

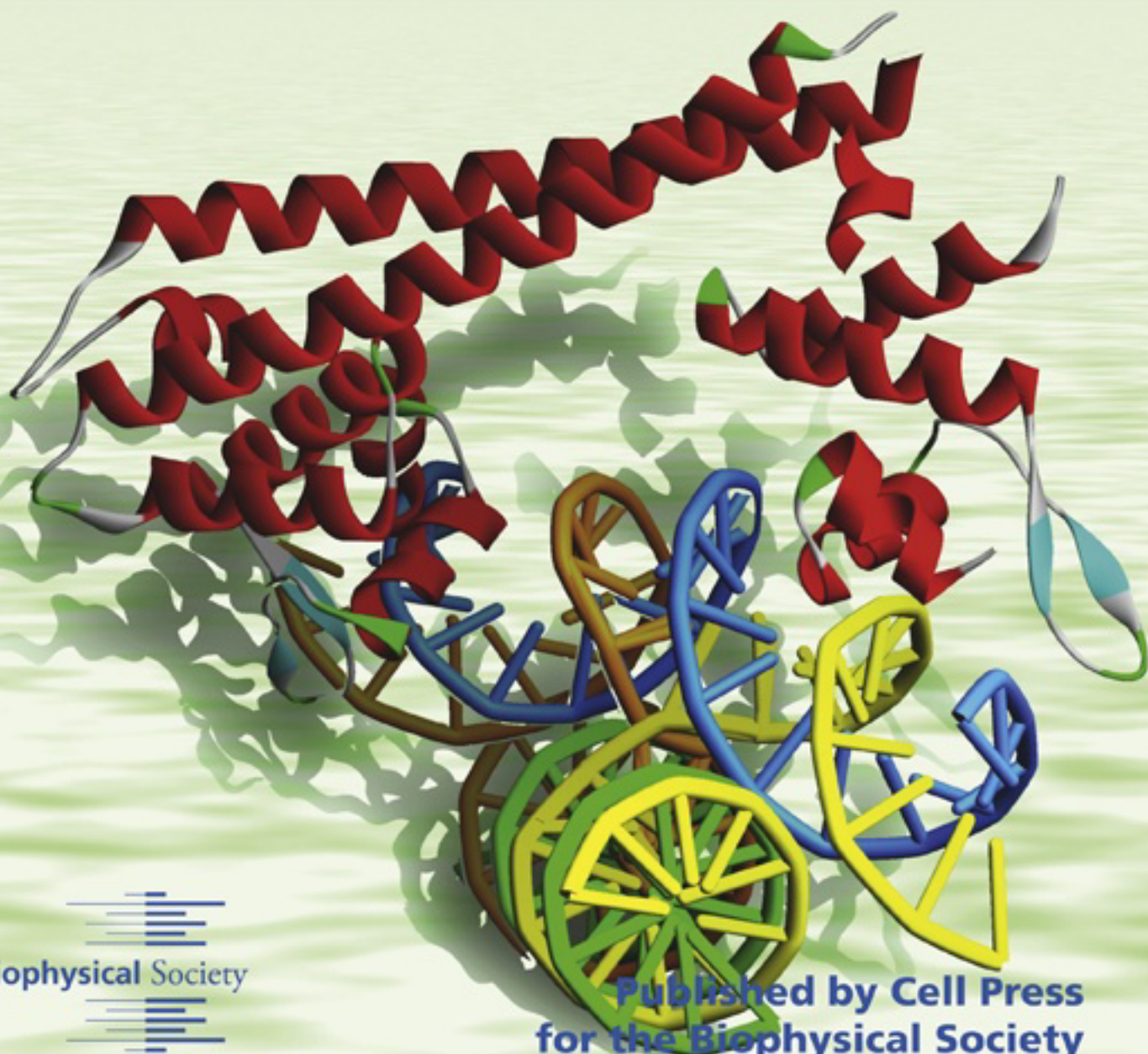
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Single-Molecule Study of Metalloregulator CueR-DNA Interactions Using Engineered Holliday Junctions

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ABSTRACT To maintain normal metal metabolism, bacteria use metal-sensing metalloregulators to control transcription of metal resistance genes. Depending on their metal-binding states, the MerR-family metalloregulators change their interactions with DNA to suppress or activate transcription. To understand their functions fundamentally, we study how CueR, a Cu¹⁺-responsive MerR-family metalloregulator, interacts with DNA, using an engineered DNA Holliday junction (HJ) as a protein-DNA interaction reporter in single-molecule fluorescence resonance energy transfer measurements. By analyzing the single-molecule structural dynamics of the engineered HJ in the presence of various concentrations of both apo- and holo-CueR, we show how CueR interacts with the two conformers of the engineered HJ, forming variable protein-DNA complexes at different protein concentrations and changing the HJ structures. We also show how apo- and holo-CueR differ in their interactions with DNA, and discuss their similarities and differences with other MerR-family metalloregulators. The surprising finding that holo-CueR binds more strongly to DNA than to apo-CueR suggests functional differences among MerR-family metalloregulators, in particular in their mechanisms of switching off gene transcription after activation. The study also corroborates the general applicability of engineered HJs as single-molecule reporters for protein-DNA interactions, which are fundamental processes in gene replication, transcription, recombination, and regulation.

INTRODUCTION

Metal ions are essential in biology and play key roles in the structure and function of a large number of proteins (1). Despite their importance, they can also be cytotoxic, especially at high concentrations (2,3). Intracellular metal ion concentrations and their bioavailability must therefore be tightly regulated to maintain normal cell metabolism. Bacteria, being susceptible to either limiting or toxic levels of metal ions in their living environment, have developed highly sensitive and selective metal homeostasis mechanisms (3–15). A key step in bacteria's response to varying levels of metal ions in their environment is through metal-sensing regulatory proteins (4–16). These proteins, also known as metalloregulators, respond to specific metal ions within the cell and regulate gene expression for metal-specific homeostasis (3–6).

