Modularity of eukaryotic transcription activators

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The key experiment described here showed that a eukaryotic transcriptional activator is a modular protein composed of a DNA-binding domain and an "activating region" (2466). The definition of these as distinct, separable elements led to a coherent picture for how such activators work; provided a powerful tool for the analysis of other gene-activating and -repressing functions; laid the basis for the development of the two-hybrid system for detecting protein-protein interactions; and now provides a framework for understanding how specificity is achieved in many cellular information processing networks.

Background

The key experiment we shall describe was performed in yeast, a simple eukaryote. Our understanding of gene regulation in bacteria around 1985 strongly influenced the design of the experiment. By that time, we had in hand three concepts describing how a bacterial regulatory protein could recognize a specific sequence in DNA (called an operator) and activate or repress transcription of a gene. We begin here with a brief statement of these three concepts, as exemplified by the action of lambda repressor, the protein we were studying most intensively (see EXP, Isolation of repressor†):

- A regulatory protein (repressor or activator) binds a specific sequence in DNA without significantly changing the structure of either the protein or the DNA.
- A repressor turns off (represses) a gene by excluding RNA polymerase from that gene. This effect requires that an operator site overlap the binding site for RNA polymerase (the promoter).
- An activator turns on (activates) transcription of a gene that bears an operator site immediately adjacent to the promoter. Activation is effected by a protein-protein interaction between the regulator and RNA polymerase, an interaction that we now know recruits RNA polymerase to the gene. Specific amino acids on a surface of the activator, distinct from the DNA interaction surface, are required for the interaction with RNA polymerase.
Figure 1 illustrates the third concept, the activation of transcription by lambda repressor.

The Yeast Transcriptional Activator Gal4 In the early 1980s, we and others had begun analysis of gene activation in a simple eukaryote, the yeast *S. cerevisiae*. Genetic studies had shown that the Gal4 protein is required for transcription of various Gal genes, the products of which metabolize the sugar galactose. (See, for example, 2480.) Addition of galactose to the medium induced transcription of the various *GAL* genes well over a thousand fold. This feature, taken with the ease with which experiments could be done with yeast, encouraged us to study gene activation by Gal4.

Our first important finding was that Gal4 is indeed a specific DNA-binding protein. Guarente and coworkers had shown that a fragment of about 100 base pairs, if positioned some 250 base pairs upstream of the *GAL1* gene, was sufficient to confer galactose regulation on another promoter, that of the gene *CYC1* (2471). They termed the site of Gal4 action the Galactose Upstream Activating Sequence, or UASg.

Using in vivo footprinting techniques and deletion analysis, we found that Gal4 bound specifically to repeated 17-base-pair sequences in UASg (2497, 2498, 2468). At the time, we did not know how Gal4 recognized DNA. The symmetry of each of the sites suggested that the protein bound as a dimer, but the Gal4 protein sequence did not bear the helix-turn-helix motif characteristic of the bacterial DNA-binding regulators known at that time. This contributed to our concern that protein binding to DNA in a eukaryotic nucleus, where it is wrapped in nucleosomes, might require some still undescribed mechanism. The following preliminary yeast experiment shows that this concern was unfounded.

In the mid-1980s, we began to wonder whether this picture from bacteria applied in eukaryotes as well. There were good grounds for thinking it might not. For example, unlike in bacteria, DNA in eukaryotes is wrapped around histone proteins to form nucleosomes, and the DNA is sequestered in a special compartment, the nucleus. Moreover, unlike in the lambda case, eukaryotic regulators often exert their effects over many hundreds, or even thousands, of base pairs. As we shall see, however, the lessons from bacteria stood us in good stead, and taken together with the studies from yeast we now describe, provided a unified view of many aspects of gene regulation, applying to both bacteria and eukaryotes. (2472, 2481, 2471, 3048, 100)
Setting the Stage: Repression by a Bacterial Protein in Yeast

This preliminary experiment was performed with a bacterial repressor called LexA. We and others had shown that LexA binds to well-defined DNA sites, a dimer of the protein recognizing each site (2464, 2479, 2475). Our knowledge was consistent with the idea that DNA-bound LexA works as do other repressors: it turns off transcription of a gene by competing for binding of polymerase to an overlapping promoter. We knew that LexA comprised two domains connected by a flexible linker: an amino terminal domain that contained the DNA-binding function, and a carboxyl terminal domain that contained a dimerization function. Sequence comparison indicated that LexA recognized DNA using a helix-turn-helix motif similar to that found in lambda repressor. Unlike lambda repressor, there was no evidence that LexA activated transcription in prokaryotes; as we now understand, it lacks an activating region.

