Auto-Inhibition of Ets-1 Is Counteracted by DNA Binding Cooperativity with Core-Binding Factor α2

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Auto-inhibition is a common transcriptional control mechanism that is well characterized in the regulatory transcription factor Ets-1. Autoinhibition of Ets-1 DNA binding works through an inhibitory module that exists in two conformations. DNA binding requires a change in the inhibitory module from the packed to disrupted conformation. This structural switch provides a mechanism to tightly regulate Ets-1 DNA binding. We report that the Ets-1 partner protein core-binding factor α2 (CBFα2; also known as AML1 or PEBP2) stimulates Ets-1 DNA binding and counteracts auto-inhibition. Support for this conclusion came from three observations. First, the level of cooperative DNA binding (10-fold) was similar to the level of repression by auto-inhibition (10- to 20-fold). Next, a region necessary for cooperative DNA binding mapped to the inhibitory module. Third, an Ets-1 mutant with a constitutively disrupted inhibitory module did not bind DNA cooperatively with CBFα2.

We have developed a structural and mechanistic model of Ets-1 auto-inhibition based on biophysical and biochemical studies (21) (Fig. 1). Ets-1 binds DNA through the ETS domain, which consists of three α helices and four β strands folded into a winged helix-turn-helix motif (5, 15, 32, 41, 72). Two regions that flank the ETS domain work together to repress the DNA binding activity of Ets-1. Deletion of either region leads to derepression, resulting in DNA binding affinity that is 10- to 20-fold higher than that measured for full-length Ets-1 (25, 27, 35, 51, 71). Nuclear magnetic resonance analyses indicate that the flanking inhibitory regions consist of three α helices, HI-1 and HI-2 amino terminal and H4 carboxyl terminal to the ETS domain (15, 59). These data support a model in which the three inhibitory helices and HI of the ETS domain pack together in a four-helix bundle to form an inhibitory module. Upon DNA binding, helix HI-1 unfolds, causing a disruption of the inhibitory module (27, 51). We have proposed that this conformational change is responsible for the relatively low DNA binding affinity of the native Ets-1. In this study, we tested whether a partnership with a second DNA binding protein counteracts the auto-inhibition of Ets-1.

We have chosen the Ets-1–core-binding factor (CBF) partnership as a model system. CBF represents a small family of unique target genes. For example, Ets-1 is in the mesodermal compartment of several tissues in the mouse embryo (33) and in T and B lymphocytes, natural killer cells, endothelial cells, and the brain of the adult mouse (4, 7, 17, 18, 73). Targeted disruption of ets-1 in the mouse results in abnormal T- and B-cell function, as well as defective natural killer cell development (4, 8, 42). However, there are other ETS domain proteins found in these tissues. For example, T lymphocytes contain the ets proteins TEL, PU.1, Elf-1, Fli-1, Erg, Ets-2, and GABP, in addition to Ets-1 (1). Thus, some level of control beyond DNA binding must determine specificity. Auto-inhibition of DNA binding provides a regulatory framework with which to generate this specificity.

Auto-inhibition modulates a variety of transcription factor activities (21). In this regulatory mechanism, inhibitory sequences act in cis to repress such functions as DNA binding, transcriptional activation, nuclear localization, and ligand interaction. The picture emerging from the study of several transcription factors suggests that protein partnerships can counteract auto-inhibition. For example, serum response factor interacts auto-inhibition. The picture emerging from the study of several transcriptional activation, nuclear localization, and ligand interaction. The picture emerging from the study of several transcriptional activation, nuclear localization, and ligand interaction. The picture emerging from the study of several transcriptional activation, nuclear localization, and ligand interaction. The picture emerging from the study of several transcriptional activation, nuclear localization, and ligand interaction. The picture emerging from the study of several transcriptional activation, nuclear localization, and ligand interaction. The picture emerging from the study of several transcriptional activation, nuclear localization, and ligand interaction. The picture emerging from the study of several transcriptional activation, nuclear localization, and ligand interaction. The picture emerging from the study of several transcriptional activation, nuclear localization, and ligand interaction. The picture emerging from the study of several transcriptional activation, nuclear localization, and ligand interaction.
CBF family synergistically activate the TCRα and TCRβ enhancers, the immunoglobulin (Ig) heavy-chain enhancer, the osteopontin promoter, as well as the Mo-MLV enhancer in vivo (16, 19, 31, 39, 55, 65, 74). Cooperative DNA binding between Ets-1 and CBFα2 is detected in in vitro studies of the TCRα and TCRβ enhancers (19, 31, 39, 74). Furthermore, the accompanying report quantifies the level of cooperativity and demonstrates that enhancement of Ets-1 and CBFα2 DNA binding is reciprocal (23). These findings suggest that DNA binding cooperativity is a mechanism for transcriptional synergy.

