Biphasic unbinding of Zur from DNA for transcription (de)repression in Live Bacteria

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Abstract

Transcription regulator on-off binding to DNA constitutes a mechanistic paradigm in gene regulation, in which the repressors/activators bind to operator sites tightly while the corresponding non-repressors/non-activators do not. Another paradigm regards regulator unbinding from DNA to be a unimolecular process whose kinetics is independent of regulator concentration. Using single-molecule single-cell measurements, we find that the behaviors of the zinc-responsive uptake regulator Zur challenges these paradigms. Apo-Zur, a non-repressor and presumed non-DNA binder, can bind to chromosome tightly in live E. coli cells, likely at non-consensus sequence sites. Moreover, the unbinding from DNA of its apo-non-repressor and holo-repressor forms both show a biphasic, repressed-followed-by-facilitated kinetics with increasing cellular protein concentrations. The facilitated unbinding likely occurs via a ternary complex formation mechanism; the repressed unbinding is first-of-its-kind and likely results from protein oligomerization on chromosome, in which an inter-protein salt-bridge plays a key role. This biphasic unbinding could provide functional advantages in Zur's facile switching between repression and derepression.
INTRODUCTION

Transcriptional regulation in cells is generally orchestrated by regulators, which, upon binding to operator sites, either block the binding of RNA polymerase (RNAP) leading to repression (i.e., repressors) or recruit RNAP leading to activation (i.e., activators)\(^1\)-\(^2\). One mechanistic paradigm for these regulators is an on-off model in which they bind to their cognate operator sites tightly, while their corresponding non-repressor/non-activator forms have insignificant affinity to DNA and stay predominantly in the cytoplasm. Some exceptions recently emerged. For example, IscR, a member of the MarA/SoxS/Rob family of transcription regulators in *E. coli*, is a repressor in its holo-form (i.e., containing a Fe-S cluster); its apo-form, generally thought to not bind DNA, was shown to bind DNA motifs different from its holo-repressor form\(^3\)-\(^4\).

Derepression or deactivation subsequently comes from the unbinding of the regulator from the operator site. Here another mechanistic paradigm exists regarding the kinetics of regulator unbinding, which is presumed to be a unimolecular reaction (i.e., spontaneous unbinding), whose first-order rate constant is independent of surrounding regulator concentration. However, recent *in vitro* single-molecule and bulk measurements uncovered facilitated unbinding, in which the first-order unbinding rate constant increases with increasing protein concentrations\(^5\). These proteins include nucleoid associated proteins that bind double-stranded DNA nonspecifically\(^6\), replication protein A that binds single-stranded DNA nonspecifically\(^7\), and DNA polymerases\(^8\)-\(^9\). We also discovered that CueR and ZntR, two MerR-family metal-sensing transcription regulators that bind to their cognate promoter sequences specifically, also show facilitated unbinding\(^10\). Using single-molecule tracking (SMT) and single cell quantification of protein concentration (SCQPC) that connect protein-DNA interaction kinetics with cellular protein concentrations quantitatively, we further showed that the facilitated unbinding of CueR and ZntR also operate in living *E. coli* cells\(^11\). A mechanistic consensus emerged, involving multivalent contacts between the protein and DNA\(^2\), which enables the formation of ternary complexes as intermediates that subsequently give rise to concentration-enhanced protein unbinding kinetics.

Here we report a SMT and SCQPC study of Zur, a Fur-family homodimeric zinc-uptake regulator, whose Zn\(^{2+}\)-bound holo-form binds to its cognate operator site with nM affinity and represses the transcription of zinc uptake genes under zinc stress\(^12\)-\(^15\); its apo-form is a non-repressor. We found that in living *E. coli* cells, Zur's interactions with DNA challenge the above two paradigms. First, apo-Zur, long thought to not bind DNA, can bind to chromosome tightly, likely at non-consensus sites. Second and more strikingly, the unbinding of both apo- and holo-Zur from chromosome not only show facilitated unbinding with increasing cellular protein concentrations, but also exhibit repressed unbinding at lower concentrations, giving a first-of-its-kind biphasic unbinding behavior. The repressed unbinding of Zur likely stems from Zur oligomerization on DNA, where an inter-dimer salt bridge plays a key role, and it likely facilitates transcription switching between repression and depression in cells.

RESULTS

SMT and SCQPC identify a tight DNA-binding state for both holo- and apo-Zur in cells

To visualize individual Zur proteins in *E. coli* cells, we fused the photoconvertible fluorescent protein mEos3.2\(^16\), \(^17\) to its C-terminus creating Zur\(^\text{mt}^\text{ }\), either at its chromosomal locus to have physiological expression or in an inducible plasmid in a Δ*zur* deletion strain to have a wider range of cellular protein concentrations (Methods). This Zur\(^\text{mt}^\text{ }\) fusion-protein is intact and as functional a repressor as the wild-type (WT) in the cell under Zn stress growth conditions (Supplementary Fig. 1a-b).

Using sparse photoconversion and time-lapse stroboscopic imaging, we tracked the motions of photoconverted Zur\(^\text{mt}^\text{ }\) proteins individually in single *E. coli* cells at tens of nanometer precision until their mEos3.2 tags photobleached (Fig. 1a). This SMT allows for measuring Zur\(^\text{mt}^\text{ }\)'s mobility, which reports on whether the molecule is freely diffusing in the cell or bound to DNA. We repeated this
photoconversion and SMT cycle 500 times for each cell, during which we counted the number of tracked protein molecules. We then used the SCQPC protocol to quantify the remaining number of Zur\textsuperscript{mE} protein molecules in the same cell\textsuperscript{11}, eventually determining the Zur\textsuperscript{mE} concentration in each cell (i.e., [Zur\textsuperscript{mE}]\textsubscript{cell}). This single-cell protein quantitation allowed for sorting the cells into groups of similar protein concentrations and subsequently examining protein-concentration–dependent processes, without being limited by the large cell-to-cell heterogeneity in protein expression.

