Balancing between affinity and speed in target DNA search by zinc-finger proteins via modulation of dynamic conformational ensemble

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Although engineering of transcription factors and DNA-modifying enzymes has drawn substantial attention for artificial gene regulation and genome editing, most efforts focus on affinity and specificity of the DNA-binding proteins, typically overlooking the kinetic properties of these proteins. However, a simplistic pursuit of high affinity can lead to kinetically deficient proteins that spend too much time at nonspecific sites before reaching their targets on DNA. We demonstrate that structural dynamic knowledge of the DNA-scanning process allows for kinetically and thermodynamically balanced engineering of DNA-binding proteins. Our current study of the zinc-finger protein Egr-1 (also known as Zif268) and its nuclease derivatives reveals kinetic and thermodynamic roles of the dynamic conformational equilibrium between two modes during the DNA-scanning process: one mode suitable for search and the other for recognition. By mutagenesis, we were able to shift this equilibrium, as confirmed by NMR spectroscopy. Using fluorescence and biochemical assays as well as computational simulations, we analyzed how the shifts of the conformational equilibrium influence binding affinity, target search kinetics, and efficiency in displacing other proteins from the target sites. A shift toward the recognition mode caused an increase in affinity for DNA and a decrease in search efficiency. In contrast, a shift toward the search mode caused a decrease in affinity and an increase in search efficiency. This accelerated site-specific DNA cleavage by the zinc-finger nuclease, without enhancing off-target cleavage. Our study shows that appropriate modulation of the dynamic conformational ensemble can greatly improve zinc-finger technology, which has used Egr-1 (Zif268) as a major scaffold for engineering.

Significance

Many transcription factors and DNA repair/modifying enzymes must first locate the target sites through stochastic scanning of DNA in the vast presence of nonspecific sites. In this work, we demonstrate that target search by these proteins can be accelerated via engineering based on structural dynamic knowledge of the DNA-scanning process. Our biophysical data for the Egr-1 zinc-finger protein and its nuclease derivatives reveal kinetic and thermodynamic roles of the conformational equilibrium between two modes in the DNA-scanning process: one suitable for search and the other for recognition. We found that optimizing the balance between the search and recognition modes improves efficiency in zinc-finger proteins’ target search. This finding can help advance zinc-finger technology for artificial gene regulation and genome editing.

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relatively short lifetime (~0.5–1 h) (29). Our previous study suggested the presence of dynamic transitions between conformationally different states of Egr-1 during the DNA-scanning process: one allowing for rapid translocation on DNA (the search mode) and the other allowing for highly specific recognition (the recognition mode) (28). This finding is consistent with the conformational switch model (19, 25, 31–33), which some theoreticians proposed to resolve the speed-stability paradox 

To test the abovementioned hypothesis, we published online August 31, 2015 PNAS | R 15 B 1 cause local dissociation of and C E5143 original specificity to the target

Our previous NMR and computational investigations on non-specific and specific DNA complexes of Egr-1 suggested that, while originally DNA, Egr-1 undergoes the dynamic conformational ensemble in the DNA-scanning process. Our mutagenesis data on the Egr-1 ZF DNA-binding domain (ZF-DBD) and its ZFN derivative reveal the kinetic and thermodynamic roles of the search and recognition modes in target search. This work highlights the importance of structural dynamic knowledge of the DNA-scanning process at the molecular and atomic levels in engineering of transcription factors and DNA-modifying enzymes. The insight from this work should be applicable to many artificial ZF proteins because Egr-1 (Zif268) has been used as a major scaffold for ZF technology.

**Results**

Our previous NMR and computational investigations on non-specific and specific DNA complexes of Egr-1 suggested that, while originally DNA, Egr-1 undergoes the dynamic conformational ensemble between the search and recognition modes (28). We consider the conformational ensemble involving three major states in the DNA-scanning process as shown in Fig. L1. In the recognition mode, all three ZFs are bound to DNA. In the search mode, a ZF is locally dissociated from DNA and the other two ZFs are nonspecifically bound to DNA (two different search modes, a and b, are defined as shown in Fig. L1). Our previous studies suggested that the interdomain dynamics could harness the rapid conformational switch between the search and recognition modes, thereby allowing Egr-1 to overcome the speed-stability paradox (24, 28). Our current work is based on a hypothesis that the balance between the search and recognition modes is a major determinant of the kinetic efficiency in target DNA search by ZF proteins.