This study links the auto-inhibition phenomenon of Ets-1 to the DNA binding cooperativity of the Ets-1 and CBFα2 partnership. The regions of Ets-1 necessary for cooperativity and auto-inhibition overlap. In addition, disruption of the inhibitory module abrogates the effect of CBFα2 on Ets-1 DNA binding. Finally, the binding sites can be topologically uncoupled and widely spaced, suggesting that through-DNA effects do not mediate cooperativity. These findings support a mechanistic model of the Ets-1 and CBF protein partnership in which protein interactions counteract the auto-inhibition mechanism.

**MATERIALS AND METHODS**

**Protein synthesis and purification.** Full-length Ets-1 was synthesized in bacteria and purified from the insoluble fraction by conventional chromatography (27). Ets-1Δ2820 (wild type and L429A mutant) and Ets-1ΔN331 were produced in bacteria and purified from the soluble fraction by conventional chromatography (51). CBFα2 with amino acids 1 to 331 of the 451-amino-acid native CBFα residues, termed CBFα2Δ351, was produced in Sf9 insect cells with a baculovirus expression system and purified by affinity chromatography with FLAG and His tag methodology (23). Purified Ets-1 and CBFα2 proteins were incubated first with 1/10 volume of fresh 0.1 M dithiothreitol at 4°C for 30 min before each use to reverse any artificial oxidation. Activity of protein preparations was determined by DNA titration experiments (23, 27).

**Synthetic oligonucleotides.** Oligonucleotides were synthesized (Applied Biosystems model 394 or 3948 apparatus) and purified either by gel purification as described previously (27) or by reverse-phase chromatography on an automated DNA synthesizer (Applied Biosystems model 3948) followed by gel filtration (BioSpin-6; Bio-Rad). Radiolabeling of 5’ termini was performed with T4 polynucleotide kinase and [γ-32P]ATP (7,000 Ci/mol) prior to annealing of complementary oligonucleotides, as described previously (27). Synthetic complementary oligonucleotides containing the composite Ets-1 and CBF binding sites were used for the DNA binding assays. The sequences of the Mo-MLV enhancer (ets/cbf) oligonucleotides were 5’-GATCCAAAAAGGAT ATCTGGTTAAGCA-3’ (top strand) and 5’-GATCTGTTAACACAGATA TCCTGTGTTG-3’ (bottom strand). Sequences of the SCI/core oligonucleotides were 5’-GGCCAAAGCCGAAATGTGTTAATACACATTT-3’ (top strand) and 5’-AAAGTTTACACACACACGCTCGCAG-3’ (bottom strand). Mutations of these duplexes were listed in Table 1 and 2 with only the top strand presented. In the case of the oligonucleotides in which the binding site orientation was reversed, 9 and 14 bp for the Ets-1 and CBF binding sites, respectively, were reversed to accommodate both requisite core sequences (underscored in Table 2) and the preferred flanking sequences. This engineering altered the native spacing. Nicked duplexes were generated by annealing two separate oligonucleotides that represented the top strand to an intact complementary bottom strand. This DNA duplex has a nick at the junction of the two upper-strand oligonucleotides. Furthermore, the 5’ phosphate in the nick (junction) of the two upper strands is missing.

**EMSA.** Equilibrium dissociation constants (Kd) of CBFα2Δ351 and Ets-1 were determined by electrophoretic mobility shift assays (EMSA) using conditions described previously (47). In brief, the reaction mixtures were in TGEK buffer, which consisted of 25 mM Tris-Cl (pH 7.9), 10% glycerol, 6 mM MgCl2, 0.5 mM EDTA, and 60 mM KCl with 0.5 mM dithiothreitol and 200 μg of bovine serum albumin per ml. Following a 20-min incubation on ice (40 min for Ets-1Δ2820), reaction products were loaded onto a native polyacrylamide gel (acylamide:bisacylamide, 30:0.8, 45 mM Tris-borate [pH 8.3], 1 mM EDTA). The full-length Ets-1 protein was resolved on 6% polyacrylamide gels (16 cm), while the truncated Ets-1 proteins were resolved on 8% gels (20 cm). The gels were dried and exposed to a PhosphorImager screen (Molecular Dynamics), and relative radioactivity was quantified as the volume integration of individual bands. When protein titrations were used, the concentrations were in a range that resulted in approximately 0 to 100% binding. For experiments in which CBFα2Δ351 was added in saturating amounts, the concentration was at least 10-fold higher than the Kd of CBFα2Δ351 for its specific site (c~ 2 x 10^-8 M). This ensured >90% DNA occupancy. In all assays, the DNA concentrations were at least 10-fold below the