We first examined Zur\textsuperscript{mE} whose regulatory Zn-binding site was mutated (i.e., C88S) to make it permanent apo and a non-repressor\textsuperscript{15} (Supplementary Fig. 1b). To quantify its mobility in cells, we determined the distribution of its displacement length $r$ between successive images and the corresponding cumulative distribution function (CDF) of $r$ for each cell group having similar cellular Zur\textsuperscript{mE} concentrations (Fig. 1b-c). Global analysis of these CDFs across all cellular protein concentrations resolved minimally three Brownian diffusion states with effective diffusion constants of $\sim 5.0 \pm 0.5, 0.82 \pm 0.05$, and $0.040 \pm 0.003 \mu\text{m}^2 \text{s}^{-1}$ (Fig. 1b-c; Methods). No subcellular localization or protein aggregation was observed; therefore, these two aspects are not the reasons for the presence of these three diffusion states. On the basis of their diffusion constants and previous studies of transcription regulator diffusion in \textit{E. coli} cells\textsuperscript{11,18-21}, we assigned the fastest diffusion state as Zur\textsuperscript{apo} proteins freely diffusing (FD) in the cytoplasm, the medium diffusion state as those nonspecifically bound (NB) to and moving on chromosome, and the slowest state as those tightly bound (TB) to the chromosome, whose small effective diffusion constant ($\sim 0.040 \mu\text{m}^2 \text{s}^{-1}$) reflects chromosome dynamics\textsuperscript{19, 22} and measurement uncertainties. Control measurements on the free mEos3.2 further support the assignment of the FD state, as we reported\textsuperscript{11}.

The resolution of CDFs of $r$ also gave the fractional populations of the three states across the range of cellular protein concentrations (Fig. 1d). With increasing [Zur\textsuperscript{mE}]\textsubscript{cell}, the fractional population of the FD state increases, while that of the TB state decreases. These trends further support their assignments because, with increasing cellular protein concentrations, more proteins compete for the limited number of tight binding sites on chromosome, leading to smaller fractional populations of the TB state and larger fractions of the FD state.

The presence of a significant fraction of the tight DNA-binding state, even at low cellular protein concentrations, is surprising for Zur\textsuperscript{apo} (e.g., $\sim 32\%$ at [Zur\textsuperscript{apo}]\textsubscript{cell} $\sim 60 \text{nM}$; 1 nM in an \textit{E. coli} cell corresponds to $\sim 1$ protein copy), as apo-Zur is a non-repressor. Furthermore, previous gel shift assay showed that \textit{E. coli} apo-Zur does not bind to operator sites (i.e., $K_D > 300 \text{nM}$ at the \textit{znuABC} promoter)\textsuperscript{15}, and for \textit{B. subtilis}, its apo-Zur’s binding affinity to operator sites is $\sim 1000$ times weaker than its holo-form\textsuperscript{23}. We hypothesized that the TB state of Zur\textsuperscript{mE} likely comes from its binding to nonoperator sites (i.e., non-consensus sequence sites; see later).

We next examined Zur\textsuperscript{mE} in cells stressed with 20 $\mu\text{M}$ Zn\textsuperscript{2+} in the medium. This Zn\textsuperscript{2+} concentration can evoke maximal repression of zur regulons (Supplementary Note 2.3). Therefore, most of Zur proteins in the cell should be metallated, mimicking the holo repressor form (i.e., Zur\textsuperscript{mE}). The same three diffusion states are resolved in the CDFs of $r$ across all cellular protein concentrations (Supplementary Note 4.2). In contrast to the case for Zur\textsuperscript{apo}, the TB state of Zur\textsuperscript{mE} is expected here because holo-Zur binds specifically to consensus operator sites within Zur-regulated promoters. Expectedly, the fractional population of the FD state of Zur\textsuperscript{mE} increases with increasing [Zur\textsuperscript{mE}]\textsubscript{cell}, whereas that of the TB state decreases (Fig. 1d).

\textbf{Concentration-dependent biphasic unbinding kinetics of Zur from DNA}

To probe Zur–DNA interaction dynamics, we examined the $r$ versus time $t$ trajectories of individual Zur proteins inside cells. These trajectories show clear transitions between large and small $r$ values (Fig. 2a): the small $r$ values are expected to be dominated by instances of Zur tightly bound to chromosome (i.e., TB state). We set an upper threshold $r_0 (= 0.2 \mu\text{m})$, below which $>99.5\%$ of the TB
states are included based on the resolved distributions of $r$ (Fig. 1b), to select these small displacements and obtain estimates of the individual residence time $\tau$ of a single Zur protein at a chromosomal tight binding site (Fig. 2a). Each $\tau$ starts when $r$ drops below $r_0$ and ends when $r$ jumps above $r_0$ (e.g., $\tau$'s in Fig. 2a), which are expected to reflect dominantly protein unbinding from DNA, or when the mEos3.2-tag photobleaches/blinks.