A large class of bacterial metalloregulators belongs to the MerR-family; they respond to metal ions such as Hg²⁺, Pb²⁺, and Cu¹⁺ with high selectivity and sensitivity (4–6, 16–22). All MerR-family regulators are homodimers with two DNA-binding domains. They regulate gene transcription via a unique DNA distortion mechanism (5,17,18,23,24), in which both the apo- and the holo-regulator bind tightly to a dyad-symmetric sequence in the promoter region, with one DNA-binding domain binding to each half of the dyad sequence. In the apo-regulator bound form, DNA is slightly bent and the transcription is suppressed. Upon metal binding, the holo-regulator further unwinds DNA slightly, and transcription is activated. As the regulator-DNA interactions

dictate the transcription process, we are interested in defining the associated protein-DNA interactions quantitatively as a fundamental step to understand their detailed structure-dynamics-function relationships.

Single-molecule fluorescence resonance energy transfer (smFRET) measurements are powerful in studying protein-DNA interactions and associated structural changes of proteins and DNA (25–27). Owing to both the FRET mechanism and the fluorescent probes suitable for single-molecule detection, smFRET relies largely on detecting nanometer-scale distance changes (25,26). The structural changes associated with MerR-family regulator-DNA interactions are mainly on the angstrom scale, however (23,24). To detect small structural changes, we recently developed engineered DNA Holliday junctions (HJs) as generalizable single-molecule reporters in smFRET measurements for protein-DNA interaction studies (28).

Our method builds on the intrinsic structural dynamics of DNA HJs and the ease of following the dynamics by smFRET. In the presence of Na⁺ and Mg²⁺, each HJ molecule folds into two X-shaped stacked conformers that interconvert dynamically at room temperature (conf-I and conf-II, Fig. 1) (28–33). With a FRET donor-acceptor pair labeled at the ends of two HJ arms, the two conformers have distinct FRET signals, one having high FRET efficiency (E_{FRET}) and the other low E_{FRET} , and their interconversion dynamics are reflected by their two-state FRET fluctuation behaviors (28,29,31). To use a HJ as a single-molecule protein-DNA interaction reporter, we encode in its arms the dyad-symmetric sequence recognized by a metalloregulator (Fig. 1). Because the encoded sequence has distinct spatial orientations in the

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Technologies (Coralville, IA), and dissolved in 10 mM Tris buffer pH 7.3 with 100 mM NaCl. HJC2 was assembled by annealing strands *a*, *b*, and *c* first at 50°C; after slow cooling to 37°C, strand *d* was added. The solution was then incubated for 30 min at 37°C before cooling down to room temperature. The annealed HJC2 was purified by electrophoresis in 20% polyacrylamide gel.

Single-molecule fluorescence experiments and data analysis

A prism-type total internal reflection microscope based on an Olympus IX71 inverted microscope (Olympus, Melville, NY) was used for single-molecule fluorescence measurements. The Cy3 probe on HJC2 was directly excited by a continuous-wave circularly polarized 532-nm laser (GCL-025-L-0.5%, CrystaLaser, Reno, NV) of ~6 mW focused onto an area of $\sim 150 \times 75 \mu\text{m}^2$ on the sample. The fluorescence of both Cy3 and Cy5 was collected by a $60\times$ NA 1.2 water-immersion objective (UPLSAPO60XW, Olympus), with an extra $1.6\times$ magnification, and split by a dichroic mirror (635DCXR) into two channels using a Dual-View system (Optical Insights, Santa Fe, NM). HQ550LP filter was used to reject the excitation laser light and each channel of fluorescence was further filtered (HQ580-60m or HQ660LP) and projected onto half of the imaging area of a camera (Ixon EMCCD, DV887DCS-BV, Andor, Belfast, N. Ireland), controlled by the Andor IQ software. The time resolution for all the single-molecule experiments is 30 ms. A custom IDL program was then used to extract individual fluorescence trajectories of Cy3 and Cy5 for each HJC2 from the fluorescence movie recorded by the camera.

Single-molecule experiments were carried out using a flow cell, formed by double-sided tapes sandwiched between a quartz slide (Technical Glass, Opus, Snoqualmie, WA) and a borosilicate coverslip (Gold Seal coverslip, Thomas Scientific, Swedesboro, NJ). All samples were in 10 mM Tris buffer, pH 7.3 with 10 mM NaCl and 2 mM MgCl_2 . To minimize nonspecific protein adsorption on glass surfaces, quartz slides were first amine-functionalized (Vectabond, Vector Laboratories, Burlingame, CA) and then coated with polyethylene glycol (PEG) polymers (100 mg/mL m-PEG-SPA-5000, SunBio, Santa Clara, CA; and 1 mg/mL biotin-PEG-NHS-3400, JenKem Technology, Allen, TX) (26). One-percent of the PEG polymers contain a biotin terminal group to form biotin-streptavidin (Molecular Probes, Eugene, OR) linkages for immobilizing biotinylated HJC2 molecules. Oxygen scavenging system (0.1 mg/mL glucose oxidase (Sigma Chemical, St. Louis, MO), 0.025 mg/mL catalase (Roche, Hoffman-LaRoche, Basel, Switzerland), 4% glucose (Sigma Aldrich, St. Louis, MO)), and 1 mM Trolox (Sigma) were added into the sample solution just before each experiment to prolong the lifetime and suppress the blinking of the fluorescence probes (40).

E_{FRET} trajectories were obtained from Cy3 and Cy5 intensity trajectories as $E_{\text{FRET}} = I_{\text{Cy5}}/(I_{\text{Cy3}} + I_{\text{Cy5}})$, a good approximation for FRET efficiency. Photobleaching and blinking of either dye were first removed before performing threshold analysis on each trajectory based on the distribution of the E_{FRET} values to obtain individual waiting times (29). The average waiting time was then calculated from all trajectories obtained at a given protein concentration with the standard error of the mean as the error bar.

