

Design of Artificial Transcriptional Activators with Rigid Poly-L-proline Linkers

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Abstract: Typical eukaryotic transcriptional activators are composed of distinct functional domains, including a DNA binding domain and an activating domain. Artificial transcription factors have been designed wherein the DNA binding domain is a minor groove DNA binding hairpin polyamide linked by a flexible tether to short activating peptides, typically 16–20 residues in size. In this study, the linker between the polyamide and the peptide was altered in an incremental fashion using rigid oligoproline “molecular rulers” in the 18–45 Å length range. We find that there is an optimal linker length which separates the DNA and the activation region for transcription activation.

Introduction

Transcriptional activators typically bind near a gene and recruit the transcriptional machinery to a nearby promoter, thereby stimulating the expression of the gene. These activators comprise two regions: the DNA binding domain and the activating domain. The former defines the promoter address in the genome where the activating region is to be delivered for recruitment of the transcriptional machinery.^{1,2} We have previously reported efforts to replace the natural protein activators with nonnatural components smaller in size than nature’s proteins. Artificial transcription activators comprised of three synthetic modules, a hairpin polyamide (PA) DNA binding domain (DBD) and a short peptide activation domain (AD) (typically 16–20 amino acid residues in size) connected by flexible linkers which vary in length, have been shown to initiate transcription at targeted promoter sites in cell-free systems.^{3,4} Modeling studies on these polyamide–peptide conjugates with flexible linkers suggested a distance of 20–40 Å between the DNA and the activating region for transcription activation. We sought to determine whether the length of the linker that

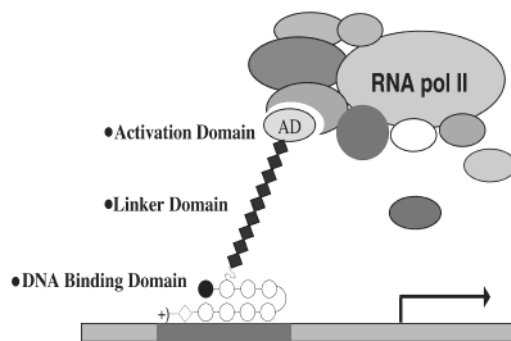


Figure 1. Activation of gene transcription by artificial transcription factors. The artificial activator is composed of three separate functional domains. The DNA binding domain consists of minor groove binding pyrrole/imidazole polyamides. The DNA binding domain is tethered to the activation domain (AD), a peptide, by a linker domain.

projected the activating region away from the DNA would have any affect on the degree of activation (Figure 1). In the present study, we replaced the flexible linkers between the DBD and AD with rigid oligoprolines of varying incremental lengths (18–45 Å). Our results suggest that there is an optimal window within which activating regions of the type used here can function efficiently.

Results

In this study, we replaced the flexible linker between the activating peptide and the hairpin polyamides with 6, 9, 12, and 15 L-proline residues (Pro₆–Pro₁₅) (Figure 2). Poly-L-proline linkers were chosen as the “molecular rulers” for the present study since a stretch of proline residues forms a stable helical structure (the polyproline II helix). Addition of each proline residue increases the length of this helix in a predictable manner, approximately 3 Å per proline residue. Thus, the oligoproline linker projects the activating region peptide away from the DNA

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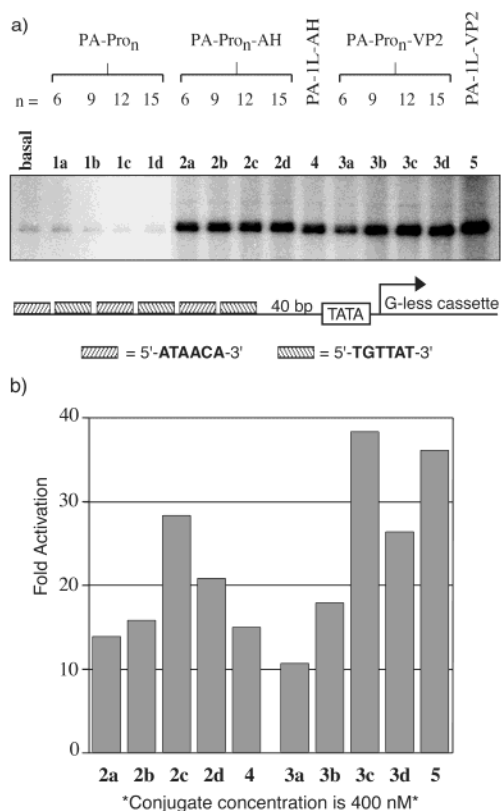