**Shifting the Conformational Equilibrium During DNA Scanning via Ion-Pair Engineering.** To test the abovementioned hypothesis, we shifted the equilibrium between the search and recognition modes and examined the kinetic consequences of the shifts. For this strategy, we used mutagenesis to modulate (i) nonspecific interactions between a ZF and DNA and (ii) interdomain interactions between ZFs. Weakening these interactions can shift the conformational equilibrium toward the search mode, whereas strengthening these interactions can shift it toward the recognition mode. To maintain the proteins’ original specificity to the target sequence, the mutations are limited to residues that are not in contact with bases of the DNA. Previously, we found that weaker ZF-DNA and ZF-ZF interactions cause local dissociation of ZF1 from DNA, while ZF2 and ZF3 are nonspecifically bound to DNA (28). Based on this finding, we modulated ZF-DNA and ZF-ZF interactions via engineering of the ion pairs at the ZF-DNA and ZF-ZF interfaces (Fig. 1B). The ZF-DNA interactions were modulated via mutations of T23 and K79. Although T23 and K79 are located at corresponding positions of ZF1 and ZF3, respectively, K79 of ZF3 forms an intermolecular ion pair with DNA phosphate and T23 of ZF1 does not. Therefore, the intermolecular ion pair with DNA can be added or removed from the ZF-DNA interfaces by the T23K or K79T mutation. We confirmed it by using an NMR method that was recently developed in our laboratory (36) for measuring scalar coupling between lysine side-chain 15N and DNA 31P nuclei across a hydrogen bond (Fig. S1). The ZF-ZF interactions were modulated via mutations of Q32 and E60. The ZF2-ZF3 interface involves an ion pair of R55 and E60, stabilizing the interface via the hydrogen bonds and electrostatic interactions (37), whereas the ZF1-ZF2 interface lacks such ionic interactions between corresponding residues Q32 and R27 (Fig. 1B).

**Fig. 1.** Modulation of the search and recognition modes during DNA scanning by the Egr-1 ZF-DBD. (A) Search and recognition modes. (B) Mutation sites used to modulate the balance between the search and recognition modes. (C) Backbone amide 15N R1 relaxation rates measured at the 1H frequency of 800 MHz for the nonspecific complexes of the wild-type and mutant proteins of Egr-1 ZF-DBD with 28-bp DNA duplex, dGTACCGATTGCAGATTCCGAACC-TTCAG, which contains neither specific nor semispecific sequences.
Therefore, the ZF-ZF interfaces can be stabilized or destabilized by the Q32E or E60Q mutation.

By combining these mutations, we shifted the equilibrium between the search and recognition modes. We generated the T23K/Q32E double mutant (referred to as “type 1”) to shift toward the recognition mode, the T23K/Q32E/E60Q/K79T quadruple mutant to shift toward the search mode (“type 2”), and the E60Q/K79T mutant (“type 3”) to enhance both the search modes a and b. Our previous NMR 15N relaxation, residual dipolar coupling (RDC), and paramagnetic relaxation enhancement (PRE) data clearly indicated that ZF1 is mainly dissociated from DNA, whereas ZF2 and ZF3 remain bound to DNA in the nonspecific DNA complex of wild-type (“type 0”) Egr-1 ZF-DBD (28). Because the most remarkable signature for this local dissociation was the 15N R1 relaxation data, we used 15N R1 relaxation to investigate the impact of these mutations on the conformational equilibrium between the search and recognition modes. Local dissociation of a ZF in the search modes should shorten the domain’s rotational correlation time because of the independent tumbling and thereby increases its overall 15N R1 relaxation rates. The nonspecific DNA complexes of the four constructs showed clearly different 15N R1 profiles, as shown in Fig. 1C, ZFI in the type 0 and type 3 complexes showed clearly elevated 15N R1 rates, suggesting a major presence of the search mode a; ZF3 of the type 2 and type 3 complexes exhibited elevated 15N R1 rates, suggesting a major presence of the search mode b. No ZF showed elevated 15N R1 rates for the type 1 complex, suggesting a major presence of the recognition mode during DNA scanning.