Fluorescence anisotropy

Fluorescence of Cy3-labeled double-strand DNA was measured using a Cary Eclipse fluorescence spectrophotometer (Varian, Palo Alto, CA). The CueR titration was in 10 mM Tris buffer with 10 mM NaCl and 2 mM MgCl_2 at pH 7.3, and PbrR691 titration in 10 mM Tris buffer with 100 mM NaNO_3 at pH 7.2. The sequence of the double-strand DNA for PbrR691 titration is 5'-TGACTCTATATCTACTAGAGGTT-3', where the PbrR691-specific dyad-symmetric sequence is underlined. The fluorescence was excited at 532 nm. Anisotropy (r) was calculated as $r = (I_{\parallel} - G * I_{\perp}) / (I_{\parallel} + 2 * G * I_{\perp})$, where I_{\parallel} and I_{\perp} are the fluorescence intensity parallel and perpendicular to the excitation polarization, respectively, and G is the correction factor for the instrument's different responses

to light of parallel and vertical polarizations. The fluorescence anisotropy titration curves were fitted with (41)

$$r = r_{\text{D}} + (r_{\text{PD}} - r_{\text{D}}) \times \frac{[\text{D}]_{\text{T}} + [\text{P}]_{\text{T}} + K_{\text{D}} - \sqrt{([\text{D}]_{\text{T}} + [\text{P}]_{\text{T}} + K_{\text{D}})^2 - 4[\text{P}]_{\text{T}}[\text{D}]_{\text{T}}}}{2[\text{D}]_{\text{T}}}, \quad (1)$$

where r_{D} and r_{PD} are the anisotropy values for free and protein-bound DNA, respectively, $[\text{D}]_{\text{T}}$ is the total DNA concentration, $[\text{P}]_{\text{T}}$ is the total protein concentration, and K_{D} is the dissociation constant of the protein-DNA complex.

RESULTS AND ANALYSIS

CueR-specific engineered HJ

Fig. 2A shows the design of the engineered Holliday junction, HJC2, targeting the metalloregulator CueR and using four oligo-DNA strands. The sequence of strand *a* is taken from the wild-type promoter that CueR binds, and it contains the CueR-specific dyad-symmetric sequence, which spans the arms *M* and *N*. The ends of arms *M* and *Q* are labeled with the FRET pair, Cy3 (donor) and Cy5 (acceptor), to distinguish between the two stacked conformers of HJC2: conf-I has a higher E_{FRET} and conf-II has a lower E_{FRET} (see Fig. 1). A biotin is attached at the end of arm *P* for surface immobilization. The assembly of HJC2 is confirmed by gel electrophoresis in reference to a characterized HJ (28), and by its absorption spectrum, in which the absorption bands of Cy3 and Cy5 indicate their 1:1 labeling ratio (Fig. 2A, inset).

The intrinsic structural dynamics of a single HJC2 molecule is clear from its anticorrelated two-state fluorescence intensity fluctuations in both the Cy3-donor and the Cy5-acceptor channel (Fig. 2B, upper). The corresponding E_{FRET} trajectory shows a two-state fluctuation between a high E_{FRET} (~0.59) and a low E_{FRET} (~0.17) state, corresponding to the structural interconversions between conf-I and conf-II (Fig. 2B, bottom). Past studies have shown that the two stochastic waiting times in the E_{FRET} trajectory, τ_{I} and τ_{II} , follow exponential distributions and the exponential decay constants are the interconversion rate constants (28,29). Therefore, $\langle \tau_{\text{I}} \rangle^{-1}$, where $\langle \dots \rangle$ denotes averaging and which represents the time-averaged single-molecule rate of conf-I \rightarrow conf-II transition, equals k_1 , the rate constant for conf-I \rightarrow conf-II transition. In addition, $\langle \tau_{\text{II}} \rangle^{-1}$, which represents the time-averaged single-molecule rate of conf-II \rightarrow conf-I transition, equals k_{-1} , the rate constant for conf-II \rightarrow conf-I transition (Supporting Material, Section A). For HJC2, the rate constants determined are $\langle \tau_{\text{I}} \rangle^{-1} = k_1 = 5.1 \pm 0.1 \text{ s}^{-1}$ and $\langle \tau_{\text{II}} \rangle^{-1} = k_{-1} = 0.85 \pm 0.02 \text{ s}^{-1}$.

Apo-CueR-HJC2 interaction dynamics

In the presence of CueR without Cu^{1+} bound, i.e., apo-CueR, significant perturbations are observed in the E_{FRET} trajectory of individual HJC2 molecules (Fig. 3A), indicating that

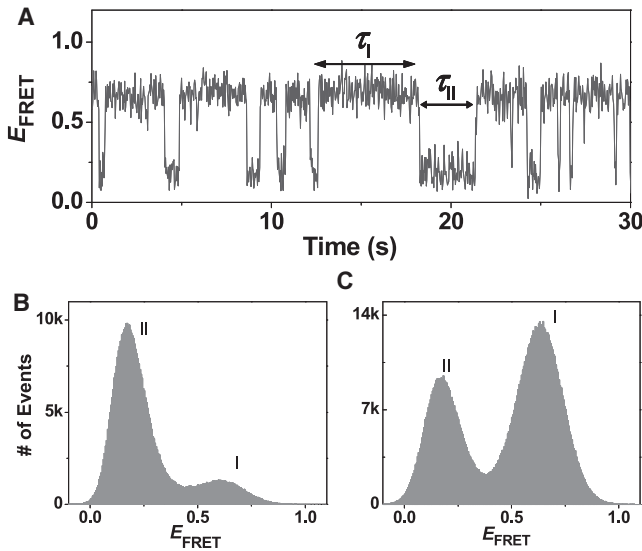


FIGURE 3 (A) Exemplary HJC2 E_{FRET} trajectory in the presence of 1.0 μM apo-CueR. (B and C) Histograms of HJC2 E_{FRET} trajectories in the absence (B) and presence (C) of 1.0 μM apo-CueR. Bin size: 0.005. Approximately 250 molecules were analyzed for each histogram. Histograms in the presence of other apo-CueR concentrations are given in Fig. S2.

apo-CueR binding alters HJC2 structural dynamics. The E_{FRET} trajectory shows a shift toward the high E_{FRET} state, i.e., conf-I. This shift in structural equilibrium is clearer in the E_{FRET} histogram (Fig. 3, B and C), where the intensity of the peak corresponding to conf-I increases relative to that of conf-II. This equilibrium shift reports the preferential binding of apo-CueR to conf-I over conf-II.