Figure 5. In vitro transcription reactions with the compounds listed in Figure 3. The transcription reaction conditions are described in the Experimental Section. (a) Storage phosphor autoradiogram of the reactions which were performed with a 400 nM concentration of each conjugate. The template configuration showing the sequence and number of the polyamide binding sites is depicted below the gel. (b) Fold activation was determined by comparing the amount of transcription elicited by conjugates 2a–d, 3a–d, 4, and 5 with that of the basal level.

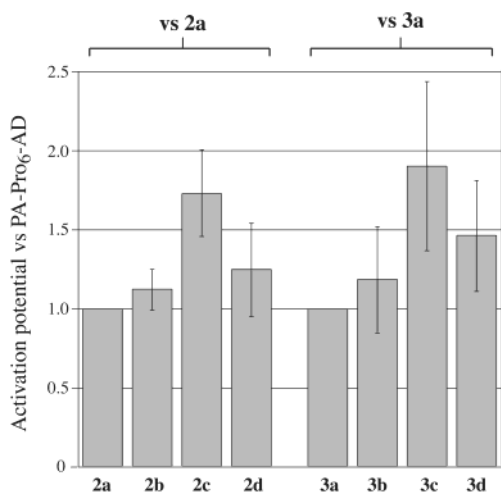


Figure 6. Summary of four independent in vitro transcription reactions showing the relative potency of each compound in comparison to PA-Pro₆-AD (2a, AD = AH; 3a, AD = VP2).

quantitative DNase I footprinting titrations.^{8,9} These studies revealed that conjugates bound with similar affinities and specificities to their sites on the promoter DNA (Figure 7 and Table 1). In control experiments polyamide–proline conjugates

Table 1. Equilibrium Dissociation Constants (nM)

conjugate	K_D (nM)	conjugate	K_D (nM)
PA-Pro ₆ -VP2 (3a)	4.8(±1.4)	PA-Pro ₁₅ -VP2 (3d)	2.3(±1.3)
PA-Pro ₉ -VP2 (3b)	2.0(±1.3)	PA-Pro ₁₂ -AH (2c)	7.7(±1.7)
PA-Pro ₁₂ -VP2 (3c)	3.3(±1.1)		

1a–d lacking the activation transcription peptides did not activate.

The use of poly-L-proline linkers as molecular rulers was inspired by the seminal work of Stryer and co-workers, who utilized fluorescence resonance energy transfer (FRET) analysis to show that oligomers composed of up to 12 proline residues can retain a rigid helical structure.⁵ To ascertain that an oligomer composed of 15 proline residues does not deviate from the predicted structure, we repeated the FRET analysis on poly-L-proline linkers used in our experiments. (See the Supporting Information for details.) To perform this analysis, we synthesized poly-L-proline linkers bearing Oregon Green (OG) energy donor and tetramethylrhodamine (TAMRA) energy acceptor moieties on each end (conjugates 7a–d, Figure S1). In the Stryer study, naphthyl and dansyl groups were used as the energy donor and the energy acceptor moieties, respectively. The different dyes were used in the present study because the Förster radius (the distance at which energy transfer is 50% efficient) for the dansyl–naphthyl pair ($R_0 = 27 \text{ \AA}$) may be too low to allow for efficient FRET when the energy donor and the acceptor are spaced by the Pro₁₅ linker (45 \AA). The Förster radius (R_0) for the OG–TAMRA pair is predicted to be 55 \AA .¹⁰ FRET measurements indicate that the oligomer composed of 15 proline units does retain a rigid structure (Supporting Information, Figure S2a). The plot of distance versus FRET efficiency provides a linear relationship between the conjugates, and importantly, the experimentally observed FRET efficiency is in good agreement with the predicted r^{-6} dependence (Supporting Information, Figure S2b).⁵ Thus, the FRET studies show that the dependence of transcription activation on the number of L-prolines is not due to a change in the linker structure.

