Synergistic transcriptional enhancement does not depend on the number of acidic activation domains bound to the promoter

(transcriptional activation/eukaryotic promoters/cooperativity/enhancers/Fos and Jun oncoproteins)

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ABSTRACT Many eukaryotic transcriptional activator proteins contain a DNA-binding domain that interacts with specific promoter sequences and an acidic activation region that is required to stimulate transcription. Transcriptional enhancement by such activator proteins is often synergistic and promiscuous; promoters containing multiple binding sites for an individual protein or even for unrelated proteins can be 10-100 times more active than promoters with single sites. It has been suggested that such synergy reflects a nonlinear response of the basic transcription machinery to the number and/or quality of acidic activation regions. Here, we determine the transcriptional activity of Jun–Fos heterodimers containing one or two GCN4 acidic activation regions on promoters containing one or two Ap-1 target sites. Surprisingly, heterodimers with one or two acidic regions activate transcription with similar efficiency and are equally synergistic (10- to 15-fold) on promoters containing two target sites. Thus, transcriptional synergy does not depend on the number of acidic activation regions but rather on the number of proteins bound to the promoter. This suggests that synergy is mediated either by cooperative DNA binding or by alternative mechanisms in which the DNA-binding domain plays a more direct role in transcription (e.g., changes in DNA structure, nucleosome displacement, or direct interactions with the transcriptional machinery).

Eukaryotic transcription factors contain distinct DNA-binding and transcriptional activation functions that are generally located in separate regions of the protein (1–3). Transcriptional activation domains are often defined by short acidic regions that function autonomously when fused to heterologous DNA-binding domains (3–5). Although many acidic sequences can serve as transcriptional activation regions and negative charge is clearly important (3–8), the level of transcriptional stimulation is influenced by other structural features such as the length of the region and possibly the α-helical character (8, 9). It has been hypothesized that the DNA-binding domain serves merely to bring the protein to the DNA target, whereupon the acidic activation region can interact with a component(s) of the basic transcription machinery. Since acidic regions are necessary for yeast activator proteins to function in mammalian cells (10, 11) and for mammalian activator proteins to function in yeast cells (5, 12), it is likely that they contact some part of the basic transcription machinery that is conserved functionally throughout the eukaryotic kingdom.

Transcriptional enhancement by activator proteins is synergistic in that promoters containing multiple binding sites upstream of a “TATA” element are often 10–100 times more active than analogous promoters containing single binding sites (for review, see ref. 13). Moreover, transcriptional synergy is frequently observed when the multiple binding sites are recognized by distinct, and even evolutionarily distant, proteins. For example, the combination of the mammalian glucocorticoid receptor and the yeast GAL4 protein stimulates transcription much more effectively than either protein alone. Such promiscuous synergy between activator proteins is a fundamental aspect of eukaryotic transcription and constitutes an important basis for the extraordinary diverse patterns of gene expression mediated by enhancers.

Cooperative DNA binding of transcription factors to adjacent promoter sites represents a simple mechanism that is likely to account for at least some instances of transcriptional synergy. Such cooperative binding has been observed in vitro for some proteins (14–16) and generally reflects highly specific protein–protein interactions that contribute to the overall stability of the protein–DNA complexes (17, 18). However, the apparent requirement for highly specific protein–protein interactions makes it difficult to invoke cooperative DNA binding for explaining the promiscuous nature of transcriptional synergy. Moreover, synergistic activation has been observed in vitro under conditions where the binding sites for a given activator protein are fully occupied (19, 20). For all of these reasons, it seems very likely that there must be alternative mechanisms of synergy beyond cooperative DNA binding.

One such alternative mechanism is that acidic activation regions associated with DNA-binding proteins that are bound to adjacent promoter sequences interact synergistically with a common target of the basic transcription machinery (for review, see ref. 13). In such a model, the common target would respond functionally in a nonlinear fashion to the number and/or quality of acidic activation regions. Here, we test this proposal by determining whether transcriptional synergy depends on the number of acidic activation regions. To vary the number of acidic regions without changing the number of DNA-bound proteins, we compare the level of transcriptional activation mediated by DNA-binding heterodimers carrying either one or two acidic regions. In contrast to the predictions of the above model, the results indicate that synergistic enhancement does not depend on the number of acidic activation domains but rather on the number of proteins bound to the promoter.

