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Tunable and Multifunctional Eukaryotic Transcription Factors Based on CRISPR/Cas

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ABSTRACT: Transcriptional regulation is central to the complex behavior of natural biological systems and synthetic gene circuits. Platforms for the scalable, tunable, and simple modulation of transcription would enable new abilities to study natural systems and implement artificial capabilities in living cells. Previous approaches to synthetic transcriptional regulation have relied on engineering DNA-binding proteins, which necessitate multistep processes for construction and optimization of function. Here, we show that the CRISPR/Cas system of Streptococcus pyogenes can be programmed to direct both activation and repression to natural and artificial eukaryotic promoters through the simple engineering of guide RNAs with base-pairing complementarity to target DNA sites. We demonstrate that the activity of CRISPR-based transcription factors (crisprTFs) can be tuned by directing multiple crisprTFs to different positions in natural promoters and by arraying multiple crisprTF-binding sites in the context of synthetic promoters in yeast and human cells. Furthermore, externally controllable regulatory modules can be engineered by layering gRNAs with small molecule-responsive proteins. Additionally, single nucleotide substitutions within promoters are sufficient to render them orthogonal with respect to the same gRNA-guided crisprTF. We envision that CRISPR-based eukaryotic gene regulation will enable the facile construction of scalable synthetic gene circuits and open up new approaches for mapping natural gene networks and their effects on complex cellular phenotypes.

KEYWORDS: synthetic transcription factors, CRISPR/Cas9, RNA-guided multiplex gene regulation, synthetic gene regulation, endogenous gene regulation, synthetic biology

Complex and sophisticated phenotypes in eukaryotic cells manifest from layered regulatory networks and specific expression programs involving the regulated transcription of many genes. As major players in these networks, eukaryotic transcriptional factors (TFs) can integrate multiple signals and perform complex, combinatorial functions on promoters, where regulatory information is encoded in the form of binding sites for TFs and interactions between TFs, to modulate gene expression patterns. Rewiring endogenous transcriptional networks by natural or synthetic TFs is a powerful strategy for interrogating cellular functions and controlling cellular phenotypes. Previously, natural DNA-binding domains (DBDs, mainly from bacterial sources, such as TetR, LacI, and LexA) have been used to recruit effector (e.g., activator and repressor) domains to the regulatory regions of eukaryotic genes in order to modulate their transcription. This necessitates the placement of DBD-specific operator site(s) in the cis-regulatory region of the promoters for specific genes which is a labor- and time-intensive process, especially if the regulation of multiple genes is desired. Moreover, engineering and modulating complex transcriptional networks requires tunable, extensible, and orthogonal transcription factors. However, only a few orthogonal variants of natural DBDs are well-characterized and changing their specificity has proven to be challenging. As such, the use of natural DBD-based TFs for wiring complex transcriptional networks and synthetic gene circuits has been limited.

To address these limitations, synthetic TFs based on Zinc Fingers (ZF s) and Transcriptional Activator-Like Effectors (TALEs) have been developed. The ability to program the specificity of ZFs and TALEs to potentially target any sequence makes these DBDs appealing for designing libraries of orthogonal transcription factors. Synthetic ZF- and TALE-based TFs have been shown to work in a wide range of
eukaryotes; however, obtaining a TF for a given target site requires tedious selection processes or multistage DNA assembly protocols. Furthermore, the scale of regulation that can be achieved by these TFs is potentially limited by the metabolic burden imposed on the cells and the number of TFs that can be simultaneously encoded in a given cell.