The structural equilibrium shift of HJC2 caused by apo-CueR binding is accompanied by changes in the interconversion kinetics. The time-averaged single-molecule rate of conf-I \rightarrow conf-II transition, $\langle\tau_I\rangle^{-1}$, depends on the apo-CueR concentration, [apo-CueR], decreasing asymptotically to zero with increasing [apo-CueR] (Fig. 4 A). This dependence indicates that apo-CueR binding slows down conf-I \rightarrow conf-II structural transition, lengthening the lifetime of conf-I. In contrast, the time-averaged single-molecule rate of conf-II \rightarrow conf-I transition, $\langle\tau_{II}\rangle^{-1}$, increases initially

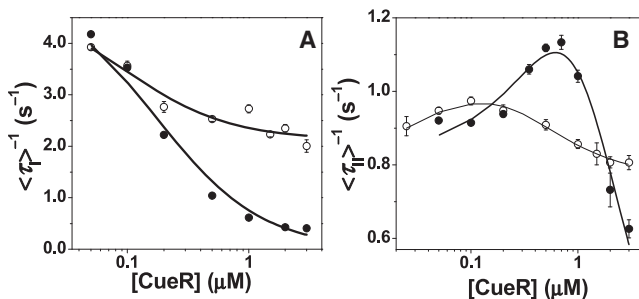


FIGURE 4 Apo-CueR (●) and holo-CueR (○) concentration dependence of $\langle\tau_I\rangle^{-1}$ (A) and $\langle\tau_{II}\rangle^{-1}$ (B). Each data point is an average of the waiting times from ~ 250 trajectories. The solid lines are the fits with Eq. 2 (apo) and Eq. 4 (holo) for panel A and Eq. 3 (apo) and Eq. 5 (holo) for panel B. Results from the fit are summarized in Table 1.

with increasing [apo-CueR], but decays at higher [apo-CueR] after reaching a maximum (Fig. 4 B). This biphasic behavior of $\langle\tau_{II}\rangle^{-1}$ indicates that the initial apo-CueR binding facilitates conf-II \rightarrow conf-I transition whereas a higher-order apo-CueR interaction slows it down.

The [apo-CueR] dependence of $\langle\tau_I\rangle^{-1}$ can be described by a simple kinetic mechanism in which apo-CueR (P) binds to conf-I to form a complex (P-I) that does not lead to structural transition to conf-II (Fig. 5 A, red box). Based on this kinetic scheme and following a single-molecule kinetic analysis (Supporting Material, Section B), we get

$$\langle\tau_I\rangle^{-1} = \frac{k_1}{1 + [P]/K_{P-I}} \quad (2)$$

Here $K_{P-I} (= k_{-2}/k_2)$ is the dissociation constant for the apo-CueR-conf-I complex, and k_2 and k_{-2} are the protein binding and unbinding rate constants to conf-I, respectively. Equation 2 predicts that with increasing protein concentration, $\langle\tau_I\rangle^{-1}$ decreases asymptotically to zero, consistent with the experimental results (Fig. 4 A). Using k_1 determined from the free HJC2 and fitting the data in Fig. 4 A give $K_{P-I} = 0.17 \pm 0.02 \mu\text{M}$.

To account for the biphasic [apo-CueR] dependence of $\langle\tau_{II}\rangle^{-1}$, we considered a two-step kinetic mechanism. Apo-CueR initially binds to conf-II to form a complex (P-II) that can lead to structural transition to conf-I; this P-II complex can then bind a second protein molecule to form a tertiary complex (P_2 -II) that does not lead to structural transition to conf-I (Fig. 5 A, green box). Based on this two-step interaction scheme between apo-CueR and conf-II and following a single-molecule kinetic analysis, we get

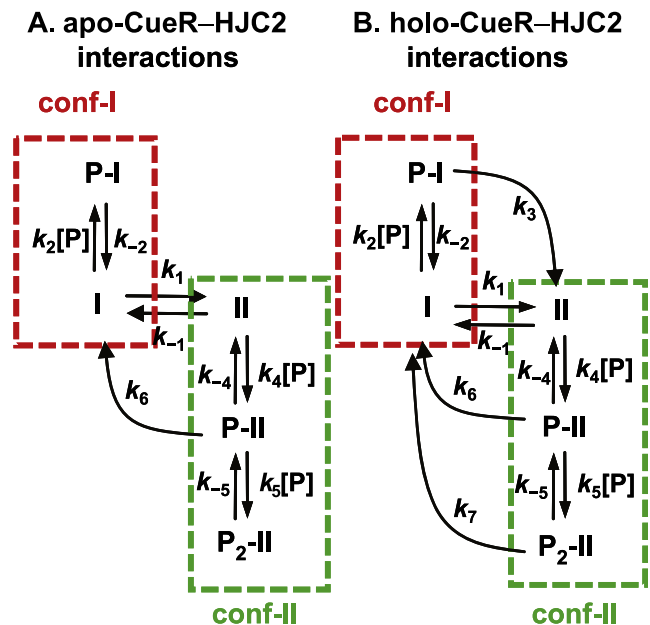


FIGURE 5 Kinetic schemes for HJC2 interactions with apo-CueR (A) and holo-CueR (B). I, conf-I; II, conf-II; P, apo-CueR or holo-CueR; and k -values, rate constants.

$$\langle \tau_{\text{II}} \rangle^{-1} = \frac{k_{-1} + k_6[\text{P}]/K'_{\text{P-II}}}{1 + [\text{P}]/K'_{\text{P-II}} + [\text{P}]^2/(K'_{\text{P-II}}K_{\text{P}_2\text{-II}})}. \quad (3)$$

Here $K'_{\text{P-II}} = (k_{-4} + k_6)/k_4$, $K_{\text{P}_2\text{-II}} = k_{-5}/k_5$, and k -values are the rate constants defined in Fig. 5 A. Equation 3 predicts that at low protein concentrations, $\langle \tau_{\text{II}} \rangle^{-1}$ increases with increasing protein concentration because the formation of complex P-II facilitates the structural transition to conf-I, and at higher protein concentrations, $\langle \tau_{\text{II}} \rangle^{-1}$ decreases because the formation of complex P₂-II slows down the transition to conf-I. Using k_{-1} determined from the free HJC2 and fitting the data in

$$\langle \tau_{\text{II}} \rangle^{-1} = \frac{k_{-1} + [\text{P}] \left(k_{-1}k_7 / (k_6K'_{\text{P}_2\text{-II}}) + k_6/K'_{\text{P-II}} \right) + [\text{P}]^2k_7 / (K'_{\text{P-II}}K'_{\text{P}_2\text{-II}})}{1 + [\text{P}] \left(k_7 / (k_6K'_{\text{P}_2\text{-II}}) + 1/K'_{\text{P-II}} \right) + [\text{P}]^2 / (K'_{\text{P-II}}K'_{\text{P}_2\text{-II}})}, \quad (5)$$

Fig. 4 B give: $k_6 = 4 \pm 3 \text{ s}^{-1}$, the rate constant for P-II \rightarrow I transition; $K'_{\text{P-II}} = 3 \pm 3 \mu\text{M}$, the apparent protein dissociation constant of complex P-II; and $K_{\text{P}_2\text{-II}} = 0.5 \pm 0.5 \mu\text{M}$, the first dissociation constant of complex P₂-II. The determined k_6 is higher than k_{-1} , indicating that the initial binding of apo-CueR to conf-II facilitates its structural transition to conf-I. Table 1 summarizes all the fitting results.

