNA-CprB complex gives us credibility in using the model A (see Experimental section) for analysis of fluorescence anisotropy decay kinetics.

Since X-ray crystallography data is not available for the biologically relevant OPB sequence we utilized the comparison of the motional dynamics pattern of CS with the OPB sequence to generate a structural footprint for the cognate OPB sequence. The analysis shows that both the sequences exhibit a similar dynamic signature, with the 5th and the 23rd position exhibiting a higher overall change in the generalized order parameter as compared to the central 12th position. Therefore, it can be concluded that the overall global structure of the OPB sequence is most likely similar to that of CS. However, it appears that the CS sequence has an overall character of being more dynamically restrained by the protein as compared to the OPB sequence. The restraint is enhanced in the 5th and 23rd positions as these positions are in close proximity of the surrounding protein. However, a close look at the values observed at the individual positions revealed that unlike the CS sequence where the motion at the 5th position is most restrained in the OPB sequence it is the 23rd position which exhibits the maximum constrain in motion. Thus the order of restriction in local dynamics is 5th $>$ 23rd $>$ 12th in the CS sequence is now changed to 23rd $>$ 5th $>$ 12th for the OPB sequence. This is most likely because...
both the CS and the OPB sequences exhibit a small degree of difference in the recognition elements that are presented to the protein. For example as compared to CS, OPB sequence is more GC rich. Though there exists a high degree of similarity in the general mode of binding, fine differences could arise from certain preferred interactions of one DNA over the other, by virtue of the dissimilarities in their sequences. Hence, it is noteworthy that the dynamic study can reveal even the fine tuning of interactions by the protein on two quite similar sequences that is otherwise difficult to extract. The results are very significant in the case of proteins such as CprB that are known to bind a multitude of sequences. Though obtaining the crystal structure of complexes with every cognate DNA of the protein is not always feasible, it is however possible to delineate the mode of binding for all the cases using 2AP as a site-specific probe.

3.5. Dynamics study with CprB mutants

To analyze the influence of disrupting interactions at the CprB–DNA interface, conserved residues F48 and K43 whose side chains are involved in contact with the bases were mutated to alanine residue. Both F48 and K43 make hydrophobic interactions with the adenine bases at the 5th and 12th positions respectively. Fluorescence anisotropy results show that in the absence of these individual interactions binding of CS_5 and CS_12 to mutants F48A and K43A resulted in a marginal decrease in the S2 by 0.04 and 0.05 respectively (Table 2, Supporting Information Fig. S3). The decrease indicates the role of these residues in the restriction of the corresponding bases in the sequence.

However, since the decrease in the value of S2 due to the mutations are not drastic it asserts the fact that single point removal of a side chain does not affect the local and overall binding of the protein to the DNA by a large extent and the effect is most likely cumulative. These results are in accord with gel shift experiments performed on an earlier report where single mutations did not exhibit reduced binding affinity, whereas double mutants showed a marked reduction in binding affinity [13]. Our attempts to perform experiments with the double mutants resulted in noisy un-interpretable data mostly because the double mutants were intrinsically unstable and could not be obtained in concentrations required for a proper dynamic study.

4. Conclusions

In this study we investigated the alteration in the dynamics of 2-AP incorporated at certain selected positions in two DNA sequences, CS and OPB, on binding of the protein CprB. The aim was to illustrate how the method can be successfully used to study the mode of binding of the protein to various regions of the DNA site-specifically. In the study, 2-AP in all double stranded DNA exhibited a very short lifetime that was probably caused mainly from electron transfer to bases either adjacent to 2-AP or in the opposite strand. The influence of the neighboring bases was evident from the comparison of the contribution of the shortest lifetime species to the mean lifetime with the 2-AP of CS_23 standing out due to the absence of guanine as an immediate neighbor or in the opposite strand.

In the absence of the structure of CprB with the biologically relevant OPB sequence, a combination of structural and motional dynamics information was employed to generate a structural footprint. In terms of the restriction in the local motion, the protein binding clearly causes an increase in the motional restriction of bases at all positions within the signature binding sequence. The DNA restriction parameters exhibited a similar overall binding pattern, with a larger restriction at the 5th and the 23rd position as compared to the middle 12th position. The results have been accounted for from the interactions observed in the structure of CprB in complex with the consensus sequence. However, due to the differences in base composition of the two sequences local differences in restriction at individual positions were observed. The analysis suggested that unlike in the CS sequence where the base at the 5th position is strongly anchored, in the OPB sequence it is the base at the 23rd position that is preferably stabilized. Therefore, this approach apart from providing global structural correspondence can also give information about fine details of individual interactions. Hence, we can use it to deduce a structural footprint for individual protein–DNA interactions for proteins that bind to a multitude of sequences (such as the pleiotropic regulator CprB).

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Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.bbagen.2015.08.008.

References