MATERIALS AND METHODS

DNA Manipulations and Construction of Yeast Strains. DNAs encoding the various chicken Jun derivatives were derived from YCp88, a ara3 vector that utilizes the ded1 promoter for expression of the proteins (3). JunGa was constructed by inserting the XbaI–EcoRI fragment encoding the Jun DNA-binding domain in place of the yeast GCN4 DNA-binding domain of LexA-gcn4ΔA20 (8) and then replacing the Escherichia coli LexA region with a BssHII–Sal I fragment containing the ded1 promoter fused to AGC-TACGGGTACAAAAAGAAAAATGAGATTTTCTTTCGAGGG-TAAAAGCAAAAAGATT that provides the AUG initia-

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tion codon and 12 additional amino acids in-frame with the GCN4 coding sequence. JunA4 and JunA9 were obtained from the LexA derivatives (5) by substitution of the LexA region as described for JunGa. The mouse Fos molecules (cloned in a derivative of YCp88 that contains trp1 as the selectable marker) were constructed by combining the BssHII–EcoRI/filled-in fragment carrying the ded1 promoter and the above oligonucleotide sequence, the Fsp I–Xho II fragment (to which a Sal I linker was added at the Xho II site) from v-fos (21), and the Sal I–EcoRI fragments of GCN4 deletions that either do (N125) or do not (N71) contain the GCN4 activation domain (3).

To construct yeast strains KY371 and KY372, the relevant DNAs were introduced into the his3 locus of KY329 (22) by gene replacement. The plasmid carrying the two optimal GCN4 binding sites (his3-282) was obtained by cloning the oligonucleotide shown in the bottom line of Fig. 1 as an EcoRI–Sac I fragment into YIp55-Sc4099 (23), a derivative with a deletion between positions −447 and −83 of the wild-type his3 promoter. The plasmid containing one optimal GCN4 binding site (his3-281) was obtained by deletion of the above DNA between the EcoRI and Sal I sites. Plasmid DNAs containing the various Jun and Fos derivatives were introduced into KY371 and KY372 by selecting for the appropriate markers.

RNA Analysis. RNAs from KY371 and KY372 derivatives were hybridized to completion with an excess of 32P-labeled oligonucleotides for his3 and for ded1 (the internal control) and treated with S1 nuclease as described (24). His3 RNA levels, normalized to ded1 RNA levels in the same lane, were quantitated by densitometry; the level of transcription activated by GCN4 in strain KY372 was defined as 100.

DNA-Binding Experiments. The EcoRI–Sac I fragment from the his3-282 promoter, which contains two GCN4 optimal binding sites, was end-labeled and incubated with various amounts of GCN4 protein produced in E. coli cells and purified to near homogeneity as described (25). The resulting protein–DNA complexes were separated from unbound DNA by electrophoresis in a 5% native polyacrylamide gel. The percentage of input DNA found in complex I (contains one bound GCN4 dimer) or complex II (contains two bound GCN4 dimers) was quantitated by densitometry. The protein composition of the two electrophoretically distinct complexes was inferred from the observation that only complex I is formed on the analogous fragment containing one optimal binding site (data not shown). Independent binding of GCN4 to the adjacent sites can be seen most easily under conditions of moderate occupancy (50 and 100 ng) where the fraction of DNA bound in complex II equals the square of the fraction bound in complex I.

RESULTS

Experimental Design. The crucial issue to be addressed in this paper is whether synergy depends on the number of acidic activation domains or on the number of proteins bound upstream of the TATA element. Normally, increasing the number of protein binding sites upstream of the TATA element results in the concomitant increase of both the number of acidic regions and DNA-binding proteins at the promoter. To alter the number of acidic domains independently of the number of DNA-binding proteins, we devised an experimental situation in which transcription was activated by DNA-binding heterodimers. In this way, acidic activation regions could be fused to one or both partners of the heteromeric complex.

