Bacterial gene control by DNA looping using engineered dimeric transcription activator like effector (TALE) proteins

Nicole A. Becker, Tanya L. Schwab, Karl J. Clark and L. James Maher, III*

Department of Biochemistry and Molecular Biology, Mayo Clinic College of Medicine and Science, 200 First St. SW, Rochester, MN 55905, USA

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ABSTRACT

Genetic switches must alternate between states whose probabilities are dependent on regulatory signals. Classical examples of transcriptional control in bacteria depend on repressive DNA loops anchored by proteins whose structures are sensitive to small molecule inducers or co-repressors. We are interested in exploiting these natural principles to engineer artificial switches for transcriptional control of bacterial genes. Here, we implement designed homodimeric DNA looping proteins (‘Transcription Activator-Like Effector Dimers’; TALEDs) for this purpose in living bacteria. Using well-studied FKBP dimerization domains, we build switches that mimic features of the natural switches in living bacteria by implementing TALEDs to engineer artificial switches for transcriptional control of bacterial genes. Here, we implement designed homodimeric DNA looping proteins (‘Transcription Activator-Like Effector Dimers’; TALEDs) for this purpose in living bacteria. Using well-studied FKBP dimerization domains, we build switches that mimic features of the natural switches in living bacteria by implementing TALEDs to engineer artificial switches for transcriptional control of bacterial genes. Here, we implement designed homodimeric DNA looping proteins (‘Transcription Activator-Like Effector Dimers’; TALEDs) for this purpose in living bacteria. Using well-studied FKBP dimerization domains, we build switches that mimic features of the natural switches in living bacteria by implementing TALEDs to engineer artificial switches for transcriptional control of bacterial genes. Here, we implement designed homodimeric DNA looping proteins (‘Transcription Activator-Like Effector Dimers’; TALEDs) for this purpose in living bacteria. Using well-studied FKBP dimerization domains, we build switches that mimic features of the natural switches in living bacteria by implementing TALEDs to engineer artificial switches for transcriptional control of bacterial genes. Here, we implement designed homodimeric DNA looping proteins (‘Transcription Activator-Like Effector Dimers’; TALEDs) for this purpose in living bacteria.

INTRODUCTION

Regulated control of gene expression is a key feature of living cells. Classical examples include the Escherichia coli lactose, galactose and tryptophan operons (1–3). These operons are regulated by genetic ‘off’ or ‘on’ switches that prevent or enhance RNA polymerase access to the gene promoter, respectively. Regulatory control typically does not involve changing the concentration of regulatory proteins, but rather changing concentrations of small molecules that induce allosteric changes in these regulatory proteins. We are interested in engineering new gene regulatory switches in bacteria based on natural principles.

DNA looping is exploited as a fundamental principle in bacterial gene repression (4–12). DNA looping represses transcription initiation by at least two mechanisms. First, for bidentate repressor proteins like Lac repressor (LacI), protein saturation of an operator DNA sequence overlapping the RNA polymerase binding site can be increased by increasing the total local repressor concentration by contributions from repressors looping from a distant site (5,11,13,14) compare Figure 1A and B). Second, trapping of a promoter within a strained DNA loop may be intrinsically repressive (15–18). While small molecule-induced allosteric effects in natural gene control switches typically alter the affinity of regulatory proteins for DNA (Figure 1C), here we explore switches where it is protein dimerization that is regulated by the small molecule (Figure 1D).

We describe the design and testing of a regulated gene repression system controlled by dimerization of engineered sequence-specific Transcription Activator-Like Effector Dimers (TALEDs) expressed in living bacteria (Figure 1D). Transcription Activator-Like Effectors (TALEs) are remarkable protein products of plant pathogenic bacteria (19). Bacterial injection of these proteins into hosts is believed to foster infection through transcriptional activation of host genes. TALEs are composed of repeats of 34-amino acid domains carrying a repeat variable diresidue (RVD) encoding specificity for a target base pair. TALE proteins engage the DNA major groove, forming a right-handed protein spiral that binds at a specific DNA target sequence (20,21), bringing a C-terminal transcription activation domain to the protein binding site in chromatin. TALE proteins have previously been fused to nucleases (‘TALENs’) for genetic engineering. Here we replace the TALE activation domain with a dimerization domain [Figure 1E; (19,22)]. In the present work we create new gene control switches in living bacteria by implementing TALEDs to mimic features of the natural lac, gal and trp gene control systems of E. coli. In particular, we borrow from the gal operon switch the concept that repression can be partly dependent on repressor protein dimerization.