Holo-CueR-HJC2 interaction dynamics

Holo-CueR, i.e., Cu¹⁺-bound CueR, causes perturbations on HJC2 structural dynamics similar to apo-CueR. With increasing [holo-CueR], $\langle \tau_{\text{I}} \rangle^{-1}$, the time-averaged conf-I \rightarrow conf-II transition rate, decreases gradually (Fig. 4 A), and $\langle \tau_{\text{II}} \rangle^{-1}$, the time-averaged conf-II \rightarrow conf-I transition rate, increases initially and then decays at higher [holo-CueR] (Fig. 4 B). However, significant differences also exist: neither $\langle \tau_{\text{I}} \rangle^{-1}$ nor $\langle \tau_{\text{II}} \rangle^{-1}$ decays to zero at high [holo-CueR], in contrast to those of apo-CueR-HJC2 interactions. The nonzero values of $\langle \tau_{\text{I}} \rangle^{-1}$ and $\langle \tau_{\text{II}} \rangle^{-1}$ at high [holo-CueR] indicate that the relevant holo-CueR-HJC2 complexes can still allow transitions from one conformer of HJC2 to the other.

TABLE 1 Kinetic parameters for CueR-HJC2 interaction dynamics

		Free HJC2	
k_1		$5.1 \pm 0.1 \text{ s}^{-1}$	
k_{-1}		$0.85 \pm 0.02 \text{ s}^{-1}$	
		Apo-CueR	Holo-CueR
k_3		—	$2.3 \pm 0.2 \text{ s}^{-1}$
k_6		$4 \pm 3 \text{ s}^{-1}$	$1.2 \pm 0.3 \text{ s}^{-1}$
k_7		—	$0.77 \pm 0.02 \text{ s}^{-1}$
$(k_{-2} + k_3)/k_2$		$0.17 \pm 0.02 \mu\text{M} (K_{\text{P-I}})^*$	$0.05 \pm 0.02 \mu\text{M} (K'_{\text{P-I}})$
$(k_{-4} + k_6)/k_4$		$3 \pm 3 \mu\text{M} (K'_{\text{P-II}})$	$0.1 \pm 0.1 \mu\text{M} (K'_{\text{P-II}})^\dagger$
$(k_{-5} + k_7)/k_5$		$0.5 \pm 0.5 \mu\text{M} (K_{\text{P}_2\text{-II}})^\ddagger$	$0.2 \pm 0.2 \mu\text{M} (K'_{\text{P}_2\text{-II}})$

* $k_3 = 0$ for apo-CueR.

[†] k_{-4} is set to zero for holo-CueR (Supporting Material, Section C).

[‡] $k_7 = 0$ for apo-CueR.

To account for these differences observed for holo-CueR-HJC2 interactions, we added two kinetic transitions on top of the kinetic mechanism of apo-CueR-HJC2 interactions (Fig. 5 B). One transition connects P-I to II (i.e., k_3), and the other connects P₂-II to I (i.e., k_7). Based on this kinetic mechanism, the corresponding equations connecting $\langle \tau_{\text{I}} \rangle^{-1}$ and $\langle \tau_{\text{II}} \rangle^{-1}$ with the kinetic parameters and the protein concentration are (Supporting Material, Section C)

$$\langle \tau_{\text{I}} \rangle^{-1} = \frac{k_1 + [\text{P}]k_3/K'_{\text{P-I}}}{1 + [\text{P}]/K'_{\text{P-I}}}, \quad (4)$$

where $K'_{\text{P-I}} = (k_{-2} + k_3)/k_2$, $K'_{\text{P-II}} = k_6/k_4$, and $K'_{\text{P}_2\text{-II}} = (k_{-5} + k_7)/k_5$. The individual kinetic parameters are defined in Fig. 5 B. In deriving Eq. 5, we assumed $k_{-4} = 0$ to obtain a clean analytical expression; this assumption does not affect our analyses of $\langle \tau_{\text{I}} \rangle^{-1}$ (Eq. 4), from which $K'_{\text{P-I}}$ and k_3 can be obtained and interpreted quantitatively.

Equation 4 predicts that with increasing [holo-CueR], $\langle \tau_{\text{I}} \rangle^{-1}$ decreases and eventually approaches k_3 , the rate constant for the P-I \rightarrow II transition, consistent with experimental observations (Fig. 4 A) and resulting from that high [holo-CueR] drives the conversion of I \rightarrow P-I. Using k_1 determined from the free HJC2 and fitting the data in Fig. 4 A give $k_3 = 2.3 \pm 0.2 \text{ s}^{-1}$ and $K'_{\text{P-I}} = 0.05 \pm 0.02 \mu\text{M}$, which is the apparent dissociation constant of P-I. The determined k_3 is smaller than k_1 , consistent with the expectation that holo-CueR binding stabilizes conf-I and slow down its structural transition to conf-II.