In contrast, these mutations did not affect the 15N R1 relaxation profiles for the specific complexes with the target DNA (Fig. S2), indicating that the mutant proteins are also locked in the recognition mode when they are bound to the target. Although one might expect that the mutations would impact the interdomain dynamics in the free state, our 15N R1 relaxation data for the Egr-1 ZF-DBDs in the free state showed otherwise (Fig. S3). This was also supported by small-angle X-ray scattering (SAXS) data (Fig. S3B). The ZF-ZF packing in the free state might be too loose because of the lack of the C-terminal capping of ZF3 α-helices in the absence of DNA (38). Our NMR data indicate that the mutations affect the interdomain dynamics only for nonspecific complexes and successfully shift the equilibrium between the search and recognition modes during the DNA-scanning process.

**Trade-Off Between Search Efficiency and Binding Affinity.** Taking advantage of these shifts, we examined the roles of the search and recognition modes during DNA scanning by Egr-1 ZF-DBD. Using fluorescence methods (26, 27), we compared the four types of Egr-1 ZF-DBDs in terms of the binding affinities and search kinetics. Fig. 2A shows binding affinities measured for specific and nonspecific DNA duplexes (also see Table S1 for the dissociation constants). The type 1 construct, which is primarily in the recognition mode during DNA scanning, exhibited the strongest affinities among the four constructs. The type 3 construct, for which both search modes a and b are enhanced, exhibited the weakest affinities. We also compared the target search kinetics by the stopped-flow fluorescence assay in which the protein was mixed at 150 mM KCl with a solution of fluorescence-labeled DNA (2.5 nM) and sonicated calf thymus DNA (56 μM base pairs) as competitors in large excess (Fig. 2B). By monoeponential fitting to the fluorescence time-course data, we measured apparent pseudo-first-order rate constants k\text{app} at distinct concentrations of the protein, from which the apparent second-order rate constants k\text{a} for the target association were determined (27) (Fig. 2B). The protein-concentration dependence data ensured that contribution of the backward first-order process to k\text{app} (26, 27, 39) was negligible for each construct of Egr-1 ZF-DBD. The target search by the type 3 construct was fastest among the four constructs, and that by the type 1 construct was slowest. A shift of the conformational equilibrium toward the recognition mode resulted in higher affinities for both specific and nonspecific DNA sites but a lower efficiency in target search. This represents a kinetic defect in target search at the expense of high affinity, which renders the protein trapped on nonspecific DNA before reaching the target.

In contrast, a shift toward the search modes resulted in lower affinities for DNA but a higher efficiency in target search (Figs. 2A and B). As a result, the target search by the type 3 construct was 15-fold faster than that by type 1 construct in the stopped-flow assays.

**Impact on Protein Sliding, Dissociation, and Intersegment Transfer.**
To understand how the shifts toward the search modes accelerate the target search, we investigated kinetics of translocation of the type 0-3 Egr-1 ZF-DBD proteins on nonspecific DNA. For this purpose, we used the stopped-flow-based assays together with a discrete-state stochastic kinetic model for protein translocation on DNA, which we previously developed (26, 27). In these assays, the time courses of the FAM fluorescence were...
monitored upon mixing the Egr-1 ZF-DBD protein with FAM-labeled probe DNA (2.5 nM) and nonspecific 28-bp DNA (1–16 μM) as competitor in large excess. The probe DNA duplexes used were 33, 48, 63, 88, 113, and 143 bp (sequences shown in ref. 26), each containing a single target sequence (GGGTGGGCG) near the 5′-terminal FAM probe. From these kinetic data, we determined the 1D diffusion coefficients $D_i$ for sliding on DNA, the rate constants $k_{IT,N}$ for dissociation from a nonspecific site on DNA, and the rate constants $k_{IT,N}$ for intersegment transfer between two nonspecific sites on distinct DNA duplexes (Fig. 2C). The data for the four constructs show that the shifts toward the search modes causes the faster dissociation, sliding, and intersegment transfer, which collectively accelerates the target search; the shift toward the recognition mode causes the opposite effects. These data suggest that the balance between the search and recognition modes during DNA scanning is an important determinant of the kinetic and thermodynamic properties of the ZF-DBD.