We analyzed trajectories from many cells of similar cellular Zur concentrations to obtain their corresponding distribution of $\tau$ (Fig. 2b). We used a quantitative three-state model (i.e., FD, NB, and TB states; Fig. 2c) to analyze the distribution of $\tau$, in which the contributions of FD and NB states are deconvoluted (Eq. (4); approximations and validations of this model in Supplementary Note 5)\textsuperscript{11}. This model also accounts for mE photobleaching/blinking kinetics, determined from the fluorescence on-time distribution of SMT trajectories (Supplementary Fig. 8). This analysis gave $k_a$, the apparent first-order unbinding rate constant of Zur from a tight binding site on the chromosome, for each group of cells having similar cellular Zur concentrations.

Strikingly, $k_a$ for Zur\textsubscript{apo}\textsuperscript{mE} shows a biphasic, repressed-followed-by-facilitated behavior: it initially decreases with increasing free (or total) cellular Zur concentration (i.e., repressed), reaching a minimum at \textasciitilde130 nM; it then increases toward higher protein concentrations (i.e., facilitated; Fig. 2d, left, blue points). This biphasic behavior is also apparent in the simple averages of residence time ($\langle \tau \rangle$) or by analyzing the distributions of $\tau$ that merely takes into account mE photobleaching/blinking (Supplementary Fig. 9a). The facilitated unbinding of Zur\textsubscript{apo}\textsuperscript{mE} is analogous to those of CueR and ZntR, two MerR-family metalloregulators that we discovered \textit{in vitro} and in living cells\textsuperscript{10, 11}; the repressed unbinding of Zur\textsubscript{apo}\textsuperscript{mE} is a \textit{first-of-its-kind} discovery, however.

In contrast, $k_a$ for Zur\textsubscript{apo}\textsuperscript{Zn} only shows the facilitated unbinding within the accessible cellular protein concentration range (\textasciitilde30 to \textasciitilde900 nM) — it increases consistently with increasing cellular protein concentrations (Fig. 2d, left, red points). The different behaviors of Zur\textsubscript{apo}\textsuperscript{Zn} from that of Zur\textsubscript{apo}\textsuperscript{mE} indicate that we could indeed observe the behaviors of the holo-repressor.

**Mechanism of biphasic unbinding of Zur from DNA**

Amid the biphasic unbinding of Zur from DNA (Fig. 2d, left), the concentration-facilitated unbinding at higher protein concentrations is analogous to those of CueR and ZntR\textsuperscript{11}. There it stems from an assisted dissociation pathway, in which an incoming protein from solution helps an incumbent protein on DNA to unbind, or a direct substitution pathway, in which the incoming protein directly replaces the incumbent one (Fig. 2e, lower)\textsuperscript{10, 11}. The rates of both pathways depend linearly on the free protein concentration, and both likely occur through a common ternary protein—DNA complex, in which the two homodimeric proteins each use one DNA-binding domain to bind to half of the dyad recognition sequence\textsuperscript{5, 24}. As Zur is also a homodimer, Zur could also form this ternary complex and undergo assisted dissociation or direct substitution, leading to its concentration-facilitated unbinding from DNA.

Regarding the repressed unbinding of apo-Zur in the lower concentration regime, we propose that it likely results from protein oligomerization around the DNA binding site, in which the number of proteins in the oligomer increases with increasing protein concentration and the resulting protein-protein interactions contribute to additional stabilization, thereby repressing protein unbinding rate (Fig. 2e, upper). (The facilitated unbinding later takes over when the protein concentration reaches a high enough level.) Two evidences support our oligomerization proposal: (1) Crystallography showed that two \textit{E. coli} Zur dimers can bind to a short cognate DNA sequence\textsuperscript{15}. (2) DNA footprinting showed that \textit{S. coelicoror} Zur forms oligomers around its recognition sites, containing greater than 4 dimers\textsuperscript{25}.

To further support this oligomerization proposal, we examined the spatial distribution in the cell of Zur's residence sites at its TB state; these residence sites correspond to the $r_0$-thresholded small displacements (Fig. 2a; Supplementary Note 8). For comparison, we further simulated an equal number of sites randomly distributed in a cell of the same size (Supplementary Note 8.1). We then examined
their pair-wise distance distributions (PWD), in which Zur oligomerization at chromosomal binding
sites should lead to more populations at shorter pair-wise distances. This PWD for Zur\textsuperscript{mE}\textsubscript{apo} indeed shows
a higher population at distances shorter than ~500 nm relative to the simulated random sites (Fig 3a).
However, at the distance scale of a few hundred nanometers, the compaction of chromosome also
contributes to the PWD of residence sites\textsuperscript{11}. To decouple the contribution of protein oligomerization
from chromosome compaction, we examined the fraction of residence sites within a radius threshold R.
At small R (e.g., <100 nm), the contribution of Zur oligomerization to this fraction should dominate
over chromosome compaction, as oligomerization is at molecular scale whereas the most compact
chromosome in a E. coli cell is still around hundreds of nanometer in dimension\textsuperscript{11,26}. At any specified
R (e.g., 200 nm), the fraction of Zur\textsuperscript{mE}\textsubscript{apo} residence sites within the radius R increases expectedly with
increasing cellular protein concentrations (Fig 3b, red points), because higher protein concentrations
gave higher sampling frequency of residence sites. More important, at lower R (e.g., 100 nm), the
fraction of Zur\textsuperscript{mE}\textsubscript{apo} residence sites is larger than that of simulated random sites (Fig 3b, red vs. blue
points), and their ratio is larger at lower protein concentrations (Fig 3b, green points). The average ratio
of the fraction of Zur\textsuperscript{mE}\textsubscript{apo} residence sites over that of the simulated random sites is always greater than
1, and it becomes larger at smaller R down to <70 nm (Fig. 3c; note our molecular localization precision
is ~20 nm; Supplementary Note 3), supporting Zur\textsuperscript{mE}\textsubscript{apo} oligomerization at chromosomal tight binding
sites at the nanometer scale.