To obtain such heterodimers, we fused the GCN4 acidic activation region (3, 8) to the DNA-binding domains of the Jun and/or Fos oncoproteins (JunGa and FosGa; Fig. 1). As controls, we utilized Jun deletions that remove part (JunΔ4) or all (JunΔ9) of the Jun activation region (5) as well as a deletion of Fos (FosΔ1). The Jun and Fos DNA-binding domains peripherally form a heterodimer that binds specifically to a transcription factor AP-1 site (26–29), a dyad-symmetric sequence that is essentially identical to the GCN4 recognition site (22, 23). Fos is unable to form homodimers and can only bind DNA as a heterodimer with Jun. Jun homodimers can be formed, but in yeast cells they are very inefficient in activating transcription from promoters containing AP-1/GCN4 sites, presumably due to relatively weak dimerization and DNA-binding activity (5, 22). Thus, the transcriptional activity of Jun–Fos heterodimers containing one or two GCN4 acidic regions can be measured directly on promoters that contain a required AP-1 site(s) without significant interference from the contributions of Jun homodimers and Fos homodimers.

Transcriptional Activity of Fos–Jun Heterodimers on Promoters Containing One AP-1 Site. Plasmid DNAs capable of expressing the Jun and Fos derivatives were introduced into yeast strain KY371, which is deleted for the GCN4 gene and contains his3-281, an allele with one optimal AP-1 binding site.
upstream of the TATA element and mRNA coding region. The resulting strains were assayed for their levels of his3 RNA (Fig. 2A). In the absence of any Jun or Fos derivative, his3 transcription was extremely low and in fact the cells were unable to grow in the absence of histidine; this reflects the absence of GCN4 protein as well as the upstream pro-

![Figure 2: Analysis of his3 transcription. RNAs from KY371 derivatives, which contain a his3 promoter with one optimal binding site (A), or KY372 derivatives, which contain a his3 promoter with two optimal binding sites (B) were hybridized to 32P-labeled his3 and ded1 probes; positions corresponding to the ded1 and his3 +1 and +13 transcripts are indicated. His3 RNA levels, normalized to ded1 RNA levels in the same lane, are shown in C (100 is defined as the level achieved by GCN4 in strain KY372).](image)

moter elements necessary for constitutive his3 transcription (30, 31). When tested alone, JunGa showed some activation of his3 expression, JunΔ4 was barely effective, and JunΔ9 or either of the Fos derivatives was inactive. Thus, with the possible exception of JunGa (see below), his3 transcription cannot be significantly induced by any of the Jun or Fos homodimers. In addition, the results rule out the possibility of endogenous yeast proteins that can heterodimerize with the Jun or Fos derivatives to produce a transcriptionally active complex with the AP-1 binding site.

The fact that heterodimers containing a single GCN4 acidic region can activate his3 transcription was most clearly illustrated by the combination of FosGa and JunΔ9 (Fig. 2A; compare lanes 5 and 6 with lane 12). Neither protein can activate transcription alone because FosGa is not able to bind DNA and JunΔ9 lacks an activation region. Activation by the FosGa–JunΔ9 heterodimer depends primarily on the single GCN4 acidic region fused to FosGa because only 20–30% of much his3 transcription was observed in cells containing JunΔ9 and FosΔ1. Thus, dimerization of the GCN4 acidic region is not necessary for transcriptional activation.

Interestingly, all the Fos–Jun combinations that contain either one or two activation regions stimulated his3 transcription to comparable levels (Fig. 2A, lanes 8–12). This result also confirmed the expectation that Jun homodimers do not significantly contribute to transcription in cells containing both Jun and Fos derivatives. Specifically, similar RNA levels were obtained when either JunGa or JunΔ9 was combined with FosGa even though JunGa homodimers could contribute to his3 expression, whereas JunΔ9 homodimers could not contribute and might possibly interfere with his3 expression by binding to the AP-1 site. These observations indicate that the activation domain is a monomeric structure and that the number of acidic regions on a DNA-bound protein does not significantly affect the level of transcription.