Computational Analysis of the Impact on Intersegment Transfer. Our previous studies on the target DNA search process of Egr-1 demonstrated the importance of intersegment transfer (also known as direct transfer) between two nonspecific DNA sites (26–28). Efficient intersegment transfer was also experimentally observed for some other proteins (40–44). To gain further insight into the roles of the search modes, we also conducted coarse-grained molecular dynamics simulations for investigating the intersegment transfer processes. This computational approach is well established and was previously used for several proteins, including Egr-1 (28, 45–47). In these simulations, two B-form DNA duplexes (100 bp) were placed in parallel with an interaxial distance of 60 Å, and we monitored how protein translates on DNA for the four types of Egr-1 ZF-DBD constructs. The trajectories of these simulations showed that intersegment transfer can occur via the “monkey-bar” mechanism (45, 47) in two distinct manners: one is via ZF1, which captures the second DNA, whereas the other domains remain on the initial DNA (Fig. 3A); and the other via ZF3 (Fig. 3B). Each case involves an intermediate with a protein molecule transiently bridging two DNA duplexes via different ZFs. For the type 0 and type 3 constructs, where the search mode $a$ is enhanced, intersegment transfer via ZF1 occurred more frequently than for the type 1 and type 2 constructs (Fig. 3C). Likewise, for the type 2 and type 3 constructs, for which the search mode $b$ is enhanced, intersegment transfer via ZF3 occurred more frequently than for the type 0 and type 1 constructs (Fig. 3D). The coarse-grained simulations show that the different Egr-1 ZF-DBD constructs have different linear diffusion coefficients of the following order $D_3(\text{type } 3) > D_1(\text{type } 0) > D_2(\text{type } 2) > D_1(\text{type } 1)$. These computational data suggest that the shifts toward the search modes $a$ and $b$ enhance intersegment transfers via ZF1 and ZF3, respectively, by increasing the probability of forming intermediates with a protein molecule bridging two DNA duplexes via multiple ZFs.

**Improvement of ZFN via Optimizing Balance Between the Search/Recognition Modes.** We examined whether these shifts of the conformational equilibrium also influence the kinetics of sequence-specific DNA cleavage by ZFNs. ZFNs are fusion proteins of a ZF-DBD with the FokI ND, and cleave double-stranded DNA in a sequence-specific manner via dimer formation of FokI ND at two target sites separated by several base pairs (4, 5). We constructed ZFNs comprising type 0, 1, 2, or 3 Egr-1 ZF-DBD and FokI ND and compared the kinetics of their sequence-specific DNA cleavage at physiological ionic strength (150 mM KCl). In our in vitro kinetic assays of ZFN, we monitored the site-specific cleavage of a 4-kbp linear DNA (1 nM; containing a target cleavage site for Egr-1 ZFN) in the presence of sonicated calf thymus DNA (56 μM base pairs) in large excess. This reaction produces 3- and 1-kbp DNA fragments (Fig. 4A; also see Fig. S4). With FokI ND’s catalytic rate constant ($k_{\text{cat}}$) taken into consideration (48), the target search process should be the rate-limiting step for the sequence-specific DNA-cleavage reaction under our experimental conditions. Fig. 4B shows the observed rates for the sequence-specific DNA cleavage by 5 nM Egr-1 ZFNs in this assay. Despite its highest affinity, the type 1 Egr-1 ZFN exhibited the slowest DNA cleavage among the four constructs. In contrast, despite its lowest affinity, the type 3 Egr-1 ZFN exhibited the fastest DNA cleavage. The site-specific DNA cleavage by the type 3 Egr-1 ZFN was 14-fold faster than that by the type 1 Egr-1 ZFN, and these results are consistent with the abovementioned results from the stopped-flow assays for the Egr-1 ZF-DBDs. At a higher concentration of ZFN and a longer reaction time, we observed off-target DNA cleavage but virtually to the same degree for all four ZFN constructs (Fig. S5). Although high affinities are pursued in ZF engineering (1–3), our data show that a high affinity does not warrant kinetically optimal ZF proteins because they can be trapped at nonspecific sites more strongly, which hampers target search. This is also consistent with some previous reports that ZFNs containing a larger number of ZFs are less active in sequence-specific DNA cleavage (13, 14).