*To whom correspondence should be addressed. Tel: +1 507 284 9041; Email: maher@mayo.edu

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Figure 1. Engineering DNA regulatory loops by analogy with natural bacterial repressors. (A) Constitutive ‘off’ switch. ‘On’ state (left) involves an unoccupied operator (grey box) at the promoter (broken arrow) allowing transcription of the downstream gene (filled arrow). ‘Off’ state (right) involves a repressor protein (shown as a tetramer) binding at the operator, preventing promoter access by RNA polymerase. (B) Enhancing effective repressor concentration by DNA looping. Two unoccupied operators (grey boxes) flank the promoter to be regulated (left). The effective concentration of repressor (shown as a bidentate tetramer) at the proximal regulatory operator is the sum of contributions due to free, [R]F, and tethered, [R]L (distal) repressors (right). (C) Inducible ‘off’ switch. ‘Off’ state (left) involves DNA looping anchored by a bidentate tetrameric repressor protein. ‘On’ state (right) is induced by tetramer destabilization into dimers, removing the contribution of tethered (distal) repressors. Residual repression is indicated by grey arrow. (D) Analogous inducible ‘off’ switch involving a designed TALED. A small molecule (small circles) decreases dimer stability. (E) TALED design. TALE protein (gray) with sequence-specific RVD cassettes highlighted and the amino acid sequence of one cassette illustrated (below) with specificity diresidue underlined. DD: dimerization domain. (F) Controlled TALED dimerization. TALE-FKBP(F36M) dimerization variant (left) is broken by FK506 (small circle). Polypeptide polarity is shown below. (G) TALE-FKBP(F36V) dimerization variant (right) is induced by AP20187.

**MATERIALS AND METHODS**

**DNA looping reporter constructs**

DNA looping constructs (Supplemental Table S1) were based on plasmid pJ992 (23), created by modifications of pFW11-null (24,25). See supplemental methods for full details.

**TALE-FKBP protein expression**

Assembly and cloning of genes encoding TALEs has been facilitated by semi-automated methods (26). TALE-FKBP protein expression plasmids were created using modified versions of pJ1034 (low expression promoter) and pJ1035 (moderate expression promoter) (25). Plasmid pJ1035 contains the bacterial UV5 promoter with complete –10 and –35 boxes. The low expression plasmid pJ1034 contains the –10 box of the UV5 promoter with a non-ideal –35 box (24). See supplemental methods for full details.

**E. coli β-galactosidase reporter assay**

LacZ expression was measured using a liquid β-galactosidase colorimetric enzyme assay (27). Repression quantitated in terms of repression ratio (RR):

\[
RR = \frac{U_0 - U_{TALE-FKBP}}{U_0 + U_{TALE-FKBP}}
\]

with the contribution to the repression ratio due to free repressor defined as RRF:

\[
RR_F = \frac{[U_0 - U_{TALE-FKBP}]}{U_0 + U_{TALE-FKBP}}
\]

and the contributions to the repression ratio due to DNA looping defined as RR_L:

\[
RR_L = \frac{RR_T}{RR_F}
\]

See supplemental methods for full details.

**Chemically induced dimerization of FKBP(F36V) variants**

Bacterial stains containing TALE-FKBP(F36V) variants were subcultured in the presence or absence of 1 μM dimerizing agent, AP20187 (Clontech), dissolved in methanol, followed by standard β-galactosidase assay. See supplemental methods for full details.

**Chemical disruption of FKBP(F36M) variant dimerization**

Bacterial strains containing the TALE-FKBP expression plasmids pJ2307 (low protein expression) or pJ2309 (moderate protein expression) were grown in LB medium containing ampicillin. For reporter assay, ~1.2 × 10^7 bacterial cells were subcultured into 1.1 ml LB medium in the presence or absence of 3 μg/ml PMBN (Sigma). Also included at the time of subculture was dimer disrupting FK506 (Sigma), dissolved in DMSO to yield either 1 or 5 μM final concentration. See supplemental methods for full details.
RESULTS

TALED design

Designed TALEDs (Figure 1E and Supplemental Figure S1) were created by modification of a semi-automated method where clusters of modules with proper RVDs are encoded in DNA segments assembled by a Golden Gate procedure (26). In the original assembly system, RVDs are cloned into a plasmid with a C-terminal nuclease in place of the natural transcriptional activation domain. For production of TALEDs, the nuclease domain was replaced with various FKBP modules to facilitate dimerization. Fusion to a C-terminal FKBP(F36M) mutant domain allows constitutive dimerization (Figure 1F) that can be disrupted by small molecule FK506. FKBP(F36M) dimer affinity has been reported to be 30 μM in vitro (28). In contrast, fusion to a C-terminal FKBP(F36V) domain creates monomers whose homodimerization depends on the additional of a small molecule chemical dimerizer such as AP20187 (Figure 1G). In this case, dimer affinity has been reported to be in the nM range in vitro (29). Details are provided in Supplemental Figure S1.