Discussion

In this study we examine the role of the distance separating the DBD and AD in our artificial transcription factors. This study was prompted by previous observations in which we found that activating regions activated transcription to different degrees on the basis of the point of attachment as well as the nature of the linker separating them from the DNA binding polyamide.⁴ This dependence on spatial presentation led us to test the role of linker length in the degree of activation elicited by two acidic activating peptides. The proline linker domain is conjugated to an internal pyrrole of the polyamide, rather than to the C-terminus, which allows the linkers to project out of the minor groove away from the DNA helix. Our data suggest that for the two acidic activating regions (AH and VP2) the strength of activation increases as the linker reaches a length of 36 \AA . The increase, while modest, is reproducible and displays a clear trend (Figures 5 and 6). The effects are muted in part due to three features that were incorporated in the experimental design: the use of multiple DNA binding sites to elicit robust activation, the presence of a flexible hinge tethering the proline linker to

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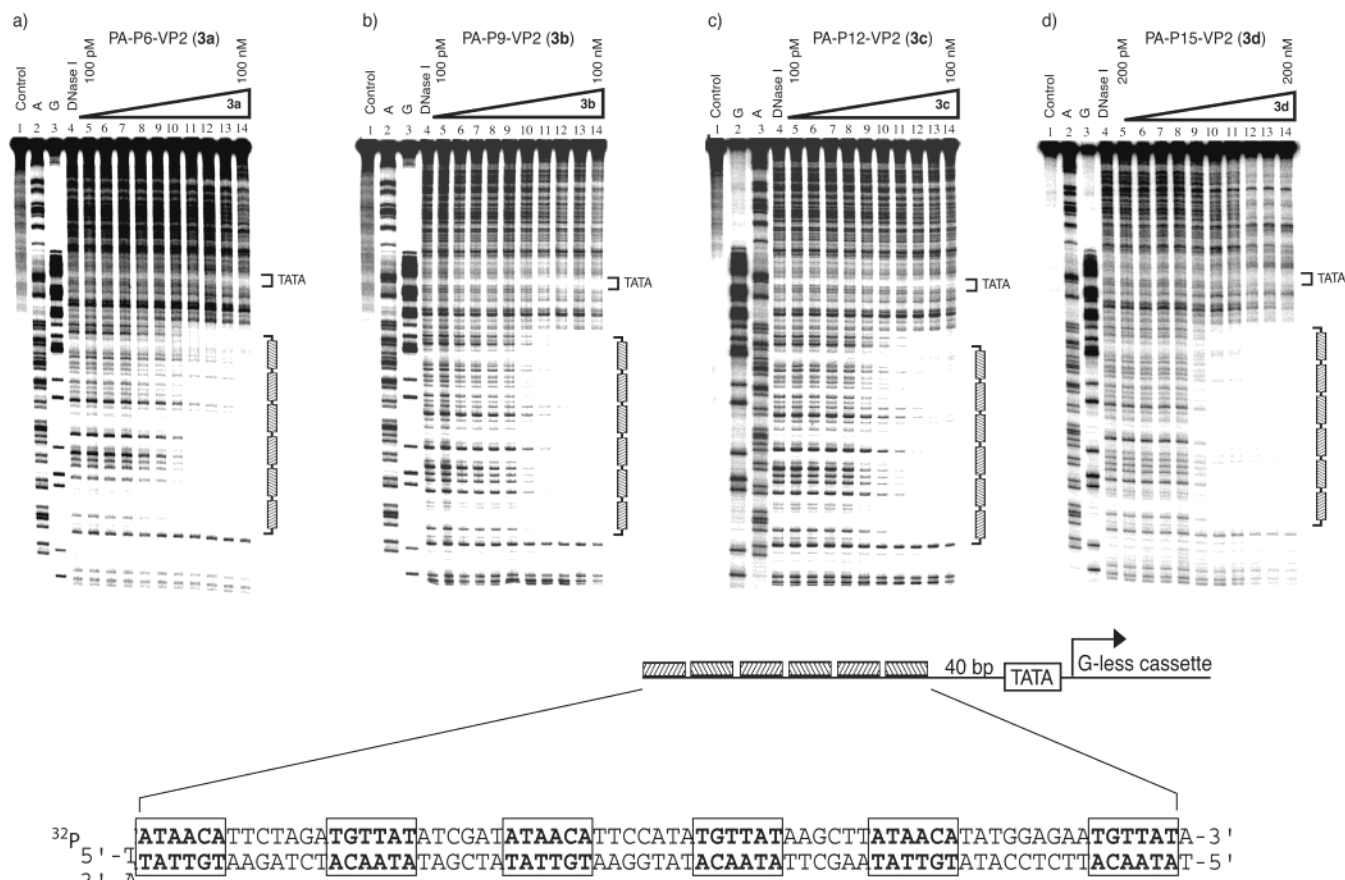