**Target Search Requiring Displacement of Other Proteins.** The results shown above were obtained from assays in the absence of other proteins. In the nuclei, however, many different proteins compete for particular sites on DNA. For example, upon cellular stimuli such as neuronal signals and vascular stress, the inducible transcription factor Egr-1 displaces the constitutive transcription factor Sp1 (which binds to DNA via 3 ZFs) from their overlapping target sites in some gene promoters (29, 49, 50). To investigate the extent to which the shifts of the search/recognition modes influence Egr-1’s target search kinetics in the presence of Sp1, we also conducted the kinetic assays for the four Egr-1 ZFNs using the Egr-1–Sp1 overlapping sites that Sp1 ZF-DBD initially occupied (Fig. 5A). In this assay, unless Sp1 is displaced from the overlapping sites, the Egr-1 ZFNs would not be able to carry out the site-specific DNA cleavage. Interestingly, the type 3 Egr-1 ZFN exhibited the fastest sequence-specific DNA cleavage.
in this assay as well. To gain more insight into the displacement process, we also conducted stopped-flow experiments using 5′-FAM-labeled 117-bp DNA that contains Egr-1/Sp1 overlapping target sites to which Sp1 ZF-DBD was initially bound. The presence of Sp1 ZF-DBD at the overlapping target site made the target association of Egr-1 ZF-DBDs >100-fold slower. For all constructs of Egr-1 ZF-DBDs, the target association kinetics in the presence of Sp1 ZF-DBD showed hyperbolic (rather than linear) dependence on the concentration of Egr-1 ZF-DBD (Fig. 5B). This suggests that the rate-limiting step at high concentrations is a first-order process, which is likely to be the displacement of Sp1 by Egr-1 upon formation of the encounter complex. In this case, the asymptote of the hyperbolic dependence corresponds to the first-order rate constant $k_{\text{disp}}$ for displacement of Sp1 ZF-DBD by Egr-1 ZF-DBD nonspecifically bound to DNA. Interestingly, the $k_{\text{disp}}$ constant for each construct of Egr-1 ZF-DBD was significantly larger than the rate constant for spontaneous dissociation of Sp1 from its target in the absence of Egr-1 (Fig. 5B; also see Fig. S6 and Table S2). These results suggest that in the displacement process, Egr-1 enhances Sp1’s dissociation from the overlapping target site, rather than passively awaiting its spontaneous dissociation. The first-order rate constants $k_{\text{disp}}$ were comparable for the four constructs of Egr-1 ZF-DBDs. Nonetheless, especially at relatively low concentrations, the overall displacement process for type 3 Egr-1 ZF-DBD was significantly faster than those for the other three constructs, largely because this construct can reach the overlapping target sites more rapidly than the other constructs do.

### Discussion

Since Riggs et al. discovered amazingly rapid target location by the E. coli lac repressor in 1970 (51), the mechanisms that allow DNA-binding proteins to efficiently locate their target DNA sites have been studied both experimentally and theoretically (15–22). Despite its long history over four decades, the field has remained descriptive, lacking the maturity required to enable engineering to improve the kinetic properties of natural or artificial proteins. Our current work breaks through this situation, demonstrating that improving the kinetic properties of ZF proteins is possible via engineering based on structural dynamic knowledge of the DNA-scanning process at molecular and atomic levels. Our key approach is modulation of the dynamic conformational ensemble in molecular recognition process, which has been proposed as a potentially effective strategy for protein engineering in general (52). As demonstrated above, this strategy is effective for engineering of ZF proteins to improve their kinetic properties. Although our current work is limited to Egr-1 ZF-DBD and its ZFN derivatives, we expect that our strategy is directly applicable to many artificial ZF proteins because Egr-1 (Zif268) has been used as a major scaffold for ZF technology (1–3). It is also noteworthy that interdomain dynamics are known for other natural C2H2-type ZF proteins (53–55). Although conventional engineering of artificial proteins as ZFN and TALEN (4, 5, 7, 8) focuses on affinity and specificity, a simplistic pursuit of these properties can lead to kinetically deficient proteins that are easily trapped at nonspecific sites on DNA, as seen for our type 1 constructs. Understanding DNA-scanning mechanisms at molecular and atomic levels enables creation of proteins that locate their target sites more rapidly. In fact, we found that the sequence-specific DNA cleavage by the type 3 Egr-1 ZFN was 14-fold faster than that by the type 1 construct, which exhibits the highest affinity for the target. Balanced design in terms of target search efficiency and binding affinity may greatly improve genome editing and gene control by artificial proteins and boost the applications of such proteins to basic science, therapeutics, and biotechnologies.