We formulated a quantitative kinetic model to describe the biphasic unbinding of Zur\textsuperscript{mE}\textsubscript{apo}. It
considers both oligomerization at a TB site and facilitated unbinding via a ternary protein-DNA
complex (Fig. 2c and e; Supplementary Note 6). The microscopic unbinding rate constant \(k_1^{(n)}\) from a
TB site with n Zur\textsuperscript{mE}\textsubscript{apo} dimers bound as an oligomer comprises three terms:
\[
k_{1}^{(n)} = k_0 + k_1 \alpha^n + k_1 [P]_{FD}
\]
(1)
k\(_0\) is a first-order intrinsic unbinding rate constant. The \(k_1 \alpha^n\) term accounts for the repressed unbinding
from protein oligomerization, where a first-order rate constant \(k_0\) is attenuated by the factor \(\alpha\) (0 \(<\alpha\) \(<\text{1})\) to the exponent of \(n\), which depends on the cellular protein concentration and has a maximal value
of \(n_0\), the oligomerization number. The third term describes the facilitated unbinding, with \(k_1\) being a
second-order rate constant and \([P]_{TD}\) being the concentration of freely diffusing Zur dimers in the cell,
as reported for CueR/ZntR\textsuperscript{11}. In the limit of weak oligomerization and low free protein concentrations,
the apparent unbinding rate constant \(k_a\) from any TB site is:
\[
k_a = \left(\frac{k_{1}^{(n)}}{K_m}\right) = k_{0}^{\text{off}} + k_1 \left(e^{-[P]_{TD}/K_m} - 1\right) + k_I [P]_{FD}
\]
(2)
\(K_m = \frac{k_0^{\text{off}}}{k_1(1-\alpha)}\): it has the units of protein concentration, reflecting the effective dissociation constant of the
protein oligomer on the chromosome. \(k_0^{\text{off}} = k_0 + k_1\); it is a first-order spontaneous unbinding rate
constant at the limit of zero cellular protein concentration. Equation (2) satisfactorily fits the biphasic
unbinding kinetics of Zur\textsuperscript{mE}\textsubscript{apo} (Fig. 2d, left), giving the associated kinetic parameters (Table 1 and
Supplementary Table 6). In particular, \(K_m\) of Zur\textsuperscript{mE}\textsubscript{apo} is ~5 nM, indicating that apo-Zur can oligomerize
on chromosome at its physiological concentrations in the cells (Fig. 4a).

The same model also allowed for analyzing the relative populations of FD, NB, and TB states
of Zur across all cellular protein concentrations, giving additional thermodynamic and kinetic
parameters (Table 1, and Supplementary Table 6). Strikingly, the dissociation constant \(K_{d1}\) of Zur\textsuperscript{mE}\textsubscript{apo} at
TB sites of DNA is ~11 nM, merely ~2 times weaker than that of Zur\textsuperscript{mE} (\(K_{d1}\sim5\) nM). This is not
expected because apo-Zur, in both E. coli and B. subtilis, was shown to have no significant affinity to
the consensus sites recognized by holo-Zur\textsuperscript{15,23}. Therefore, the high affinity of Zur\textsuperscript{mE}\textsubscript{apo} at the TB state
suggests that inside cells, apo-Zur likely bind tightly to other, non-consensus sites in the chromosome.
This likelihood is supported by a ChIP-seq analysis in *B. subtilis*, which showed Zur can bind tightly to many locations in the chromosome that do not share consensus with the known recognition sites (although it was undefined whether the detected bindings there were by apo- or holo-Zur). 

**Molecular basis of repressed unbinding**

Our model of Zur oligomerization at TB sites was based partly on the structure of two holo-Zur dimers bound to a cognate DNA, which showed two inter-dimer D49–R52 salt bridges. To probe the role of these salt bridges in Zur oligomerization, we made the D49A mutation, known to disrupt the interactions. For apo-Zur, the resulting mutant Zur<sup>apo</sup><sub>mE</sub>,D49A still exhibits the biphasic unbinding behavior, however the minimum of the apparent unbinding rate constant <i>k<sub>d</sub></i> shifted to a higher cellular protein concentration (Fig. 2d, right). Its <i>K<sub>m</sub></i> is 16.2 ± 7.5 nM, three times larger than that of Zur<sup>apo</sup> <sub>mE</sub> (Table 1), indicating a weakened oligomerization affinity and thus a significant role of these salt bridges.

More strikingly, for Zur<sup>mE</sup> <sub>Zn</sub>, which only showed facilitated unbinding (Fig. 2d, left), the resulting mutant Zur<sup>mE</sup> <sub>Zn</sub>,D49A clearly shows biphasic unbinding with <i>K<sub>m</sub></i> = 3.2 ± 1.9 nM (Fig. 2d, right; Table 1). Therefore, holo-Zur also possesses repressed unbinding kinetics — it was invisible for Zur<sup>mE</sup> <sub>Zn</sub> likely because its <i>K<sub>m</sub></i> is smaller than the low limit of accessible cellular protein concentrations (~3 nM), but emerges after the D49A mutation, which further supports the importance of the salt bridges in Zur oligomerization and repressed unbinding behaviors.