Figure 7. Quantitative DNase I footprinting titration of conjugates **3a–d** shows that all conjugates bound to their target sites with similar affinities. (Top) Storage phosphor autoradiogram of a quantitative DNase I footprinting titration of **3a–d** on a 363 bp $5'$ - ^{32}P -labeled PCR fragment containing both the promoter region and 140 bp of the G-less cassette reporter: (a) lane 1, undigested DNA; lane 2, A reaction; lane 3, G reaction; lane 4, DNase I standard; lanes 5–14, 100 pM, 200 pM, 500 pM, 1 nM, 2 nM, 5 nM, 10 nM, 20 nM, 50 nM, and 100 nM **3a**, respectively; (b) lane 1, undigested DNA; lane 2, A reaction; lane 3, G reaction; lane 4, DNase I standard; lanes 5–14, 100 pM, 200 pM, 500 pM, 1 nM, 2 nM, 5 nM, 10 nM, 20 nM, 50 nM, and 100 nM **3b**, respectively; (c) lane 1, undigested DNA; lane 2, G reaction; lane 3, A reaction; lane 4, DNase I standard; lanes 5–14, 100 pM, 200 pM, 500 pM, 1 nM, 2 nM, 5 nM, 10 nM, 20 nM, 50 nM, and 100 nM **3c**, respectively; (d) lane 1, undigested DNA; lane 2, G reaction; lane 3, A reaction; lane 4, DNase I standard; lanes 5–14, 200 pM, 500 pM, 1 nM, 2 nM, 5 nM, 10 nM, 20 nM, 50 nM, 100 nM, and 200 nM **3d**, respectively. (Bottom) Design of the plasmid DNA template used for the DNase I footprinting titration experiments. The promoter region contains six cognate binding sites for the hairpin polyamide upstream of a G-less cassette reporter.

the polyamide, and the unstructured nature of the activating peptides, which themselves may sample a significant amount of solvent space when fully extended.

We did not find a significant contribution of poly-L-proline linkers themselves to the level of activation in the absence of tethered activating regions. It has been reported that proline-rich (rather than poly-L-proline) activating domains can function to activate transcription even in yeast extracts, though this activation is not as robust as that elicited by acidic activators.¹¹ Presumably the proline oligomer is not a sufficiently strong activator; its effects are therefore not detected in our studies. An alternative possibility is that the ability of proline-rich activating domains to elicit transcription may be more sensitive to their spatial location.

As the principle of recruitment implies, for a DNA-tethered activator to function efficiently, it must sample sufficient nuclear solvent space to bind and recruit various complexes that participate in transcriptional initiation to a given promoter. However, beyond a certain distance the activating region would not function to recruit, as the local concentration of the machinery at a given promoter would not be tremendously

enhanced by binding a distant activating region. It has been shown that eukaryotic activating regions that are not tethered to DNA—even when presented at exceedingly high concentrations such that it would bind to its targets in the transcriptional machinery—do not improve the level of transcription from a given promoter. In fact, as expected, it “squashes” the ability of a DNA-bound activator from functioning—presumably by binding the targets in the machinery.^{2,3,12} Thus, for efficient recruitment in the context of our *in vitro* studies with artificial activators on nonchromatinized templates using yeast extracts, we find that a spacing of 36–45 Å serves as an optimal distance between the DNA and the activating regions to elicit transcriptional activation.

This paper is a step forward toward the goal of engineering and integrating at the molecular level the components for functional artificial transcription factors. The field is at an early stage, and one can imagine other components replacing the design reported here, such as triple-helix-forming oligonucleotides or PNAs tethered to activating peptides.^{13,14} Long-term

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goals for the field would be the replacement of the activating peptide with nonpeptide constructs and the activation (or repression) of endogenous genes in cell culture experiments.