Transcriptional Activity of Fos–Jun Heterodimers on Promoters Containing Two AP-1 Sites. When the Jun–Fos combinations were introduced into KY372, a strain containing the his3-282 promoter, which has two adjacent AP-1 sites, his3 RNA levels were approximately 10-fold higher than observed in the corresponding KY371 derivatives that contain one AP-1 site (Fig. 2B, lanes 8–12). Again, heterodimers containing one acidic region activated transcription almost as well (60–90%) as heterodimers containing two acidic regions (Fig. 2C). Thus, there was a dramatic difference in transcriptional activation when two acidic domains were located on two DNA-binding proteins as opposed to the situation when the same two acidic regions were located on a single DNA-bound molecule. This demonstrates that transcriptional synergy does not depend on the number of acidic activation regions but instead depends on the number of proteins bound to the promoter. Although the length, quality, and probably number of acidic domains clearly contribute to the level of transcriptional activation (3, 6–9), they do not appear to be responsible for the synergistic effects.

Several other observations should be noted. (i) Some synergistic activation was observed with JunGa and JunΔ4 homodimers as well as with JunΔ9–FosΔ1 heterodimers (the residual activity was probably provided by the FosΔ1 moiety), even though the absolute levels of transcription were low. (ii) Activation by Jun–Fos heterodimers from a single AP-1 site resulted in his3 transcription initiated equally from the +1 and +13 sites, a pattern typically associated with constitutive his3 transcription (32). In contrast, Jun–Fos activation through two AP-1 sites was initiated with a strong preference for the +13 site, the pattern observed during activation by GCN4 (Fig. 2A; ref. 32) or by GAL4 (33).

Synergistic Activation by GCN4. Parallel experiments carried out in strains KY371 and KY372 indicated that wild-type GCN4 protein also synergistically activated transcription;
his3 RNA levels were 15-fold higher on the promoter containing two AP-1 sites (Fig. 2). Transcription activated by GCN4 was 4–7 times more efficient than transcription activated by the Jun–Fos heterodimers. Although this could be due to differences in DNA-binding affinity, it more likely reflects the fact that only a portion of the GCN4 activation region was fused to the Jun and Fos DNA-binding domains. Given the homodimeric nature of GCN4, it is not possible to explicitly separate the contributions of the DNA-binding domain and the transcriptional activation region. However, since the number of GCN4 activation regions present on the Jun–Fos heterodimers does not account for synergistic enhancement, it seems unlikely that the number of GCN4 acidic domains would explain the synergy mediated by GCN4 itself.

GCN4 Binds Noncooperatively to Adjacent Sites. As mentioned in the introduction, cooperative DNA-binding is a simple mechanism to account for transcriptional synergy. To examine this possibility for the case of synergy by GCN4, we carried out standard DNA-binding titrations using E. coli-produced GCN4 protein and an oligonucleotide derived from the his3-281 promoter that contains two adjacent binding sites. Complexes containing one or two bound GCN4 molecules were distinguished by their electrophoretic mobility in native acrylamide gels. As shown in Fig. 3, the data are entirely in accord with noncooperative binding. Independent binding of GCN4 to the adjacent sites could be seen most easily under conditions of moderate occupancy (50 and 100 ng) where the fraction of DNA bound in complex II equaled the square of the fraction bound in complex I. Similar results using DNase I footprinting on other promoters containing multiple GCN4 binding sites have been obtained in this laboratory by C. R. Wobbe (unpublished data).

**DISCUSSION**

The GCN4 Acidic Activation Domain Is Monomeric. Detailed analyses of the GCN4 activation domain indicate that the level of transcriptional stimulation is moderately correlated with the number of acidic residues but is influenced by other structural features such as the length of the region and possibly α-helical character (3, 8). For several reasons including the fact that GCN4 binds to DNA as a dimer (34), it was suggested that activation might require dimerization of the acidic region (8). However, the basic observation that Fos-Jun heterodimers containing a single GCN4 acidic region are efficient transcription factors indicates that the GCN4 activation domain is a monomeric structure. Furthermore, the results indicate that proteins lacking an activation region can stimulate transcription if they can associate into heteromeric DNA-binding complexes with partners that contain an activation domain. Conversely, the ability of a gene product to stimulate transcription does not necessarily indicate that the protein itself contains an activation function. In fact, conventional mapping of the transcriptional activation function on such a protein would instead uncover a motif necessary for oligomerization and/or DNA binding. Most importantly, however, the monomeric nature of acidic activation domains increases the complexity of regulation that can be mediated by protein families that contain common dimerization motifs such as the leucine zipper (35) and the helix–loop–helix (36).