Equation 5 predicts the observed [holo-CueR] dependence of $\langle \tau_{\text{II}} \rangle^{-1}$, with an initial rise followed by a decay. At high [holo-CueR], $\langle \tau_{\text{II}} \rangle^{-1}$ approaches k_7 , the rate constant of the P₂-II \rightarrow I transition, as high [holo-CueR] drives the formation of P₂-II. Using k_{-1} determined from the free HJC2 and fitting the data in Fig. 4 B give $k_7 = 0.77 \pm 0.02 \text{ s}^{-1}$, $K'_{\text{P-II}} = 0.1 \pm 0.1 \mu\text{M}$, which is the apparent dissociation constant of complex P-II, and $K'_{\text{P}_2\text{-II}} = 0.2 \pm 0.2 \mu\text{M}$, which is the apparent first protein dissociation constant of P₂-II. $K'_{\text{P-II}}$ for holo-CueR here is smaller than that ($3 \pm 3 \mu\text{M}$) for apo-CueR; this is consistent with that the maximum of $\langle \tau_{\text{II}} \rangle^{-1}$ occurs at lower [holo-CueR], as compared with the maximum in apo-CueR-HJC2 interactions (Fig. 4 B). The kinetic parameters are summarized in Table 1.

To confirm that the interactions between CueR and HJC2 are specific, i.e., due to the encoded specific dyad-symmetric sequence in HJC2, we studied the structural dynamics of HJC2 in the presence of another DNA-binding protein, PbrR691, which is also a MerR-family metalloregulator; no

noticeable perturbation was observed (Fig. S1). Additionally, we studied another HJ that does not contain CueR-targeting sequence; expectedly, in the presence of CueR, no noticeable perturbation on this HJ's structural dynamics was observed (28).

CueR-imposed HJC2 structural changes

Our single-molecule kinetic analyses on the structure dynamics of HJC2 indicate that both apo- and holo-CueR can bind to the two conformers of HJC2 to form complexes P-I, P-II, and P₂-II (Fig. 5). The changes in the E_{FRET} values of conf-I and conf-II in these complexes relative to those of free HJC2 can inform the structural changes of HJC2 imposed by the CueR. For interactions with conf-I, at high [apo-CueR] (e.g., 2 μM) where I is converted to P-I, $E_{\text{conf-I}}$ increases from ~ 0.59 to ~ 0.64 (Fig. 6, A and B). This increase of $E_{\text{conf-I}}$ indicates that in the P-I complex, apo-CueR causes a shortening of the distance between the ends of arms *M* and *Q* where Cy3 and Cy5 are located (see Fig. 1, left). This protein-induced structural change of conf-I also confirms the binding of apo-CueR. Similarly, holo-CueR binding also increases $E_{\text{conf-I}}$ (Fig. 6 C), indicating a similarly shortened distance between the ends of arms *M* and *Q* in conf-I.

To determine structural changes of conf-II upon CueR binding, we studied an alternatively labeled HJC2, referred to as HJC2a, which has the Cy5 placed at the end of arm *N*

instead of arm *Q* (see Fig. 1). This alternative labeling makes conf-II of HJC2a the high E_{FRET} state, rendering its E_{FRET} value ($E_{\text{conf-II}}$) more sensitive to structural changes imposed by protein binding. At low [apo-CueR] (e.g., 0.3 μM), where the complex P-II dominates the population of all forms of conf-II, no significant decrease in $E_{\text{conf-II}}$ is observed compared to that of free HJC2a (Fig. 6, D and E). At high [apo-CueR] (e.g., 3 μM), where P₂-II dominates, $E_{\text{conf-II}}$ increases significantly (Fig. 6 F). This clear increase indicates that in the P₂-II complex, where two apo-CueR molecules are bound, the arms *M* and *N* are brought closer to each other. These protein-induced structural changes of conf-II also confirm the binding of apo-CueR. For holo-CueR and conf-II interactions, in contrast, no significant changes of $E_{\text{conf-II}}$ were observed.

Ensemble CueR-DNA affinity determination

For apo-CueR interaction with conf-I of HJC2, the dissociation constant $K_{\text{P-I}} (= k_{-2}/k_2)$ directly reflects the binding affinity of apo-CueR to conf-I (Eq. 2). For holo-CueR, only the apparent dissociation constant $K'_{\text{P-I}} (= (k_{-2} + k_3)/k_2)$ is obtainable from analyzing the waiting times (Eq. 4), and it gives an upper limit for the dissociation constant of holo-CueR-conf-I interactions, as $(k_{-2} + k_3)/k_2 > k_{-2}/k_2$. The binding affinity to conf-I is important, as conf-I has its arms *M* and *N*, which contain the dyad-symmetric sequence,