lac looping model systems

We and others have been studying engineered repression by DNA looping using elements of the natural E. coli lac operon (17,18,23,30–32). To create an in vivo model for emulation using designed TALEDs, we assembled components of the lac control switch and analyzed quantitatively their behavior. This design is different from our previous published experiments because gene induction is accomplished by elimination of functional LacI rather than by addition of IPTG inducer. This scheme allows the results to be directly compared to an analogous system where TALEDs substitute for LacI. In one series (Figure 2A), a promoter-proximal pseudo-palindromic operator (O₂) controls RNA polymerase access to the promoter, but is recognized weakly by LacI, while a palindromic operator that strongly binds LacI (O₃ₛₚₚ) is placed at four distances (70.5, 72.5, 75.5, 77.5 bp) upstream. Design details are shown in Supplemental Figure S2A. Resulting measurements of DNA repression in vivo are shown in Figure 2B and interpreted in Figure 2C. The repression ratio (RR; see methods) is defined as the level of expression of the lacZ reporter gene in the absence of any functional LacI (full ‘on’ state), divided by the corresponding level of lacZ expression when LacI is expressed at normal physiological levels (full ‘off’ state). As shown in Figure 2C, repression attributable to free repressor binding at O₂ (RRF) has a value of 3, while increased local repressor concentration due to looping from O₃ₛₚₚ gives RRL as high as 43, increasing repression by a factor of as much as 14, for a total RR of up to 129. For comparison we also studied a lac looping switch where both proximal and distal palindromic O₃ₛₚₚ operators bind LacI strongly (Figure 2D and Supplemental Figure S2B). Here, repression is much more complete and is dominated by the strong binding of free repressor at the proximal operator. Nonetheless, increasing local repressor concentration by DNA looping still contributes a factor of up to 5.4 to overall repression (Figure 2E and F). Thus, these model systems illustrate that optimal operator spacing of natural components of the lac control switch allow DNA looping contributions to repression (RRL) in the range of 5–40 (Supplemental Table S2).

TALED-based lac repression in vivo

To test whether DNA looping by designed TALEDs could lead to an enhancement of gene repression comparable to that caused by looping by LacI, we designed a TALE protein against a 15-bp sequence within the O₃ₛₚₚ operators present in the test construct (Figure 3A). Operator symmetry dictates that this TALE protein can bind in either of two orientations (Figure 3B), unavoidably creating four potential competing DNA repression loops with different geometries (Supplemental Figure S3). Four operator spacing constructs and a control bearing only a proximal operator were tested in living E. coli cells after expression of E. coli LacI (compare Figures 2F and 3D). This is true despite the potential for four competing DNA–protein loops of different geometries for each operator spacing. Im-
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**Figure 3.** A designed TALED represses gene expression by DNA looping. (A) Test system involving Osym operators at four spacings. (B) Designed TALE protein binding sites (gray) and orientations within each symmetrical Osym operator. (C) Measured repression ratios (RR) comparing repression by a single proximal Osym operator with DNA loops of four lengths between distal Osym and proximal Osym. (D) Interpretation of results showing respective contributions of free (RRf) and tethered (RRt) TALEs, whose product is RRT. Here TALED-induced DNA looping enhances repression by 4.4-fold. (E and F) Comparable data for experiments with TALE monomers.

Importantly, significant enhancement of repression is not observed for TALE fusions involving FKBP(F36V) domains, which do not dimerize (Figure 3E and F). Qualitatively similar results were obtained when TALEDs were expressed at lower levels (Supplemental Table S3). Importantly, we also showed that TALED binding to the distal Osym operator at various distances from a promoter lacking a proximal operator had no effect on gene expression (Supplemental Table S4). This result confirms that repression enhancement is due to DNA looping, as designed. Together, these results demonstrate for the first time engineered in vivo transcriptional regulation by DNA looping anchored by a designed TALED.

**Figure 4.** Tuning operator affinity and TALED repression looping. Data (A, C) and interpretation (B, D) for gene repression by designed TALED loops between strong (Osym) distal operators and weak Ob (A, B) and OinvB (C, D) operators.

**Tuning TALED-controlled gene repression**

As shown in Figure 2, quantitative enhancement of gene repression by DNA looping with LacI is a function of operator affinity and loop geometry. To test this in the context of TALEDs, we designed a series of variant operators and tested repression from these operators when they were placed alone in the proximal position in cells expressing the Osym-specific TALED fused to FKBP(F36M). Unlike recognition of palindromic Osym, most of the variant operators are asymmetric and are recognized in only a single orientation. TALED binding to these isolated variant operators indeed led to weaker repression, as designed (Supplemental Figure S4).