### Materials and Methods

#### Preparation of ZF Proteins.

The human Egr-1 ZF-DBD comprising three ZF domains (Egr-1 residues 335–423) was prepared as described previously (26–28, 56). Numbering of the DBD residues used in this paper is according to...
Pavletich and Pabo (37). The synthetic gene of the Egr-1 ZFN comprising Egr-1 ZF-DDB and FokI ND was expressed as a GST-fusion protein in E. coli. The GST-fusion protein was purified by a glutathione Sepharose FF column (GE Healthcare). After cleavage of the fusion protein by HRV-3C protease (Genway), Egr-1 ZFN was further purified via a HiTrap heparin HP column (GE Healthcare). The plasmids for the mutant Egr-1 ZF-1 proteins were prepared using the QuikChange Lightning mutagenesis kit (Stratagene). The mutant Egr-1 proteins were expressed in E. coli and purified using the same methods as those for the wild-type proteins. Complete amino acid sequences of the Egr-1 ZF-DDB and ZFN constructs are given in Fig. S7. The human Sp1 ZF-DDB (Sp1 residue 616-709) comprising three ZFs was expressed in E. coli strain BL21(DE3) via a pET-49b-derivative plasmid and chromatographically purified using cation-exchange and size-exclusion chromatography as described (57). For Egr-1 ZF-DDBs, protein concentrations were measured with Pierce BCA Protein Assay kit (ThermoFisher) (Pierce Biotechnology) by using bovine serum albumin as the standard. The protein concentrations were measured with UV absorbance at 280 nm together with the extinction coefficients of 28,880 cm⁻¹ M⁻¹ for Egr-1 ZFNs and 14,000 cm⁻¹ M⁻¹ for Sp1 ZF-DDBs.

NMR Spectroscopy for Nonspecific and Specific DNA Complexes of Egr-1 ZF-DDBs. NMR experiments were performed using Bruker Avance III spectrometers equipped with a cryogenic probe operated at the ¹H frequency of 800 MHz. NMR samples of the specific and nonspecific complexes of [H]-labeled Egr-1 ZFN and Sp1 ZF-DDB proteins together with the DNA were prepared and analyzed as previously described (28). For each sample of the complexes, the molar ratio of protein to DNA was 1:2 to ensure the DNA-bound states of the isotope-labeled proteins. For the nonspecific and specific DNA complexes of the wild-type Egr-1 ZF-DDB, resonances were assigned using ¹H/¹C/¹5N triple resonance spectra (28). Based on the assignment for the wild-type complexes, backbone ¹H and ¹N resonances of the mutant complexes were assigned using 3D ¹H/¹5N-edited NOEY and 3D ¹F/¹H/¹5N double ¹H-edited NOEY spectra. Backbone amide ¹5N relaxation parameters were measured for the complexes as described (58). Information about NMR experiments to analyze the intermolecular ion pairs of lysine side-chain NH₅ and DNA phosphate groups is given in the legend of Fig. S1.