**DISCUSSION**

We have uncovered that the Fur-family Zn<sup>2+</sup>-sensing transcription regulator Zur exhibits two unusual behaviors that challenge conventional paradigms of regulator-chromosome interactions. First, apo-Zur, the non-repressor form and a long-presumed non-DNA binder, can actually bind to chromosome tightly, likely at different locations from the consensus sequence recognized by holo-Zur, the repressor form. This tight chromosome binding by apo-Zur challenges the paradigm of regulator on-off model for transcription repression (or activation)~1,2. Second, the unbinding kinetics of both apo- and holo-Zur not only exhibit facilitated unbinding, a newly discovered phenomenon for a few DNA-binding proteins~6,7,9,28, but also show repressed unbinding, a *first-of-its-kind* phenomenon that likely results from Zur oligomerization on chromosome, facilitated by inter-dimer salt bridges. Overall, Zur has biphasic unbinding kinetics from chromosome with increasing cellular protein concentrations, which challenges the paradigm of protein unbinding being typically unimolecular processes whose first-order rate constants do not depend on the protein concentration.

To probe whether the biphasic unbinding of Zur occurs within the physiological cellular protein concentrations, we quantified cellular Zur<sup>mE</sup> concentration when it is encoded only at the chromosomal locus (Fig. 4a). In minimal medium without Zn stress, the cellular Zur<sup>mE</sup>, which is mostly in the apo-form, ranges from ~24 to 108 nM (mean = 50 ± 14 nM), within which apo-Zur unbinding from TB sites is in the repressed unbinding regime and slows down by ~42% from the lowest to the highest protein concentration (Fig. 4b). When stressed by 20 μM Zn<sup>2+</sup>, the cellular Zur<sup>mE</sup>, now mostly in the holo-form, ranges from ~26 to 124 nM (mean = 63 ± 20 nM), reflecting an average of ~28% protein concentration increase induced by Zn stress. In this protein concentration range, holo-Zur is already in the facilitated unbinding regime, and its unbinding rate from a recognition site can increase by ~36% (Fig. 4b).

Within the physiological protein concentration range, the opposite dependences of unbinding kinetics on the cellular protein concentration between apo- and holo-Zur could provide functional advantages for an *E. coli* cell to repress or de-repress Zn uptake genes. When cell encounters environmental Zn stress that demands strong repression of Zn uptake, the cellular concentration of Zur swings upward and it becomes dominantly in the holo-repressor form. The unbinding of holo-repressor from recognition sites could be facilitated by its increasing concentration (Fig. 5a), but the facilitated unbinding via direct substitution by another holo-repressor has no functional consequence while
facilitated unbinding via assisted dissociation will be immediately compensated by a rebinding of a
holo-repressor (the rebinding would occur within ~0.014 s; Supplementary Note 7). For those cellular
Zur in the apo non-repressor form, its unbinding from DNA slows down, keeping them longer (i.e.,
stored) at non-consensus chromosomal sites (Fig. 5b). On the other hand, when cell transitions to a Zn-
deficient environment that demands derepression of Zn uptake, the cellular Zur protein concentration
goes down. Here unbinding of the holo-repressor would be slower (Fig. 5c), which is undesirable for
derepression, while the unbinding of the apo-form would become faster, releasing them from the non-
consensus “storage” sites on the chromosome into the cytosol (Fig. 5d). If the cytosolic apo-Zur could
possibly facilitate the unbinding of holo-Zur from promoter recognition sites (e.g., through assisted
dissociation), it would give a more facile transition to derepression. To support this possibility, we
measured the apparent unbinding rate constant $k_d$ for chromosomally encoded Zur$^{mE}$ in cells that
contains a plasmid encoding an untagged Zur$^{apo}$ mutant (i.e., C88S). When the expression of this Zur$^{apo}$
mutant is induced, $k_d$ of Zur$^{mE}$ increases by ~28% at any cellular Zur$^{mE}$ concentration (Fig. 4b, green
vs. red points), indicating that apo-Zur can indeed facilitate the unbinding of holo-Zur from recognition
sites (Fig. 5c).

Multivalent contacts with DNA, which underlie the facilitated unbinding, and salt-bridge
interactions between proteins, which underlie Zur oligomerization and its repressed unbinding, are both
common for protein-DNA and protein-protein interactions, respectively$^5, 7, 10, 28, 36$. Therefore, the
biphasic unbinding behavior from DNA discovered here for Zur could be broadly relevant to many
other proteins in gene regulation.

METHODS

Bacterial strains and sample preparation

All strains were derived from the E.coli BW25113 strain as detailed in Supplementary Note 1.
Zur$^{mE}$ was either encoded at its chromosomal locus via lambda-red homologous recombination$^{37}$ or in
a pBAD24 plasmid in a Δzur deletion strain$^{38}$. Mutant forms of Zur (Zur$^{apo}$, Zur$^{D49A}$, or Zur$^{apo, D49A}$)
were generated via site-directed mutagenesis in pBAD24, which was introduced into the Δzur strain.

All cell imaging experiments were done at room temperature in M9 medium supplemented with
amino acids, vitamins, and 0.4% glycerol. 20 μM ZnSO$_4$ was used for Zn stress conditions. The cells
were immobilized on an agarose pad in a sample chamber. Details in Supplementary Note 3.