We placed LexA binding sites between the Gal4 binding sites (UASg) and the start site of transcription of a yeast reporter. We introduced these constructs into yeast along with a second construct (now called an effector construct) expressing the LexA gene under the control of a yeast promoter. LexA repressed transcription from some of these constructs. That is, we found certain positions for the LexA sites at which repression occurred, but others also between the UASg and the gene, at which LexA had no effect (2465).

The results showed that, in yeast, LexA bound its operator sites. Thus, this bacterial protein entered a eukaryotic nucleus and bound its operator sites despite DNA's presumably being wrapped in nucleosomes. The results also argued against so-called "transmission" models for activation by Gal4. According to such models, the activator transmitted some signal (for example, to unwind the DNA) through or along the DNA. Those models predicted that the repressive effects of LexA would not depend on the exact positioning of the LexA bound between the UASg and the gene.

The Key Experiment: The Domain Swap

We thus began with two proteins that bound different DNA sites in yeast. One, the yeast activator Gal4, activated transcription; the other, the bacterial repressor LexA, did not. In bacteria, lambda repressor binds immediately adjacent to the gene it activates, whereas in yeast, Gal4 typically binds some 250 base pairs away. We nevertheless imagined that the role of DNA binding might be the same in both cases, namely, to position the activator near the gene. In that case, we might be able to replace the Gal4 DNA-binding domain with that of LexA and activate transcription of a gene with LexA binding sites nearby.

We constructed and expressed in yeast a chimeric protein, LexA-Gal4. The fusion carried the DNA-binding region of LexA (residues 1-87) fused to a C-terminal portion of Gal4 (residues 74-881) that contained more than 90 percent of the Gal4 coding sequence.

The key result is shown in Figure 2. When LexA-Gal4 was expressed in yeast, it activated transcription of a reporter gene near which we had placed LexA DNA-binding sites. LexA-Gal4 did not activate if the gene lacked LexA binding sites, even if it carried a Gal4 binding site (a "17-mer").
This experiment, whose findings are illustrated in Figure 3, showed the following:

- Gal4 contains an "activating region" that can be transferred, without loss of function, to a heterologous DNA-binding domain.
- No special mode of DNA binding is required for activation in eukaryotes beyond that found in a bacterial repressor.
- Specificity of an activator is determined by its DNA-binding address: Gal4 works if a gene bears a Gal4 binding site, and LexA-Gal4 works if a gene bears a LexA site. The role of DNA binding is to tether the activating region near the gene to be activated.
- The activating region itself does not activate unless tethered to DNA.
- Eukaryotic transcriptional activators are modular proteins, surprisingly easily separated into different functional bits.
The amino portion of Gal4 (missing from the LexA-Gal4 fusion) bears the Gal4 DNA binding domain. A Gal4 fragment bearing that domain (e.g., Gal4 [1-147]) binds Gal4 sites in yeast but does not activate transcription because it lacks an activating region [2474]). This Gal4 fragment is analogous to the so-called "positive control" (pc) mutants of lambda repressor. Those mutants, which bear amino acid changes on the surface of repressor that contacts RNA polymerase, bind DNA normally but do not activate transcription (2479).

Serendipity

Before touching upon some subsequent developments, we point out three bits of good fortune that allowed the success of the LexA-Gal4 domain swap experiment.

LexA turned out to be a good DNA-binding domain in eukaryotes. For unknown reasons (possibly small differences in protein level and/or operator affinity), lambda repressor, the protein we had studied most intensively, did not work in the initial yeast experiments. Had we not been studying LexA in previous years, we would not have had a prokaryotic DNA domain that we knew bound DNA in yeast.

The LexA fragment that was used to make LexA-Gal4 lacked the C-terminal dimerization domain required for efficient DNA binding of LexA. We believed that Gal4 also bound DNA as a dimer and, in those days before the widespread use of synthetic DNA, took advantage of a well-placed restriction site to generate a very large C-terminal fragment of Gal4 fused to the amino part of LexA. Luckily, that carboxyl Gal4 fragment supplied, in addition to an activating region, a dimerization function, but just barely: we now know that the first Gal4 residue in LexA-Gal4 is the first residue of the dimerization region. The ability of the Gal4 dimerization region to effect dimerization (and hence efficient DNA binding) of the LexA DNA binding domain defines another modular, swappable, functional element.