Kinetic Studies of Association, Dissociation, Sliding, and Intergsegment Transfer of Egr-1 ZF-DDBs. The target sequence kinetics of Egr-1 ZF-DDBs were measured at 20 °C using an ISS PCR fluorescence photophysics Rx.2000 stopped-flow device as described (26, 27). In this assay, the time courses of the FAM fluorescence were monitored under mixing two solutions: (i) the Egr-1 ZF protein (20–250 nM); and (ii) FAM-labeled probe DNA (2.5 nM) and competitor DNA (28–448 M base pairs) in large excess. As the competitor DNA, sonicated calf thymus DNA (56 M base pairs) was used as described (27). For each sample of the complexes, the molar ratio of protein to DNA was 1:2 to ensure the DNA-bound states of the isotope-labeled proteins. Kinetic Assays of Sp1 Displacement by Egr-1 ZF-DBD. For the kinetic analysis, the off-target DNA cleavage should produce 3-kbp and 1-kbp fragments from the 4-kbp substrate, which was initiated by adding 5 nM Egr-1 ZFN. A 100-M base pairs DNA for the reaction mixture was sampled at distinct time points (typically, 0, 1, 2, 4, 6, 10, 20, 30, and 60 min), and the reaction of each aliquot was quenched by adding 500 M of Buffer PB (Qiagen), which denatures proteins. DNA was isolated from the quenched reaction mixtures using Qiagen spin columns. The DNA samples mixed with EZ-Vision (Amresco) were subjected to 0.9% agarose gel electrophoresis with TBE buffer (Invitrogen). DNA in the gel was visualized with fluorescence from the EZ-Vision dye using a uniform UV transilluminator, and band intensities of the 4-kbp substrate and the 3-kbp product were quantified using image software. The band of the 1-kbp product was not analyzed because of overlap with smeared bands of sonicated calf thymus DNA. The time-course data of the relative population of the 3-kbp product were analyzed via nonlinear least-squares fitting with a biexponential function. The initial apparent pseudo-first-order rate constant (i.e., the first derivative of the fitting function at time zero) and the second order rate constants were determined and used to model the constructs of Egr-1 ZFNs. MATLAB software (MathWorks) was used for the kinetic analysis. Off-target DNA cleavage (Fig. S5) was analyzed using the same experimental conditions except that 50 nM Egr-1 ZFN (instead of 5 nM) and longer reaction times were used.

Kinetic Analysis of DNA Cleavage by Egr-1 ZFNs. The target sequence of Egr1-ZFN (CCGCCCAGCTTAAAGGCTGGCGCA; Egr-1 recognition site underlined) was inserted into the pcRI-TOPO-2.1 vector using the TOPO TA Cloning kit (Invitrogen). The resultant plasmid was amplified from the culture of the E. coli strain Top10 and purified with the Qiagen Plasmid Midi kit. The circular plasmid was linearized by the restriction enzyme BglII (New England Biolab) and purified via Qiagen spin columns. Each assay of site-specific DNA cleavage by Egr-1 ZFNs was carried out at 22 °C for a 900-M solution of this linear 4-kbp substrate DNA (1 M) in a buffer containing 20 mM Tris-HCl (pH 7.5), 150 mM KCl, 0.1 mM ZnCl₂, 1 mM MgCl₂, 2 mM β-mercaptoethanol, 5% (vol/vol) glycerol, and 3 μg/ml competitor DNA (corresponding to 56 μM of base pairs). Sonicated calf thymus DNA (average length ~0.5 kbp) was used as the competitor for Fig. 4 A and 4 B and Fig. S4, whereas 28-bp nonspecific DNA was used for Fig. S5. The sequence-specific DNA-cleavage reaction, which should produce 3-kbp and 1-kbp fragments from the 4-kbp substrate, was initiated by adding 5 nM Egr-1 ZFN. A 100-M base pairs DNA for the reaction mixture was sampled at distinct time points (typically, 0, 1, 2, 4, 6, 10, 20, 30, and 60 min), and the reaction of each aliquot was quenched by adding 500 M of Buffer PB (Qiagen), which denatures proteins. DNA was isolated from the quenched reaction mixtures using Qiagen spin columns. The DNA samples mixed with EZ-Vision dye using a uniform UV transilluminator, and band intensities of the 4-kbp substrate and the 3-kbp product were quantified using image software. The band of the 1-kbp product was not analyzed because of overlap with smeared bands of sonicated calf thymus DNA. The time-course data of the relative population of the 3-kbp product were analyzed via nonlinear least-squares fitting with a biexponential function. The initial apparent pseudo-first-order rate constant (i.e., the first derivative of the fitting function at time zero) and the second order rate constants were determined and used to model the constructs of Egr-1 ZFNs. MATLAB software (MathWorks) was used for the kinetic analysis. Off-target DNA cleavage (Fig. S5) was analyzed using the same experimental conditions except that 50 nM Egr-1 ZFN (instead of 5 nM) and longer reaction times were used.