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**FIG. 1.** Ets-1 domains and conformational change. (A) Schematic of the structural and functional domains of full-length Ets-1, Ets-1Δ2820, and Ets-1ΔN331. The asterisk indicates the mutation in helix H4 of Ets-1ΔN331 as determined by nuclear magnetic resonance spectroscopy (14, 59). Shown are the tertiary structural elements of full-length Ets-1, Ets-1Δ2820, and Ets-1ΔN331 as determined by nuclear magnetic resonance spectroscopy (14, 59). Shown are the tertiary structural domains of full-length Ets-1, Ets-1Δ2820, and Ets-1ΔN331. The asterisk indicates the mutation in helix H4 of Ets-1ΔN331 as determined by nuclear magnetic resonance spectroscopy (14, 59). Shown are the tertiary structural domains of full-length Ets-1, Ets-1Δ2820, and Ets-1ΔN331. The asterisk indicates the mutation in helix H4 of Ets-1ΔN331 as determined by nuclear magnetic resonance spectroscopy (14, 59).

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**TABLE 1.** Oligonucleotides and nucleotide sequences used in EMSA. The sequences of the SCI/core oligonucleotides were 5’-GGCCAAAGCCGAAATGTGTTAATACACATTT-3’ (top strand) and 5’-AAAGTTTACACACACACGCTCGCAG-3’ (bottom strand). Mutations of these duplexes were listed in Table 1 and 2 with only the top strand presented.
estimated $K_D$ of the Ets-1 species ($\approx 10^{-11}$ M), ensuring that the total Ets-1 species concentration [$P$] was an accurate estimate of free Ets-1 species concentration [$P$].

Quantitative analysis of DNA binding. For assays containing only a single binding species, Ets-1, the $K_D$s were measured as described previously (27). Specifically, the fraction of free DNA, $[D]/[D+D_C]$, was determined by measuring the ratio of the free DNA signal analyzed at each protein concentration to the DNA signal in a control lane containing no protein. The fraction of DNA in complex with protein, $[PD]/[D+D_C]$, was derived from the relationship $[PD]/[D_C] = 1 - [D]/[D_C]$. To derive the $K_D$ with standard error (SE), the data were fit to the rearranged mass action equation, $[PD]/[D] = 1/(1 + K_D[P])$, using nonlinear least squares analyses (Kaleidagraph; Synergy Software). Multiple experiments were performed with the same range of protein concentrations to provide mean and SE values for each data point. Mean values were used for curve fitting. SEs of means are displayed as error bars. SEs of the $K_D$ values, which were provided by the curve fitting, are presented in tabular form.

To measure cooperative DNA binding, the apparent DNA binding affinities of Ets-1 species were determined in the presence of a large molar excess of the CBF$\alpha$ fragment. To perform curve fitting with the equation $[PD]/[D] = 1/(1 + K_D[P])$, several assumptions were made. (i) The disappearance of the binary complex (DNA + CBF$\alpha_2$) was the key parameter to be measured; therefore, $[D]$ was defined as the binary complex signal in a control lane that contained DNA and only CBF$\alpha_2$. (ii) The binary complex signal (DNA + CBF$\alpha_2$) was used as $[D]$ for reaction mixtures with DNA + CBF$\alpha_2$ + Ets-1 species. (iii) The fraction of DNA bound in the ternary complex was defined as $[PD]/[D]$, which was derived from $1 - [D]/[D_C]$.

RESULTS

Ets-1 binding to the Mo-MLV enhancer is repressed by auto-inhibition and enhanced by CBF$\alpha_2$. The initial goal was to provide a biological context for the hypothesis that CBF counteracts Ets-1 autoinhibition. Our approach was to document the phenomena of auto-inhibition and cooperativity on a biologically relevant site that displays synergistic transcriptional activation by Ets-1 and CBF. Our previous studies of Ets-1 DNA binding often used an artificial, high-affinity binding site, termed SC1 (47). For our initial experiments, we switched to the composite element in the Mo-MLV enhancer that binds both Ets-1 and CBF$\alpha_2$ (24, 65). To investigate the autoinhibition of Ets-1 on the Mo-MLV enhancer site, we measured the DNA binding affinity of full-length Ets-1 and the amino-terminal deletion mutant, Ets-1$^{\Delta N331}$, which lacks the amino-terminal inhibitory helices (Fig. 1). DNA binding by purified proteins was detected by EMSA. $K_D$s were determined from full binding curves. The affinity of Ets-1 was 27-fold lower than that of Ets-1$^{\Delta N331}$ (Fig. 2A; Table 1). This result is similar to the 23-fold auto-inhibition observed previously on the SC1 site (Table 1) (27). Also, as expected from previous studies, Ets-1 affinity for the Mo-MLV site was 10-fold lower than that for the high-affinity SC1 site (24, 43, 47). The detection of autoinhibition on the Mo-MLV enhancer establishes that this phenomenon is not dependent on the sequence or affinity of the binding site.