We then tested TALED-mediated enhancement of gene repression by DNA looping between distal Osym and either proximal Ob (Figure 4A and B) or proximal OinvB (Figure 4C and D) operators. These configurations involve weaker proximal operators and probably two (rather than four) competing DNA loops (Supplemental Figure S3). Maximal RRt values ranged from 3.4–5.6 in these cases. Qualitatively similar results were obtained when TALEDs were expressed at lower levels (Supplemental Table S5).

**Regulation of gene repression by TALEDs: co-repression or induction by small molecules**

A hallmark of natural gene regulation is triggering by small molecules that increase or decrease repression of gene transcription. We sought to mimic this regulation with TALEDs whose FKBP domain dimerization is sensitive to small molecules. We began with a model of co-repression where...
partial gene repression by an TALE protein monomer bearing FKBPF36V) binding at a weak OB proximal operator (Figure 5A and B) yields an RR_F value of ~23 and a very low RR_L value of ~1.5. This anticipated result is consistent with the lack of FKBPF36V) dimerization in the absence of a chemical dimerizer (Figure 1G). Addition of the cell-permeable chemical dimerizer AP20187 at a concentration found to be effective in E. coli (Supplemental Figure S5) revealed engineered co-repression as designed (Figure 5C and D). The RR_L value in the presence of co-repressor was ~2.3, raising overall repression (RR_F) to ~50 for the optimal operator spacing. Qualitatively similar results were obtained when TALEDs were expressed at lower levels (Supplemental Table S7).

Figure 5. Chemically-induced dimerization triggers TALED repression by DNA looping. Data (A, C) and interpretation (B, D) for gene repression involving designed TALE-FKBP(F36V) repressors without (A, B) or with (C, D) chemically-induced dimerization by 1 μM AP20187.

DISCUSSION

Our focus in this work has been engineered approaches to artificial control of transcription initiation in E. coli. We based our strategy on the DNA looping paradigm common in natural gene control. Here we engineer homodimeric TALEDs for this purpose, demonstrating that DNA looping increases promoter repression to extents comparable to switches involving LacI protein. Exploiting dimerization domains based on FKBP variants, we further show how small molecules can act as co-repressors or inducers in these engineered switches by regulating TALE protein dimerization. This approach is distinct from two elegant prior in vitro studies that did not explore gene control in living cells. Gowetski et al. engineered DNA loops through designed coiled-coil peptides (32). Praetorius and Dietz have recently shown in vitro assembly of complex duplex DNA shapes by DNA looping anchored by designed constitutive dimeric TALE proteins (35). In contrast, our work implements TALEDs in living cells with controlled TALE dimerization regulating gene expression.

In the present study, we have intentionally limited ourselves to adapting cis regulatory elements from the E. coli lac operon where much is known about DNA looping between natural and artificial sequence variants of the palindromic lac operator sequence. Building lac operator-specific TALEDs has the advantage that it allows the engineered system to be directly compared to lac regulation by LacI. On the other hand, the palindromic nature of natural lac operators complicates designed DNA loops anchored by asymmetric TALEDs because multiple competing loop geometries always result. Our data show that this complexity does not prevent the desired functions, but it typically obscures the expected sinusoidal dependence of repression on distance between operators (phasing). We thus attribute the absence of sinusoidal repression dependence on operator phasing to the presence of competing loops (where one loop need not dominate), together with the flexibility of the TALED proteins themselves. An additional complication to interpretation of preferred operator phasings is that the FKBP dimer interface and protein orientations for the FKBPF36M dimer are likely to differ from FKBPF36V after chemical dimerization (28). Nonetheless, for general application the profound advantage of designed sequence-specific TALE-based protein domains will easily
permit specification of unique DNA loops by selecting non-palindromic operator sequences.

Future implementations of the TALED strategy described here can be envisioned to exploit the formation of heterodimeric repressor proteins. When combined with selective heterodimer stabilization or destabilization by available small molecules, such approaches will further generalize microbial applications. Likewise, further development of small molecule triggers to which bacterial cells are highly permeable will further advance application.

It is important to emphasize that regulation of our engineered switches is achieved by controlling the probability of repressor protein dimerization rather than the allosteric control of DNA binding affinity that is pervasive in biological systems. Thus, controlled elimination of DNA looping reduces, but does not eliminate, TALE protein monomer binding at the proximal operator. Overcoming this resulting weak residual repression would involve devising TALE proteins that mimic natural repressors whose DNA affinity can be allosterically controlled by small molecules.

In summary, the present work engineers control of bacterial gene expression by mimicry of natural DNA looping mechanisms using designed repressors whose dimerization can be controlled. It is plausible that similar approaches can be implemented in the context of transcription initiation from eukaryotic promoters to create regulated transcriptional repression by TALED-directed looping of chromatin via arbitrary targeted sequences. Such approaches would greatly extend the previously-recognized principle of eukaryotic gene regulation by DNA looping (36,37).

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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