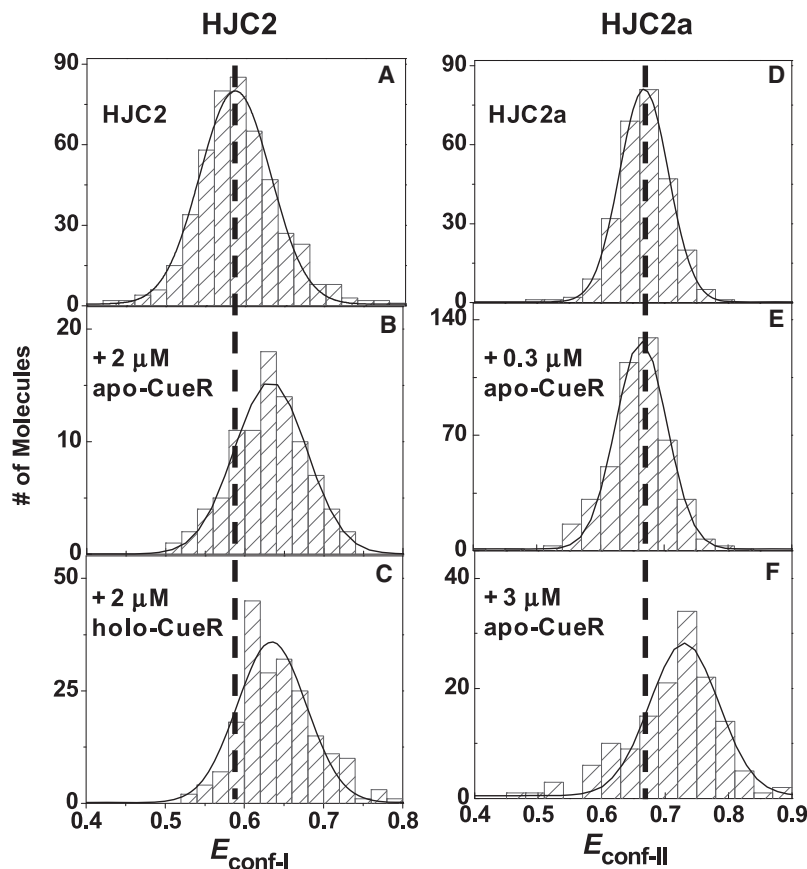


FIGURE 6 Histograms of $E_{\text{conf-I}}$ of HJC2 (A–C) and $E_{\text{conf-II}}$ of HJC2a (D–F) in the presence of various apo-CueR and holo-CueR concentrations. Solid lines are Gaussian fits centered at 0.59 ± 0.01 (A), 0.64 ± 0.01 (B), 0.63 ± 0.01 (C), 0.67 ± 0.01 (D), 0.66 ± 0.01 (E), and 0.73 ± 0.01 (F). For each molecule, its $E_{\text{conf-I}}$ or $E_{\text{conf-II}}$ was obtained by fitting the histogram of its E_{FRET} trajectory with two Gaussian functions. Histograms at another protein concentration are given in Fig. S3.

coaxially stacked as in a B-form DNA (32,33). Experimentally, K_{P-I} ($= 0.17 \pm 0.02 \mu\text{M}$) for apo-CueR is larger than K'_{P-I} ($= 0.05 \pm 0.02 \mu\text{M}$) for holo-CueR (Table 1), indicating that apo-CueR binds weaker to conf-I than holo-CueR does.

On the other hand, previous results from the gel-shift assay suggested a slightly stronger binding affinity for apo-CueR than for holo-CueR to a double-strand DNA containing the promoter sequence, with the dissociation constants $K_D(\text{apo-CueR}) = 17 \pm 2 \text{ nM}$ and $K_D(\text{holo-CueR}) = 25 \pm 7 \text{ nM}$ (36), although the error bars of these values preclude a reliable comparison. To test whether there is indeed a discrepancy that could come from the DNA used (i.e., using engineered HJ versus using a double-strand DNA), we determined the binding affinity of both apo- and holo-CueR to a double-strand DNA using ensemble fluorescence anisotropy titration, a more accurate quantitation method than the gel-shift assay. We used a Cy3-labeled 25-basepair double-strand DNA with the same sequence as that spanning the *M*, *N* arms of HJC2. The results in Fig. 7 A confirm that apo-CueR does bind weaker to DNA than holo-CueR, with $K_D(\text{apo-CueR}) = 6 \pm 2 \text{ nM}$ and $K_D(\text{holo-CueR}) = 1.9 \pm 0.8 \text{ nM}$, consistent with the results using the engineered HJC2 as a single-molecule reporter. The affinities of apo- and holo-CueR to conf-I of HJC2 are weaker than to the double-strand DNA, possibly due to the presence of the other helix or the perturbation of the junction structure of HJC2 on the protein-DNA interactions.

DISCUSSION

Nature of CueR-HJC2 interactions and relation to protein function

The single-molecule studies of CueR-HJC2 interactions indicate that both apo- and holo-CueR preferentially bind and stabilize conf-I of HJC2. This preferential interaction

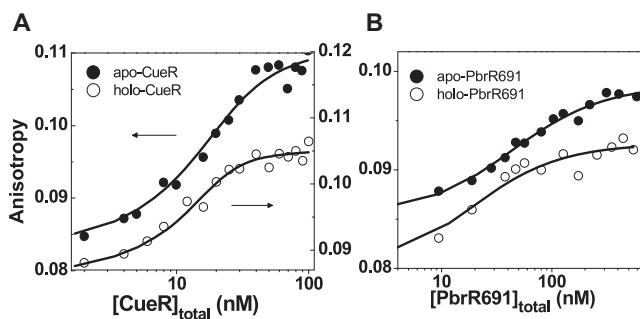


FIGURE 7 Fluorescence anisotropy titrations of CueR (A) and PbrR691 (B) binding to Cy3-labeled double-strand DNA. Solid lines are the fits with Eq. 1, with dissociation constants of $K_D(\text{apo-CueR}) = 6 \pm 2 \text{ nM}$, $K_D(\text{holo-CueR}) = 1.9 \pm 0.8 \text{ nM}$, $K_D(\text{apo-PbrR691}) = 43 \pm 9 \text{ nM}$, and $K_D(\text{holo-PbrR691}) = 23 \pm 9 \text{ nM}$. The *x* axes are total protein concentrations. Plots with *x* axes being the free protein concentration are given in Fig. S5.

is directly reflected by HJC2's structural equilibrium shift toward conf-I (Fig. 3, B and C), as well as the decrease of $\langle \tau_I \rangle^{-1}$ with increasing protein concentration due to the formation of the P-I complex (Fig. 4 A and Fig. 5). The preferential interaction with conf-I is also reflected by the protein concentration dependence of $\langle \tau_{II} \rangle^{-1}$, which shows that the P-II complex can induce structural transition from conf-II to conf-I with a larger rate constant (k_6) than that (k_{-1}) of the intrinsic II \rightarrow I transition (Table 1 and Fig. 5). All these observations are consistent with CueR's normal function as a double-strand DNA-binding protein, as conf-I mimics the natural substrate of CueR and has its arms *M* and *N*, which encodes the dyad-symmetric sequence, coaxially stacked to form a B-form DNA-like structure (Fig. 1, left) (32,33).

Both apo- and holo-CueR binding to conf-I bring the arms *M* and *Q* closer, reflected by the increase of $E_{\text{conf-I}}$ of HJC2 (Fig. 6, B and C). This structural change can be associated with the bending of the *M-N* helix of conf-I, similarly observed in our previous study of PbrR691 (28) and typical among MerR-family metalloregulators (5,23,24). For conf-II, no significant structural change is observed upon initial binding of one apo-CueR or holo-CueR molecule.