To characterize cooperative DNA binding of Ets-1 and CBF on the Mo-MLV enhancer site, Ets-1 DNA binding was measured in the presence and absence of saturating levels of CBF$\alpha_2$. We used a fragment of CBF$\alpha_2$ (spanning amino acids 1 to 331) designated CBF$\alpha_2^{\Delta C}$, which is the largest version of CBF$\alpha_2$ obtainable in a pure state from a baculovirus expression system (23). The presence of CBF$\alpha_2^{\Delta C}$ enhanced Ets-1 DNA binding affinity 20-fold (Fig. 2B and C; Table 1). This result is similar to the 10-fold enhancement of Ets-1 affinity for the artificial composite site, SC1/core, which contains the high-

### TABLE 1. Autoinhibition on the Mo-MLV and the SC1 Ets1 binding sites

<table>
<thead>
<tr>
<th>Site</th>
<th>Composite site sequence</th>
<th>CBF</th>
<th>Mean $K_D$ (nM) ± SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>MLV et/ebf</td>
<td>5'-GATCCCAAAACAGGATATCTTGTGTAAAGCA-3'</td>
<td>1</td>
<td>7.10 ± 0.58</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2</td>
<td>0.34 ± 0.028</td>
</tr>
<tr>
<td>SC1/core</td>
<td>5'-GGGCAAGCCGATATCTTGTGTAAAGCACTTTT-3'</td>
<td>3</td>
<td>8.58 ± 0.13</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4</td>
<td>430 ± 44</td>
</tr>
<tr>
<td></td>
<td></td>
<td>5</td>
<td>380 ± 55</td>
</tr>
</tbody>
</table>

### TABLE 2. Effects of altered spacing and orientation of Ets1 and CBF$\alpha_2$ binding sites

<table>
<thead>
<tr>
<th>Site</th>
<th>Composite site sequence</th>
<th>Mean $K_D$ (nM) ± SE</th>
</tr>
</thead>
<tbody>
<tr>
<td>SC1/core</td>
<td>5'-GGGCAAGCCGATATCTTGTGTAAAGCACTTTT-3'</td>
<td>0.85 ± 0.13</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.86 ± 0.0060</td>
</tr>
<tr>
<td>Nicked SC1/core</td>
<td>5'-GGGCAAGCCGATATCTTGTGTAAAGCACTTTT-3'</td>
<td>3.00 ± 0.61</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.21 ± 0.0110</td>
</tr>
<tr>
<td>invCBF</td>
<td>5'-GGGCAAGCCGATATCTTGTGTAAAGCACTTTT-3'</td>
<td>1.90 ± 0.29</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.14 ± 0.0100</td>
</tr>
<tr>
<td>invEts</td>
<td>5'-GGGCAAGCCGATATCTTGTGTAAAGCACTTTT-3'</td>
<td>0.67 ± 0.0500</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.19 ± 0.0080</td>
</tr>
<tr>
<td>$\Delta + 5$</td>
<td>5'-GGGCAAGCCGATATCTTGTGTAAAGCACTTTT-3'</td>
<td>1.90 ± 0.26</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.24 ± 0.0020</td>
</tr>
<tr>
<td>$\Delta + 10$</td>
<td>5'-GGGCAAGCCGATATCTTGTGTAAAGCACTTTT-3'</td>
<td>2.20 ± 0.23</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.25 ± 0.0030</td>
</tr>
<tr>
<td>$\Delta - 4$</td>
<td>5'-GGGCAAGCCGATATCTTGTGTAAAGCACTTTT-3'</td>
<td>2.30 ± 0.28</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.069 ± 0.0050</td>
</tr>
<tr>
<td>Nicked $\Delta - 4$</td>
<td>5'-GGGCAAGCCGATATCTTGTGTAAAGCACTTTT-3'</td>
<td>6.80 ± 0.41</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.024 ± 0.0050</td>
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</table>
affinity Ets-1 binding site SC1 juxtaposed with native spacing to the CBF site found in the Mo-MLV enhancer (Table 1) (23). DNA binding of Ets-1 and CBFW2AC also was measured on a Mo-MLV enhancer site, ets−/cbf, in which the Ets-1 recognition sequence was mutated (Fig. 2C; Table 1). Nonspecific Ets-1 binding was detected, but only at the highest concentrations of Ets-1. Dual occupancy, but no cooperative DNA binding, was observed with CBFW2AC. In conclusion, cooperative binding of Ets-1 and CBFW2AC is independent of the sequence and affinity of the binding site; however, sequence-specific DNA binding by Ets-1 is required. In summary, the Mo-MLV composite site displays both autoinhibition and DNA binding cooperativity. Thus, these initial findings link the auto-inhibition and DNA binding cooperativity phenomena to a biologically relevant enhancer element.