SMT and SCQPC

SMT and SCQPC were performed on an inverted fluorescence microscope, as reported$^{11}$
(Supplementary Note 3). For SMT, inclined epi-illuminated 405 nm and 561 nm lasers photoconverted
and excited single mEos3.2 molecules, respectively. 561 nm excitation-imaging were in stroboscopic
mode, with 4 ms laser excitation pulses separated by 40 ms time lapse, synchronized with the camera
exposure, so that the mobile proteins still appear as diffraction-limited spots. A custom-written
MATLAB software was used to identify diffraction-limited fluorescence spots and fit them with two-
dimensional Gaussian functions, giving ~20 nm localization precision$^{11, 36}$. Time trajectories of positions
and displacement length $r$ between adjacent images were then extracted.

SCQPC was performed after SMT. The remaining proteins were firstly photoconverted to the
red form by a long 405 nm laser illumination. The total cell red fluorescence was then imaged by the
561 nm laser to determine the protein copy number, provided the average fluorescence of a single
mEos3.2 from the earlier SMT. The photoconversion efficiency of mEos3.2$^{40}$ and dimeric state of Zur
were accounted for. Cell volumes were determined by fitting their optical transmission image contours
with the model geometry of a cylinder with two hemispherical caps.

Resolution of diffusion states
The effective diffusion constants and the fractional populations of diffusion states were extracted by analyzing the CDF of displacement length \( r \) per time-lapse \( (T_{tl} = 40 \text{ ms}) \), using a linear combination of three diffusion terms of CDF, as reported\(^1\) (Equation (3)). Each term is from a 2-D Brownian diffusion model\(^{18, 41, 42}\), which was regularly used to analyze SMT results of proteins in bacterial and mammalian cells\(^{18, 21, 42-46}\) (model justification in Supplementary Note 4).}

\[
\text{CDF}(r) = A_{\text{FD}} \left(1 - \exp \left(-\frac{r^2}{4D_{\text{FD}}T_{tl}}\right) \right) + A_{\text{NB}} \left(1 - \exp \left(-\frac{r^2}{4D_{\text{NB}}T_{tl}}\right) \right) + (1 - A_{\text{FD}} - A_{\text{NB}}) \left(1 - \exp \left(-\frac{r^2}{4D_{\text{TB}}T_{tl}}\right) \right)
\]

We globally fitted the CDFs across groups of cells of different cellular protein concentrations, in which the diffusion constants \((D's)\) of respective diffusion states were shared but their fractional populations \((A's)\) were allowed to vary. Three terms were always the minimal number of diffusion states to satisfactorily fit the CDF (details in Supplementary Note 4 and Supplementary Tables 4-5).

Note these diffusion constant values are not the intrinsic ones, as they are influenced by the cell confinement effect\(^7\), which decreases the magnitude of the apparent diffusion constant, and by the time-lapse effect of imaging, where longer time lapse gives apparently smaller diffusion constants; both of these effects are most significant on the FD state, less on the NB state, and negligible on the TB state, and were evaluated quantitatively in a previous study of metal-responsive transcription regulators of a different family\(^11\).

**Determination and analysis of \( k_d \)**

A three-state (FD, NB, and TB state) kinetic model, including the interconversion between states and photobleaching/blinking times (Fig. 2c) at chromosomal TB sites to extract the apparent unbinding rate constant \( k_d \). The respective residence time distribution functions \( \phi(\tau) \) for the FD, NB, and TB states with given diffusion constants \((D's)\), the unbinding rate constant from the NB state \( k_2 \), and photobleaching/blinking rate constant \( k_{bl} \) were derived to fit the \( \tau \) distribution with the overall distribution function \( \phi_{\text{all}}(\tau) \) (Eq. (4); Supplementary Note 5).

\[
\phi_{\text{all}}(\tau) = A_{\text{FD}}\phi_{\text{FD}}(\tau) + A_{\text{NB}}\phi_{\text{NB}}(\tau) + A_{\text{TB}}\phi_{\text{TB}}(\tau)
\]

\[
\phi_{\text{FD}}(\tau) = \frac{r_0^2}{4D_{\text{FD}}\tau^2} \exp \left(-\frac{r_0^2}{4D_{\text{FD}}\tau} \right) + k_{\text{eff}}^{\text{FD}} \left(1 - \exp \left(-\frac{r_0^2}{4D_{\text{FD}}\tau} \right) \right) \exp (-k_{\text{eff}}^{\text{FD}}\tau)
\]

\[
\phi_{\text{NB}}(\tau) = \frac{r_0^2}{4D_{\text{NB}}\tau^2} \exp \left(-\frac{r_0^2}{4D_{\text{NB}}\tau} \right) + k_{\text{eff}}^{\text{NB}} \left(1 - \exp \left(-\frac{r_0^2}{4D_{\text{NB}}\tau} \right) \right) \exp (-k_{\text{eff}}^{\text{NB}}\tau)
\]

\[
\phi_{\text{TB}}(\tau) = k_{\text{eff}}^{\text{TB}} \exp (-k_{\text{eff}}^{\text{TB}}\tau)
\]

Here \( k_{\text{eff}}^{\text{FD}} = k_{\text{bl}} \frac{T_{\text{int}}}{T_{\text{tl}}} \), \( k_{\text{eff}}^{\text{NB}} = k_{\text{bl}} \frac{T_{\text{int}}}{T_{\text{tl}}} + k_2 \), \( k_{\text{eff}}^{\text{TB}} = k_{\text{bl}} \frac{T_{\text{int}}}{T_{\text{tl}}} + k_d \), and \( A_i \) is the fractional population of \( i^{th} \)-state.

The dependence of \( k_d \) on the cellular free diffusing protein concentration \([P]_{\text{int}}\) was analyzed with Eq. (2), containing three terms representing spontaneous, repressed, and facilitated unbinding with the corresponding rate constants \( k_0^{\text{off}}, k_r \), and \( k_f \), respectively (derivation in Supplementary Note 6).