Experimental Section

Synthesis of Polyamide–Peptide Conjugates 1–3. Polyamide thioester **6** was transformed into conjugates **1a–d**, **2a–d**, and **3a–d** by previously reported methods.⁷ The identities of all conjugates were verified by MALDI-TOF mass spectrometry. Characterization: **1a** (PA–Pro₆), MALDI-TOF [M + H] (monoisotopic mass) calcd 2050.6, obsd 2050.0; **1b** (PA–Pro₉), MALDI-TOF [M + H] (average mass) calcd 2341.9, obsd 2341.4; **1c** (PA–Pro₁₂), MALDI-TOF [M + H] (average mass) calcd 2633.3, obsd 2633.1; **1d** (PA–Pro₁₅), MALDI-TOF [M + H] (average mass) calcd 2924.6, obsd 2925.1; **2a** (PA–Pro₆–AH), MALDI-TOF [M + H] (average mass) calcd 4354.5, obsd 4354.8; **2b** (PA–Pro₉–AH), MALDI-TOF [M + H] (average mass) calcd 4646.8, obsd 4647.5; **2c** (PA–Pro₁₂–AH), MALDI-TOF [M + H] (average mass) calcd 4937.5, obsd 4937.5; **2d** (PA–Pro₁₅–AH), MALDI-TOF [M + H] (average mass) calcd 5228.7, obsd 5229.2; **3a** (PA–Pro₆–VP2), MALDI-TOF [M + H] (average mass) calcd 3862.3, obsd 3862.7; **3b** (PA–Pro₉–VP2), MALDI-TOF [M + H] (average mass) calcd 4153.7, obsd 4154.8; **3c** (PA–Pro₁₂–VP2), MALDI-TOF [M + H] (average mass) calcd 4445.0, obsd 4445.3; **3d** (PA–Pro₁₅–VP2), MALDI-TOF [M + H] (average mass) calcd 4736.3, obsd 4736.8.

DNase I Footprinting Titration Experiments. A 363 bp 5′-³²P-labeled PCR fragment was generated from template plasmid pAZA812 in accordance with standard protocols and isolated by nondenaturing gel electrophoresis. All DNase I footprinting reactions were carried out in a volume of 400 μL. A polyamide stock solution or water (for reference lanes) was added to TKMC buffer, with final concentrations of 50 mM Tris–HCl, 50 mM KCl, 50 mM MgCl₂, and 25 mM CaCl₂, pH 7.0, and 15 kcpm 5′-radiolabeled DNA. The solutions were

equilibrated for 12–18 h at 22 °C. Cleavage was initiated by the addition of 10 μL of a DNase I stock solution and was allowed to proceed for 7 min at 22 °C. The reactions were stopped by adding 50 μL of a solution containing 2.25 M NaCl, 150 mM EDTA, 0.6 mg/mL glycogen, and 30 μM base pair calf thymus DNA and then ethanol-precipitated. The cleavage products were resuspended in 100 mM Tris–borate–EDTA/80% formamide loading buffer, denatured at 85 °C for 10 min, and immediately loaded onto an 8% denaturing polyacrylamide gel (5% cross-link, 7 M urea) at 2000 V for 2 h and 15 min. The gels were dried under vacuum at 80 °C and quantitated using storage phosphor technology.

In Vitro Transcription Assays. Template plasmid pAZA812 was constructed by cloning a 78 bp oligomer bearing three cognate palindromic sequences for conjugates **1a–d**, **2a–d**, and **3a–d** into a *Bgl*II site 30 bp upstream of the TATA box of pMLΔ53. This plasmid has the AdML TATA box 30 bp upstream of a 277 bp G-less cassette. For each reaction, 20 ng of plasmid (30 fmol of palindromic sites) was preincubated with a 400 nM concentration of the compound for 75 min prior to the addition of 90 ng of yeast nuclear extract in a 25 μL reaction volume. The reactions were performed as previously described and resolved on 8% 30:1 polyacrylamide gels containing 8 M urea. The gels were dried and exposed to photostimulatable phosphorimaging plates (Fuji Photo Film Co.). The data were visualized using a Fuji phosphorimager followed by quantitation using MacBAS software (Fuji Photo Film Co.).

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Supporting Information Available: Design and synthesis of Oregon Green–Pro_n–TAMRA conjugates for FRET studies, emission spectra, and analysis of FRET data (PDF). This material is available free of charge via the Internet at <http://pubs.acs.org>.

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