Regions required for cooperativity and auto-inhibition overlap. In studies of the Mo-MLV enhancer site, the level of auto-inhibition (10- to 20-fold) was in the same range as the level of cooperative binding. This was our first clue that CBFW2 might enhance Ets-1 DNA binding by affecting auto-inhibition. These results also suggested that both phenomena might involve the same region of Ets-1. To map the regions of Ets-1 that are necessary for cooperativity with CBFW2, the DNA binding affinities of amino-terminal deletion mutants Ets-1DNa and Ets-1ANaDN were measured with and without CBFW2AC (Fig. 1). The composite site, SC1/core, composed of a high-affinity Ets-1 site juxtaposed to the CBF site of the Mo-MLV enhancer, was used because the higher affinity leads to sharper bands on the EMSA gel and thus an increase in the precision of the assay. Ets-1DNa displayed a 10-fold enhancement of DNA binding in the presence of CBFW2AC, demonstrating cooperativity similar to that of full-length Ets-1 (Fig. 3A; Table 1) (23). In contrast, Ets-1ANaDN and CBFW2AC did not bind DNA cooperatively (Fig. 3B; Table 1). Our results indicate that sequences necessary for cooperativity lie between positions 280 and 331. Two amino-terminal inhibitory helices, HI-1 and HI-2, that play a key role in auto-inhibition are found in this region (Fig. 1B) (59). Because this region is part of a structural domain that includes these two helices as well as the carboxyl-terminal inhibitory helix, H4, and the ETS domain (Fig. 1B), we did not attempt to further delineate functional sequences. In conclusion, the structural domain required for auto-inhibition is also required for DNA binding cooperativity.

Analyses of DNA determinants for cooperativity implicate flexible protein-protein interactions. Two potential mechanisms could mediate cooperativity. CBFW2 could enhance Ets-1 DNA binding indirectly through the topological connection between the two binding sites. This would represent a through-DNA effect, with CBFW2 causing a change in the DNA conformation that affects the binding of Ets-1. Alternatively, CBFW2 could directly contact Ets-1 and mediate its effect through protein interactions. In this case, DNA could play a relatively passive role by simply tethering the two proteins for optimal interaction.

To distinguish between protein-protein interactions and through-DNA effects, we investigated the DNA determinants for cooperativity. To test the proposed through-DNA mechanism, the two sites were topologically uncoupled by nicking (KDr [P]), using nonlinear least squares analysis. (B) EMSA of equilibrium DNA binding studies of Ets-1 titrated onto DNA alone (top) or in the presence of a constant (~2 × 10−8 M) amount of CBFW2AC (bottom). The wedge indicates increasing amounts of Ets-1 in each binding assay as indicated in panel C. (C) Equilibrium DNA binding curves for Ets-1 (data from panel B), as well as from a repeat of these experiments performed on a mutant Mo-MLV site, ets−/cbf.
one strand of the composite binding site, SC1/core. This break in the phosphodiester backbone did not lower the level of cooperativity (Table 2). This result suggests that DNA binding by one protein does not indirectly influence the binding of the second protein. It is worth noting that this experiment does not address the long-distance effects of DNA conformation that have been detected in ternary complexes formed on more widely spaced binding sites (66).

The DNA determinants for cooperative binding also were explored by manipulating the orientation of the binding sites (Table 2). Both the Ets-1–DNA and CBFα2-DNA binary complexes are expected to show polarity because the Runt domain and the ETS domain each bind DNA as a monomer with no apparent pseudosymmetry. In the case of Ets-1, structural data demonstrate the asymmetry of the ETS domain-DNA interaction (5, 32, 41, 72). Structural information for CBFα2 also predicts an asymmetric complex (6, 44). Thus, a change in the orientation of either binding site could alter the putative protein-protein interface. Surprisingly, changing the orientation of the CBF site (invCBF) did not change cooperative DNA binding (Table 2). These results indicate that considerable flexibility exists in the CBF elements that participate in the cooperativity and implicate sequences outside the highly structured Runt domain as being important for cooperativity. This proposal is consistent with mapping data in the accompanying report (23). We propose that these flanking elements display sufficient flexibility or perhaps are located centrally within the structure, such that reversal of site orientation does not affect the CBFα2 and Ets-1 interactions. In contrast, reversal of the Ets-1 binding site orientation (invEts) reduced cooperativity from 10- to 3.5-fold (Table 2). We propose that the asymmetric location of the inhibitory module within Ets-1 expN280 causes this reduction (Fig. 1). Because sequences within this module are necessary for cooperativity, the optimal interface between Ets-1 and CBF may be compromised by the change in binding site orientation.