We subsequently learned that LexA and LexA-Gal4 lacked nuclear localization sequences and were not concentrated in the nucleus (2477). Both proteins were functional (even the >90-kiloDalton LexA-Gal4), which showed that such signals were not always required for nuclear entry, contrary to what we had believed up to that time.

Some Subsequent Developments

There were previous examples of protein modularity (for example, the DNA-binding and dimerization domains of lambda repressor). However, the domain swap experiment and the continuing flood of experiments to construct new multifunctional chimeric proteins vividly illustrate just how widespread is the modular principle of protein design.

Gal4, we soon learned, is a "universal" activator: it will work in any eukaryote, provided the test gene is modified by the introduction of Gal4 binding sites nearby and Gal4 is artificially expressed in that organism (2473, 2478). Gal4 is widely used to activate specific genes in specific tissues (e.g., in Drosophila; see 2499). Chimeras that contain other universal activation regions are used to turn genes on and off in response to treatment, for example, with tetracycline (2469).
Universal activating regions typically bear an excess of acidic residues, plus crucial hydrophobic residues. One can readily detect activating regions by fusing candidates to a DNA-binding domain, expressing the hybrid proteins, and assaying for activation of a reporter gene.

Tethering the activating region to the promoter does not require that the activating region be covalently linked to the DNA-binding domain. Bringing the activating region to DNA by a protein-protein interaction triggers activation of transcription. Consider, for example, the protein Gal80, which inhibits Gal4 by binding to and covering Gal4's activating region. Fusion of an activating region con- verts Gal80 to an activator when it binds to DNA-bound Gal4 (2476). Noting this, Fields and Song developed the two-hybrid system (952). That system detects interaction between proteins, one fused to a DNA-binding domain and the other fused to an activating region, by activation of transcription. Transcription by tethered activating regions (as in the two-hybrid system) is now used to map interactions among the protein complements of entire genomes.

Several lines of evidence show that the typical eukaryotic transcriptional activating region works by recruiting the transcriptional machinery to a nearby promoter. One finding consistent with this idea was that, when expressed in large amounts and not tethered to DNA, free activating regions squelched transcription (2467). The properties of activating regions also accord with this surmise. In particular, studies showing eukaryotic activating regions to be flexible in both size and sequence helped spell the end of the idea that the activation function might involve an enzymatic activity. Various lines of evidence suggest that these activating regions can touch an array of targets in the very complicated transcriptional machinery of eukaryotes (over fifty proteins), recruiting whatever is necessary to ensure transcription of a given gene. According to this general picture, the typical eukaryotic activator works by the same fundamental mechanism as does lambda repressor working as an activator: at the heart of each process lies cooperative binding to DNA of activator and target, a process that recruits the transcription machinery to the promoter.

The Legacy

Classical biochemistry taught us that substrate recognition by enzymes (those involved in intermediary metabolism, for example) was determined by highly specific interactions governed by the precise geometries of the interacting moieties. Each such enzyme typically recognizes one and only one substrate, usually a small molecule. Now we see that many enzymes, particularly those involved in regulatory processes, share a property with RNA polymerase: their specificities are not determined solely by their active sites. Rather, just as RNA polymerase has many possible genes it might work on, so too does the typical kinase, phosphatase, protease, etc., have many possible substrates. Which one is chosen, the regulatory choice, is determined in many cases by the kinds of mechanisms we describe here for transcriptional activation.

For example, modular regulatory proteins involved in proteolysis (e.g., F box proteins) direct a ubiquitylating enzyme to specific substrates by binding the enzyme to one surface, and the specific substrates to another. This strategy, imposing specificity on enzyme action by recruitment, figures prominently in the workings of the intracellular pathways that the cell uses to transmit and process information (2500).
The Authors

Mark Ptashne received his Ph.D. from Harvard University, where he was a Junior Fellow. He then became lecturer (1968-71), professor (1971-96), and chairman of the Department of Biochemistry and Molecular Biology (1980-83). Since 1997, he has occupied the Ludwig Chair of Molecular Biology at Sloan-Kettering Institute. He was scientific cofounder of the Genetics Institute in 1980. His many awards and honors include the Eli Lilly Award in Biological Chemistry (1975); le Prix Charles-Leopold Mayer, l'Academie des Sciences, Paris, France (with W. Gilbert) (1977); the U.S. Steel Foundation Award in Molecular Biology (1979); the General Motors Cancer Research Foundation Sloan Prize (1990); and the Lasker Award for Basic Research (1997). His research has concentrated on discerning general principles of gene regulation using phage lambda and yeast as model systems.

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