**Mechanism of Synergistic Transcriptional Activation.** The results here argue against the prevailing view that synergy reflects a nonlinear response of a "common target" to the number and/or quality of acidic activation regions (13). This cannot be excluded, but consider unlikely, the formal possibility that one activation region can inhibit the activity of a second activation region on the same DNA-bound molecule but not on an adjacent bound molecule. The GCN4 activation region is unstructured and physically separate from the DNA-binding domain (8), and it can function as a monomer; hence, it is likely to have considerable flexibility in finding a putative target in the transcription machinery. In this regard, the two activation regions in the JunGa–FosGa heterodimer lie in opposite configurations with respect to the DNA-binding domains. Finally, heterodimers with two GCN4 acidic regions activate somewhat better (30–60%) than heterodimers with only a single acidic region (Fig. 2B).

One explanation for synergistic activation invokes cooperative DNA binding on adjacent target sites. Although this model is probably valid for specific situations (14, 16), it does not easily account for the many examples of synergistic activation by different (and in some cases evolutionarily distant) proteins and hence is unlikely to be generally correct. Cooperative DNA-binding involves specific protein–protein interactions that are generally mediated by regions of the protein that are distinct from the DNA-binding domain (14, 17, 18). In contrast, the Fos and Jun moieties here are extensively deleted such that the potential contacts between adjacent heterodimers would have to involve the DNA-binding domains themselves or immediately adjacent regions of the protein. In this case, we suspect, but cannot demonstrate, that cooperative binding does not account for the observed synergy.

For synergistic activation by GCN4, three observations argue against a cooperative binding mechanism. (i) Such cooperativity has not been observed in DNA-binding experiments carried out in vitro using E. coli-synthesized GCN4 protein and the promoter DNAs described here (Fig. 3). (ii) Expression of the GCN4 DNA-binding domain lacking the acidic activation region results in repression of many GCN4-regulated yeast genes (3). (iii) At equivalent in vivo protein concentrations as employed here, GCN4 can repress transcription when bound at a single site located immediately downstream of the required TATA element in a gal-his3 promoter (37). These two examples of repression strongly suggest that in vivo a single binding site is frequently occupied by GCN4 and hence that cooperative binding cannot increase.

Fig. 3. Noncooperative binding of GCN4. DNA containing two GCN4 binding sites was incubated with the indicated amounts of E. coli-produced GCN4 protein, and the percentage of input DNA found in complex I (contains one bound GCN4 dimer) or complex II (contains two bound GCN4 dimers) was determined. Independent binding to the adjacent sites is best seen under conditions of moderate occupancy (50 and 100 ng) where the fraction of DNA bound in complex II equals the square of the fraction bound in complex I.
promoter occupancy by a factor of 10. Since the number of GCN4 acidic regions does not account for synergistic activation by the Fos-Jun heterodimers, it would seem unlikely to explain the synergism mediated by GCN4 itself.

In situations where transcriptional synergy is not due to cooperative DNA binding, we suggest an alternative explanation in which the DNA-binding domain plays a more direct role in transcription than simply targeting the protein to the promoter. This idea, though contrary to some suggestions (13), is supported by the existence of glucocorticoid receptor or HAPI protein derivatives that bind DNA normally but fail to activate transcription (38, 39). In addition, the GCN4 DNA-binding domain can interact selectively with RNA polymerase II in vitro (40). Given the constraints imposed by the results in this paper, several noncooperative binding models for synergistic enhancement could be imagined. The DNA-binding domain might alter DNA structure, affect nucleosome distribution on the chromatin template, or directly interact with the basic RNA polymerase II transcriptional machinery. By any of these models, the acidic activation region presumably would carry out a different function than the DNA-binding domain in the overall process of transcriptional enhancement.

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