DNA requirements were investigated further by varying the spacing between the recognition sequences for Ets-1 and CBF. To disrupt or retain helical phasing, the spacing of 4 bp between the SC1 and core sites was expanded by 5 (Δ + 5) or 10 (Δ + 10) bp (Table 2). In addition, we tested a duplex in which the spacing was eliminated (Δ = 4). The changes in spacing reflect the variability of spacing in native sites. For example, the polyomavirus enhancer and osteopontin promoters have the added 9 or 10 bp, whereas the TCRβ enhancer is identical to Δ = 4 (19, 55, 65, 74). Both the Δ + 5 and Δ + 10 duplexes displayed cooperativity between Ets-1 and CBFα2 (Table 2). (These results provide an important control for the site orientation experiments above, in which spacing was altered.) Because locally altered DNA conformation does not easily lead to long-distance effects (75), this retention of cooperativity supports the conclusion that through-DNA effects are not involved in the 10-fold enhancement of Ets-1 binding. Again, these results imply that protein elements that mediate cooperativity must accommodate considerable structural flexibility.

In contrast to results from widely spaced binding sites, the Δ = 4 duplex displayed a higher level of cooperative DNA binding than the SC1/core composite site (Fig. 1, Table 2). The 33-fold cooperativity could be mediated by additional direct contacts that would be possible due to the closer proximity of the proteins. Alternatively, the proximity of the recognition sequences could facilitate through-DNA effects. To test this latter alternative, a nick was introduced into one strand of the Δ = 4 duplex (Table 2). Nicking did not lower the level of cooperativity. The retention of cooperativity eliminates an indirect mechanism that uses the topological coupling of the binding sites. Instead, the nicked binding site displayed an increase in cooperative DNA binding, from the 33-fold effect observed with Δ = 4 to a 250-fold effect (Fig. 4; Table 2). There are several possible reasons for this increase. A torsional strain in the DNA backbone could develop due to the proximity of the sites, and the nick relieves the strain by uncoupling the sites. Alternatively, the loss of phosphate at the site of the nick could affect the energetics of binding. These findings suggest that an additional mechanism contributes to cooperativity of closely apposed sites. Future investigations will explore this possibility.

**Derepressed Ets-1 mutant is resistant to CBF cooperative effects.** The results of quantitative binding assays and mapping experiments suggested that cooperativity could function by counteracting auto-inhibition of Ets-1. Furthermore, the analysis of DNA determinants suggested that the mechanism involves protein interactions rather than through-DNA effects. Finally, considerable flexibility is expected in these protein interactions. There are two possible models to explain how the Ets-1-CBFα2 partnership counteracts auto-inhibition through such flexible protein interactions. In one scenario, CBFα2 could cause a conformational change in the folded inhibitory module. We envision this as a repositioning of structural elements such that unfolding of helix HI-1 is no longer necessary. Alternatively, CBFα2 could favor a shift in the equilibrium...
toward the disrupted inhibitory module. Unfolding and/or refolding of helix HI-1 could be affected.

To gain additional insight into the mechanism of cooperativity, we tested a version of Ets-1 in which the inhibitory module is constitutively disrupted and that displays derepressed DNA binding. Ets-1ΔN280,L429A bears an amino acid substitution in the inhibitory helix H4 (Fig. 5A). Proteolysis experiments indicate that helix HI-1 is constitutively unfolded in this mutant version of Ets-1 (data not shown). In quantitative binding assays, Ets-1ΔN280,L429A bound DNA with a 15-fold-higher affinity than Ets-1ΔN280 (Fig. 5B; Table 1). More importantly, CBFα2AC did not enhance the affinity of this mutant version of Ets-1 (Fig. 5B; Table 1). The presence of a constitutively unfolded inhibitory module resulted in insensitivity to CBFα2 effects. This finding strongly suggests that the DNA binding cooperativity mechanism involves regulating the conformation of the inhibitory module. As detailed in Discussion, these results are consistent with either mechanistic model. In conclusion, the behavior of this mutant demonstrates that the inhibitory module is required for DNA binding cooperativity, thus supporting our proposal that DNA binding cooperativity counteracts auto-inhibition.

FIG. 4. Effects of topological uncoupling of DNA binding sites for Ets-1 and CBFα2AC. (A) EMSA of equilibrium DNA binding studies of Ets-1 titrated onto nicked Δ–4 DNA alone (top) or in the presence of a constant (2 × 10^{-8} M) amount of CBFα2AC (bottom). (B) Equilibrium DNA binding curves for Ets-1 obtained from EMSA in the absence (open symbols) and presence (filled symbols) of CBFα2AC. Cooperative DNA binding was measured on either the Δ–4 (circles) or nicked Δ–4 (triangles) DNA duplex (Table 2).