Coarse-Grained Molecular Dynamics Simulations of Intergsegment Transfer. The coarse-grained molecular dynamics simulations of the intersegment transfer between two DNA duplexes were performed for type-0 Egr-1 ZF-DDBs using a reduced model (45, 60) that allows sampling of long time-scale processes. In these simulations, two 100-bp DNA duplexes in B-form were placed in parallel with a distance of 60 Å. DNA was modeled with three beads per nucleotide, representing the phosphate, sugar, and base. Each bead was located at the geometric center of the group it represents and a negative point charge was assigned to beads representing the DNA phoshate groups. The DNA remained in-place and rigid throughout the simulations. The DNA was modeled by a single bead for the sugar-phosphate backbone and by a single bead for the base, with one negative charge at the Cα of that residue. Unlike the DNA, the protein remained flexible during the simulations but was simulated at temperature where folding and unfolding events are unlikely. Non specific protein-DNA interactions were modeled by electrostatic interactions between all charged residues of the protein and the phosphate beads of the DNA using the Debye–Hückel potential, which accounts for the ionic strength of a solute immersed in aqueous solution. The mutants were modeled by introducing or eliminating charges at the corresponding Cα beads. For example, mutant type 1 was studied by adding point charges of +1 and −1 to the Cα beads of sites 23 and 32, respectively. The type 1 mutant was also studied adding a contact between the Ca beads of Q32 and R27. This contact is expected to be formed by the Q32E mutation. Similar results were observed when having a negative charge at Q32 or by adding a contact (the +1 on T23 was modeled in both cases). Similar modeling scheme was used to model mutants types 2 and 3. Further details of the simulations and the analyses can be found in previous publications of Levy and coworkers (28, 31, 45–47, 60).

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addition of Egr-1 ZF-DBD, which eventually displaced Sp1 from the overlapping sites and bound to the target. Pseudo-first-order rate constants were measured at several concentrations of Egr-1 ZF-DBD. The first-order rate constant \( k_{\text{disp}} \) for displacement of Sp1 ZF-DBD by Egr-1 ZF-DBD (i.e., transition from the encounter ternary complex to the final specific DNA complex of Egr-1) was determined from the Egr-1 concentration dependence of \( k_{\text{disp}} \) data via nonlinear least-squares fitting with \( k_{\text{disp}} = k_{\text{disp}0}/(A_{\text{tot}} + P_{\text{inh}}) \), where \( k_{\text{disp}} \) is a first-order rate constant for displacement, \( P_{\text{inh}} \) is the total concentration of Egr-1 ZF-DBD, and \( A_{\text{tot}} \) corresponds to an apparent steady-state constant for an encounter complex and is independent of \( P_{\text{inh}} \) and \( k_{\text{disp}} \). This hyperbolic expression assumes a first-order process from the encounter complex (target DNA-Egr-1) to the final complex (target DNA-Egr-1) and can be derived in a manner similar to that for the Michaelis-Menten kinetics (61, 62). The first-order rate constant \( k_{\text{disp}} \) corresponds to the asymptote of the \( P_{\text{tot}} \) dependence of \( k_{\text{disp}} \) constants. We measured the rate constants \( k_{\text{disp}} \) for all the four pairs of Egr-1 ZF-DBDs. The \( k_{\text{disp}} \) data were compared with the first-order rate constant \( k_{\text{off}+\text{Sp-1}} \) for spontaneous dissociation of Sp1 ZF-DBD from the overlapping target site in the absence of Egr-1 ZF-DBD (details are given in Fig. 56).


Morisaki T, Imanishi M, Futaki S, Sugiura Y (2008) Rapid transcriptional activity in vivo is the total concentration of Egr-1 ZF-DBD, and \( A_{\text{tot}} \) corre-