A distinct feature of CueR-HJC2 interactions is the initial-rise-followed-by-decay of $\langle \tau_{II} \rangle^{-1}$ with increasing protein concentrations (Fig. 4 B). This biphasic protein concentration dependence of $\langle \tau_{II} \rangle^{-1}$ indicates that at high protein concentrations, both apo- and holo-CueR, interact with conf-II to form the P₂-II complex, in which two protein molecules are bound (Fig. 5). This tertiary complex was not observed in the study of PbrR691 (28), suggesting that differences exist among MerR-family metalloregulators in their interactions with DNA. The formation of this tertiary complex could be related to the highly bent orientation of the two halves of the dyad-symmetric sequence in conf-II, which largely deviates from the structure of a B-form DNA (Fig. 1, right). Consequently, only half of the dyad sequence could be bound to one of the two DNA-binding domains of CueR, leaving the other half to bind another CueR molecule. The double binding of apo-CueR then leads to the *M* and *N* arms being pushed closer in conf-II, reflected by the increase in $E_{\text{conf-II}}$ of HJC2a (Fig. 6 F). However, double binding of holo-CueR does not result in observable changes in $E_{\text{conf-II}}$, indicating different interactions with DNA between apo- and holo-CueR.

To support our model that CueR can bind to half of the dyad-symmetric sequence, we constructed a Cy3-labeled double-strand DNA that encodes only half of the CueR-specific dyad-symmetric sequence. We then used fluorescence anisotropy titration to probe CueR binding. The results (Fig. S4) show that apo-CueR can indeed bind to this DNA with a K_D of $\sim 0.7 \mu\text{M}$, which is in the concentration range where the P₂-II complex forms (Fig. 4 B).

Further differences exist between apo-CueR and holo-CueR. Unlike apo-CueR, holo-CueR binding can still allow P-I \rightarrow II and P₂-II \rightarrow I transitions (k_3 and k_7 , Fig. 5 B). As conf-I and conf-II are largely different in their spatial

arrangements of the dyad-symmetric sequence (Fig. 1), being more accommodating in allowing HJC2 structural interconversion suggests that holo-CueR has more flexible conformation than apo-CueR. This conformational flexibility of holo-CueR could play important roles in its interaction with the RNA polymerase (RNAP) for transcription, as past studies on MerR, the prototype MerR-family metalloregulator, showed that the holo-MerR-DNA-RNAP tertiary complex undergoes structural rearrangements in transcription initiation (5,17,18,42).

Implications for transcriptional suppression after activation

Our single-molecule studies and the ensemble fluorescence anisotropy titration both indicate a stronger binding of holo-CueR to DNA than apo-CueR. The stronger DNA binding of the holo-protein is also observed for PbrR691 from fluorescence anisotropy titrations (Fig. 7 B), with the dissociation constants of $K_D(\text{apo-PbrR691}) = 43 \pm 9 \text{ nM}$ and $K_D(\text{holo-PbrR691}) = 23 \pm 9 \text{ nM}$. Furthermore, BmrR, another MerR-family regulator that responds to organic effectors, also has a higher affinity to its substrate in its holo-form compared to its apo-form (43).

This stronger DNA binding by the holo-protein is surprising, however, as past studies on MerR have shown that the holo-protein binds weaker to DNA than the apo-protein ($K_D(\text{apo-MerR}) = 0.14 \pm 0.04 \text{ nM}$, $K_D(\text{holo-MerR}) = 0.42 \pm 0.07 \text{ nM}$) (17). As the direct dissociation of the metal ion from the metalloregulator is believed to be difficult due to strong metal coordination, it was thought that the weaker binding of holo-protein would facilitate its replacement from the DNA by the apo-protein, thus switching off the transcription after transcriptional activation once the cell is relieved of the metal stress. Therefore, the opposite relative DNA binding affinity of apo-protein versus holo-protein for CueR (as well as PbrR691 and BmrR) suggests possible differences in the mechanism by which MerR-family regulators switch off transcription.

Moreover, unlike MerR, which might involve another protein MerD to help the dissociation of the holo-MerR-DNA complex (16), no evidence has so far been found for a coregulator role of a MerD homolog in the regulatory mechanism of CueR (16). For CueR to switch off transcription after activation, one simple scenario is a direct dissociation of holo-CueR from DNA followed by binding of apo-CueR, which would be the dominant form of CueR inside the cell after activation of Cu-resistance genes. For this scenario to be viable, the dissociation kinetics of holo-CueR from DNA has to be in a relevant timescale to gene regulation. From our single-molecule kinetic analyses, the rate constants for CueR unbinding (k_{-2}) and binding (k_2) to conf-I of HJC2 cannot be obtained. Nevertheless, as the CueR binding and unbinding are contained in the observed structural dynamics of HJC2, which we measure experimentally, CueR binding and

unbinding should occur at a comparable timescale to HJC2's structural dynamics, i.e., hundreds of milliseconds to seconds, a relevant timescale for gene expression regulation.

SUMMARY

Using the engineered HJC2 as a single-molecule protein-DNA interaction reporter, we have studied how CueR, a Cu^{1+} -responsive MerR-family metalloregulator, interacts with DNA. Both apo- and holo-CueR preferentially bind conf-I of HJC2, in which the protein-recognition sequence is arranged similarly as in a B-form DNA. This preferential binding stabilizes conf-I and slows down its structural conversion to conf-II. Both also bend the *M-N* helix of conf-I, reflecting their actions on DNA for transcriptional regulation. In their interactions with conf-II of HJC2, apo- and holo-CueR exhibit a biphasic behavior—at low protein concentrations, a 1:1 protein-conf-II complex is present, whereas at high protein concentrations a 2:1 protein-conf-II tertiary complex dominates. Many differences also exist between apo- and holo-CueR in their interactions with HJC2. Whereas apo-CueR causes clear structural changes of both conf-I and conf-II of HJC2, holo-CueR only causes measurable structural changes of conf-I. Holo-CueR is also more accommodating in allowing the structural interconversions of HJC2, suggesting its more flexible conformation, which could be important for its cooperation with RNAP in initiating transcription. Moreover, holo-CueR binds stronger to DNA than apo-CueR, a surprising finding and contrary to the behaviors of the prototype metalloregulator MerR. This contrast suggests functional differences among MerR-family regulators, in particular possible different mechanisms in switching off transcription after activation. The study also corroborates the general applicability of engineered HJs as single-molecule reporters for protein-DNA interactions, which are fundamental in gene replication, transcription, recombination, and regulation.

SUPPORTING MATERIAL

Derivations of equations and additional results are available at [http://www.biophysj.org/biophysj/supplemental/S0006-3495\(09\)01033-9](http://www.biophysj.org/biophysj/supplemental/S0006-3495(09)01033-9).

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