FIG. 5. Derepressed Ets-1 variant displays no DNA binding cooperativity with CBFα2. (A) Schematic representation of the constitutive unfolding of Ets-1ΔN280,L429A due to a leucine-to-alanine substitution in helix H4, as shown (×). The K_D value is for the SC1/core. (B) Equilibrium DNA binding curves for Ets-1ΔN280,L429A obtained from EMSA in the absence (open circles) and presence (filled circles) of a constant (2 × 10^{-8} M) amount of CBFα2AC.

DISCUSSION

Auto-inhibition provides a mechanism by which protein activity can be regulated tightly. In the case of Ets-1, this auto-regulation decreases the DNA binding affinity 10- to 20-fold. We predicted that a regulatory mechanism would counteract this auto-inhibition and enable the full DNA binding potential of the Ets-1 ETS domain to be used within a biological context. Ets-1 functions in association with other transcription factors, implicating cooperative DNA binding with other proteins as a mechanism for rescinding auto-inhibition. Both this study and the accompanying report (23) present quantitative analyses of the DNA binding cooperativity between Ets-1 and CBFα2. The quantitative approach of these experiments is distinctive, representing one of only a few such comprehensive studies of a DNA binding partnership in a eukaryotic system. These
experiments also laid the groundwork for our mechanistic model of CBF rescinding Ets-1 auto-inhibition.

**Mechanistic model of CBFα2 and Ets-1 cooperative DNA binding.** There are two general mechanisms by which CBFα2 could affect Ets-1 DNA binding. Due to the close apposition of the Ets-1 and CBF binding sites, we considered a through-DNA mechanism that would require no direct protein interactions. This model was discounted by the observations that topological uncoupling of the two sites did not disrupt cooperativity. In addition, there was considerable flexibility in the spacing and orientation of Ets-1 and CBFα2 binding sites. Consistent with these observations, selection of consensus binding sites for Ets-1 and CBFα2 also detected flexibility in site configuration. Our study provides a rigorous quantitative analysis of this type of flexibility.

Our proposed mechanism for DNA binding cooperativity is based on the model of Ets-1 auto-inhibition (Fig. 1) (21). DNA binding is accompanied by a conformational change that includes unfolding of one of the three inhibitory helices. We have proposed that this requisite conformational change reduces the DNA binding affinity of full-length Ets-1 and other repressed species such as Ets-1ΔN280. This report links autoinhibition and DNA binding cooperativity. Quantitative binding studies showed that DNA binding cooperativity requires the presence of the amino-terminal inhibitory helices. Because this region is part of the inhibitory module, we envision that CBFα2 could affect the conformational state of this structural domain. Two possible models are envisioned (Fig. 6). First, CBFα2 could alter the position of the inhibitory module such that the unfolding of HI-1 is no longer necessary for DNA binding. This repositioning model predicts a protein interface between Ets-1 and CBFα2 that would stabilize the inhibitory helices in the absence of their usual contact to the ETS domain. Such direct contacts could be made within any of the inhibitory helices. Alternatively, CBFα2 could stimulate the rate of unfolding and/or prevent refolding of helix HI-1. In contrast to the first model, this disruption model need not evoke a specific interface between the disrupted inhibitory module and CBFα2 since no structural elements are being stabilized. Either of these scenarios accommodates the flexibility of the DNA determinants for cooperativity.

The behavior of the constitutively activated mutant, the L429A derivative of Ets-1ΔN280, can be explained by both models. The inhibitory module of this mutant protein is constitutively disrupted and thus predominantly in the unfolded state. If CBFα2 alters the position of the inhibitory module by making specific contacts with structural elements, the disrupted inhibitory helices in Ets-1ΔN280,L429A would not provide the required protein interface, consistent with the repositioning model. The behavior of Ets-1ΔN280,L429A also can be explained by the model in which CBFα2 evokes a specific interface between the disrupted inhibitory module and CBF in the ternary complex. If such contacts were to exist, the L429A variant might be expected to retain the interface and, thus, cooperativity. Thus, the lack of cooperativity between CBFα2 and Ets-1ΔN280,L429A is also consistent with the disruption model. To distinguish between these alternative mechanisms, structural studies are under way to determine the conformational state of the inhibitory module within the ternary complex.

Our preliminary analyses suggest that any interaction between Ets-1 and CBFα2 at physiologically relevant concentrations occurs only on DNA. Specifically, surface plasmon resonance experiments that were performed at protein concentrations in the 10 nM range failed to detect interactions (data not shown). The interaction could have such a low affinity that it will be difficult to detect by this approach. Alternatively, DNA binding by one or the other partner may expose residues necessary for the interaction. In this regard, it is interesting that reciprocal cooperativity, which enhances CBFα2 DNA binding, was observed only under conditions in which CBFα2 was incubated with DNA before the addition of Ets-1 (23). Similarly, the enhancement of Ets-1 binding reported here and in the accompanying report (23) was detected under conditions in which CBFα2 is likely to bind before Ets-1. Although all
components were added simultaneously to the reaction mixtures, the high concentration of CBFα2 (10^{-8} M) is expected to cause a rapid association of CBFα2, facilitating preassembly of a DNA-CBFα2 binary complex. The model in Fig. 6 reflects this proposed sequence of events.