**Analysis of relative populations**

The same three-state kinetic model (Fig. 2c) was used to analyze the relative populations of FD, NB, and TB states of Zur across all cellular protein concentrations.
Oligomerization/deoligomerization of Zur at a TB site was modeled as 1-D sequential binding/unbinding, analogous to the Brunauer-Emmett-Teller multilayer-adsorption theory\textsuperscript{48} but with a limited number \( n_s \) of binding site and merely one binding rate constant \( k_1 \) (see Supplementary Note 7 for detailed derivation). Quasi-equilibrium approximation of interconversion among states was used, which approximates that the timescale of interconversion between states (~ms) are much shorter than the experimental imaging time (~hours). The kinetic parameters are then related to the relative concentrations of the proteins at three diffusion states.

\[
\left[ \frac{[PD]_{TB}}{[P]_{FD}} \right] = \frac{k_1 [D_0]_{TB}}{k_3} \frac{\partial \ln F_{TB\rightarrow FD}(x_{TB\rightarrow FD})}{\partial x_{TB\rightarrow FD}}
\]

(8)

\[
\left[ \frac{[PD]_{TB}}{[P]_{NB}} \right] = \frac{k_3 [D_0]_{TB}}{k_3 ([D_0]_{NB} - [PD]_{NB})} \frac{\partial \ln F_{TB\rightarrow NB}(x_{TB\rightarrow NB})}{\partial x_{TB\rightarrow NB}}
\]

(9)

\[
\left[ \frac{[PD]_{NB}}{[P]_{FD}} \right] = \frac{k_2 [D_0]_{NB}}{k_2 + k_2 [P]_{FD}}
\]

(10)

Here \([P]_{FD}, [PD]_{NB}, \) and \([PD]_{TB}\) are the cellular protein concentrations of FD, NB, and TB states, respectively. \( F_{TB\rightarrow j}(x_{TB\rightarrow j}) \equiv \sum_{i=0}^{n_s} x_{TB\rightarrow j} \cdot j \in \{FD, NB\} \), where \( x_{TB\rightarrow FD} \equiv \frac{k_1}{k_3} [P]_{FD} \) and \( x_{TB\rightarrow NB} \equiv \frac{k_3}{k_3 ([D_0]_{NB} - [PD]_{NB})} [PD]_{TB} \) and \( [D_0]_{TB} \) and \( [D_0]_{NB} \) are the effective cellular concentrations of TB and NB sites, respectively. Thermodynamic quantities such as the dissociation constant of TB \( (K_d1 = \frac{k_2}{k_1}) \) and NB \( (K_d2 = \frac{k_2}{k_2}) \) were also determined from this analysis.

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Author contributions

W.J. and P.C. designed research; W.J. performed experiments, derived theory, coded software, and analyzed data; W.J. and P.C. discussed the results and wrote the manuscript.

Competing interests

The authors declare no competing interest.

Additional information

Supplementary information is available for this paper.
Fig. 1 | SMT of Zur in living cells. a. Left: exemplary fluorescence image of a single Zur\textsuperscript{apo} protein in a live E. coli cell overlaid with its position trajectory (solid line). Right: overlay of many trajectories. Dash lines: cell boundary. b. Histogram of displacement length $r$ per time-lapse (40 ms) of > 1,400 tracked Zur\textsuperscript{apo} proteins at 124 ± 15 nM. Solid lines: the overall fitted distribution (black), and the resolved FD (blue), NB (green), and TB (red) diffusion states (Supplementary Note 4). Vertical dashed line: $r_0 = 0.2 \mu m$ for extracting residence times as in Fig. 2a. c. Cumulative-distribution-function (CDF) of $r$ (plotted against $r^2/(4 T_0)$) as in b. Lines: overall fit (Eq. (3)) and three resolved diffusion states with effective diffusion constants (and fractional populations): $D_{FD} = 5.0 \pm 0.5 \mu m^2 s^{-1}$ (21.7 ± 0.4%), $D_{NB} = 0.8 \pm 0.05 \mu m^2 s^{-1}$ (48.8 ± 0.4%), and $D_{TB} = 0.040 \pm 0.003 \mu m^2 s^{-1}$ (30.1 ± 0.5%). d. Fractional populations of FD, NB, and TB states for Zur\textsuperscript{apo} (half-solid squares) and Zur\textsuperscript{Zn} (half-solid circles) vs. the cellular protein concentrations.
**Fig. 2 | Biphasic unbinding kinetics of Zur from TB sites on chromosome.**

a. Time trajectory of displacement length $r$ per time-lapse from a single Zur$_{apo}^m$E protein. Two microscopic residence time $\tau$ shown in gray shades; dashed horizontal line: displacement threshold $r_o = 0.2 \mu$m (vertical dashed line in Fig. 1b).  

b. Histogram of $\tau$ for Zur$_{apo}^m$E at the cellular concentration of 124 ± 15 nM. Black line: fitting with Eq. (4). Contributions of the three diffusion states are plotted, as color-coded in Fig. 1b-c.  

c. Three-state model of a single Zur protein interacting with DNA in a cell. $k$'s are the rate constants.  

d. Protein-concentration-dependent $k_d$ for Zur$_{apo}^m$E and Zur$_{Zn}^m$E (left) and their corresponding D49A salt-bridge mutants (right). Bottom/top axis refers to free/cellular protein concentration, respectively. Lines are fits with Eq. (2). All error bars are s.d.  