There are two reports that describe direct interactions between Ets-1 and CBFα2 in the absence of DNA. In one study, the amino-terminal sequences of Ets-1 (amino acids 123 to 240), which lie outside the regions mapped here, interact directly with CBFα2 in a glutathione S-transferase pulldown assay (19). In a second study, glutathione S-transferase pulldown assays detected CBFα2 interactions with only the carboxyl-terminal half of Ets-1 (31). The concentration and folded state of the proteins were not measured in these assays. Thus, neither the affinity nor the specificity of these interactions has been rigorously determined. High-resolution structural and genetic analyses are necessary to describe the interface of Ets-1 and CBFα2 in the ternary complex.

In the accompanying report (23), which focuses on the DNA binding properties of CBF, we report that Ets-1 enhances the DNA binding activity of CBFα2 at least sevenfold. Thus, as expected from thermodynamic principles, the DNA binding cooperativity between Ets-1 and CBFα2 is reciprocal. Several studies report that CBFα2 also displays auto-inhibition (23, 30, 31). We suggest that Ets-1 counteracts the auto-inhibition of the CBFα2 Runt domain, perhaps also altering an inhibitory conformation. Interestingly, the cooperative binding of CBFα2 with Ets-1 is an alternative to the enhancement of CBFα2 DNA binding by its heterodimeric partner CBFβ. For additional discussion of the interplay between Ets-1 and CBFβ see the accompanying report (23).

By necessity our quantitative studies were performed with a truncated version of CBFα2 that could be obtained highly purified from a baculovirus expression system. We were initially concerned that this experimental design could compromise our study. However, recently a less quantitative analysis of DNA binding cooperativity between Ets-1 and CBFα2 that utilized full length CBFα2 prepared by in vitro transcription-translation was reported (31). Importantly, the two studies concur on the regions of Ets-1 that are important for DNA binding cooperativity. These findings thus lend support to our conclusions that are based on the behavior of truncated CBFα2.

Combinatorial control of transcription. Partnerships between two transcription factors are the building blocks of combinatorial control of gene expression. These types of interactions are predicted to enhance the specificity of regulatory transcription factors. In the case of Ets-1 and CBFα2, each binds a sequence-specific region of only 9 to 10 bp as a monomer, whereas the ternary complex binds to a composite site that spans 18 to 20 bp of specific sequences. This extended region of preferred sequence provides a higher degree of sequence specificity and thus could direct Ets-1 to function preferentially on enhancer elements with CBFα2 (12). Interestingly, in the case of Ets-1, there are other putative DNA binding partners, including Jun/Fos, TFE3, USF, and NF-κB (12, 37, 58, 66). Preliminary quantitative studies indicate that NF-κB has effects on Ets-1 DNA binding comparable to those of CBFα2 (12) (T. Goetz, unpublished data). This promiscuity of Ets-1 could reflect multiple pathways to transcriptional synergy, including alternative mechanisms for DNA binding cooperativity. However, these other partnerships could work also by counteracting auto-inhibition. Indeed, a recent report implicates the basic helix-loop-helix protein TFE3 in this role on the immunoglobulin η enhancer (66). The mechanistic models developed here could accommodate Ets-1 interactions with these other partner proteins.

The maximal activity of complex enhancers requires large assemblies of DNA binding proteins. This phenomenon is well illustrated by the TCRα enhancer, in which a basic helix-loop-helix protein (either USF or TFE3) and the high-mobility-group domain protein Lef-1 function with Ets-1 and CBFα2 (19, 26, 39). Therefore, the Ets-1 and CBFα2 partnerships works in the context of a much larger multiprotein complex. Another such example is the function of Ets-1 on the human immunodeficiency virus type 1 viral enhancer in collaboration with Sp1, NF-κB, Lef-1, and USF (or TFE3) (57, 64). Such complexes provide additional opportunities to enhance the specificity and affinity of regulatory transcription factors.

An additional interesting feature of this network of interactions is the family membership of Ets-1 and CBFα2. As described earlier, there are at least 20 vertebrate ets genes and three CBFα genes. Other ets proteins also synergize with CBFα2 (e.g., Fli-1, PU.1, GABP, and MEF) (38, 52, 65). Similarly, another CBFα protein, CBFα1, is functionally linked to Ets-1 (55). Furthermore, multiple members of the ets family of transcription factors display auto-inhibition of DNA binding (22). It is possible that the mechanisms of auto-inhibition and DNA binding cooperativity described here are sufficiently conserved to provide a framework for understanding many of these combinations of transcription factors.

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REFERENCES