e. Schematic molecular mechanisms for biphasic unbinding of Zur from a TB site. A bound Zur protein (dark blue) within an oligomer on DNA can unbind following either a repressed pathway (top) due to the presence of $(n-1)$ proteins nearby or a facilitated pathway (bottom) upon binding another protein (green) to form an intermediate ternary complex, which then proceeds through direct substitution or assist dissociation pathway. Black dashed lines denote salt-bridge interactions.
Fig. 3 | Spatial analysis of Zur’s residence sites. **a,** Normalized pair-wise distance distributions (PWD) of residence sites for Zur<sub>apo</sub><sup>mE</sup> and for simulated random sites in the cell (top), and the difference of Zur<sub>apo</sub><sup>mE</sup> from simulation (bottom). **b,** Fraction of residence sites within a radius threshold $R$ (= 100 nm, left axis) for Zur<sub>apo</sub><sup>mE</sup> and for simulated random sites as a function of cellular protein concentration. Their ratio (Zur<sub>apo</sub><sup>mE</sup> vs. simulation) is plotted against the right axis. **c,** Dependence of the average ratio in **b** across all protein concentrations as a function of the radius threshold $R$ for Zur<sub>apo</sub><sup>mE</sup> and Zur<sub>Zn</sub><sup>mE</sup>.

Fig. 4 | Zur behaviors within the physiological range of cellular protein concentrations. **a,** Distribution of the chromosomally expressed Zur<sub>apo</sub><sup>mE</sup> concentration in the cell with (+) and without (−) Zn stress in the medium. **b,** Dependence of $k_d$ on the protein concentration in the cell for Zur<sub>apo</sub><sup>mE</sup>, Zur<sub>Zn</sub><sup>mE</sup>, and for Zur<sub>Zn</sub><sup>mE</sup> together with a plasmid expressing Zur<sub>apo</sub> (i.e. Zur<sub>Zn</sub><sup>mE,Chr</sup>/Zur<sub>apo</sub>) when the mE-tagged Zur is only encoded on the chromosome. The blue circles and red squares for Zur<sub>apo</sub><sup>mE</sup> and Zur<sub>Zn</sub><sup>mE</sup> are part of data in Fig. 2d (left).
Fig. 5 | Functional model of holo- and apo-Zur unbinding behaviors in E.coli upon encountering zinc stress or deficiency. Upon zinc stress, unbinding of holo-Zur from operator site is facilitated (a) while that of apo-Zur from storage site is repressed (b) due to increase in cellular protein concentration. Upon zinc deficiency, the facilitated unbinding of holo-Zur is attenuated (c) while the unbinding of apo-Zur is less repressed (d) due to decrease in cellular protein concentration. Released apo-Zur into cytosol could facilitate holo-Zur to unbind (e), helping transition to de-repression of zinc uptake.

Table 1 | Kinetic and thermodynamic parameters for Zur-DNA interaction in E.coli cells

| Parameter | Zur<sup>off</sup> | Zur<sup>apo</sup> | Zur<sup>Zn</sup> | Zur<sup>apo, D49A</sup> | Zur<sup>Zn, D49A</sup> |
|-----------|-----------------|-----------------|-----------------|----------------------|---------------------|
| k<sub>i</sub> (nM<sup>-1</sup> s<sup>-1</sup>) | 1.90 ± 0.17 | 1.84 ± 0.20 | 1.10 ± 0.18 | 1.61 ± 0.58 | 1.30 ± 0.19 |
| k<sub>off</sub> (s<sup>-1</sup>) | 25 ± 12 | 22 ± 21 | 5.4 ± 0.6 | 22.1 ± 1.5 | 36 ± 41 |
| k<sub>i</sub> (s<sup>-1</sup>) | 16 ± 11 | 12 ± 20 | n/o b | 20.8 ± 1.3 | 27 ± 40 |
| k<sub>f</sub> (nM<sup>-1</sup> s<sup>-1</sup>) | 0.028 ± 0.005 | 0.044 ± 0.007 | 0.026 ± 0.033 | 0.049 ± 0.014 | 0.062 ± 0.010 |
| K<sub>i</sub> (nM) | 6.0 ± 4.0 | 4.9 ± 7.3 | n/o b | 16.2 ± 7.5 | 3.2 ± 1.9 |
| K<sub>A</sub> = k<sub>i</sub>/k<sub>i</sub> (nM) | 12.9 ± 6.2 | 11.7 ± 11.2 | 4.9 ± 1.2 | 13.7 ± 5.0 | 28 ± 20 |
| K<sub>A</sub> = k<sub>off</sub>/k<sub>i</sub> (nM) | 417 ± 35 | 348 ± 84 | 534 ± 148 | 209 ± 69 | 532 ± 134 |
| K<sub>A</sub> = k<sub>f</sub>/k<sub>i</sub> (nM) | 0.011 ± 0.002 | 0.023 ± 0.007 | 0.022 ± 0.023 | 0.032 ± 0.062 | 0.008 ± 0.006 |
| [D]<sub>0</sub> (nM) | 1144 ± 84 | 961 ± 205 | 1201 ± 287 | 858 ± 230 | 1538 ± 353 |
| [D]<sub>0</sub> (μM) | 42.56 ± 0.94 | 34.3 ± 3.2 | 54 ± 14 | 31.6 ± 5.1 | 38.8 ± 3.8 |

*<sup>a</sup> n<sub>o</sub> = 5 was used in fitting.
*<sup>b</sup> Not observed