Here, we present a strategy for modulating eukaryotic transcription at natural and synthetic promoters using programmable and tunable synthetic transcription factors based on a bacterial CRISPR (Clustered Regularly Interspaced Short Palindromic Repeats)/Cas system. Many bacteria use CRISPR-based immune systems to degrade genetic materials of invading phages. Recently, it has been shown that Cas9 can be used as a programmable tool for genome editing across various organisms. In this context, small customizable guide RNAs (gRNAs) can be used to program and target Cas9 endonuclease to specific loci in living cells to induce double (or single)-stranded breaks in DNA. Upon cleavage, error-prone or template-directed repair pathways are triggered, generating variants of the original target loci. Qi et al. showed that an endonuclease-deficient Cas9 (dCas9, with D10A H841A mutations relative to the wild-type Cas9) can be used as a programmable "CRISPRi" tool for gene silencing in Escherichia coli. When targeted to a promoter or ORF of a gene of interest, dCas9 can block progression of RNA polymerase and hence silence expression of the targeted gene. They also provide evidence that CRISPRi is functional in human cells, albeit with much lower efficiency compared with E. coli. In addition, Bikard et al. demonstrated that along with programmed transcriptional repression, transcriptional activation can be achieved in E. coli by fusing the omega subunit of RNA polymerase to the endonuclease-deficient Cas9.

Here, we achieved versatile, programmable, and multiplexable tools for gene regulation in eukaryotes by functionalized...
ing dCas9 with effector domains and targeting both natural and synthetic promoters. As a proof of concept, we made an RNA-guidable transcription factor by fusing dCas9 to an activator domain. Using this CRISPR-based transcription factor (crisprTF), we teased apart the regulatory maps of several natural eukaryotic promoters (in *Saccharomyces cerevisiae* and HEK293T cells) without the need to modify promoter architectures. Unlike previous generations of customizable DBDs (i.e., ZFs and TALEs) that require multistage design and cloning strategies, crisprTFs can be readily customized and retargeted to different loci and regulatory regions in vivo using specific gRNAs with homology to target sites (Figure 1). dCas9 thus offers a powerful tool for targeting functions of interest to specific genomic loci in living cells, which can potentially be used to regulate gene expression at will, construct scalable synthetic gene circuits, or rewire endogenous regulatory networks.

To implement crisprTFs in *Saccharomyces cerevisiae*, we fused the SV40 nuclear localization sequence (NLS) and four tandem copies of Herpes Simplex Viral Protein 16 (VP64, a commonly used eukaryotic transcription activator domain) to a codon-optimized *S. pyogenes* dCas9 (Figure 1A).31 The crisprTF cassette was then cloned under the control of pTPGI, a synthetic promoter which can be induced by growing cells in galactose + anhydrotetracycline (aTc) media23 and integrated into the yeast genome. To assess the activity of crisprTF, *gfp* was placed under the control of a minimal CYC1 promoter (pCYC1m) and the whole cassette was integrated into the yeast genome. pCYC1m retains one of the two endogenous TATA boxes of the wild-type CYC1 promoter and lacks binding sites for endogenous regulatory factors in the upstream activating sequence (UAS).42,43 gRNAs were expressed constitutively from the RNA polymerase III-dependent pRPR1 promoter and the 3′-ends of the gRNAs were defined by the pRPR1 terminator.44

The expression of gRNAs targeting different regions in the pCYC1m (as shown in Figure 1B) resulted in various statistically significant levels of reporter fluorescence compared to the no gRNA control (Figure 1C, left panel). Targeting crisprTFs to sequences upstream of the TATA box (by m1, m2, m6, and m7 gRNAs) resulted in higher *yfp* expression compared with the no gRNA control. Error bars indicate the standard error of the mean for three independent biological replicates. Asterisks (*) on each bar indicate statistically significant changes in *yfp* expression relative to the no gRNA control (based on the two-sided Welch’s t test, *p* < 0.05).

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**Figure 2.** Regulation of *yfp* expression from a minimal MLP promoter (pMLPm) by crisprTFs in HEK293T cells. (A) dCas9_VP64 is expressed in HEK293T cells by the pCMV promoter and directed to target sequences in pMLPm. The mKATE (red) and mBFP2 (blue) fluorophores act as flow-cytometry gating controls for successful plasmid transfections. (B) Map of pMLPm illustrating the relative positions of known regulatory elements. Blue lines indicate target sites for each gRNA. (C) Regulation of *yfp* expression from pMLPm by crisprTFs based on the gRNAs shown in (B). HEK293T cells were cotransfected with the plasmids shown in (A), with specific gRNAs labeled as shown in the x-axis. Targeting crisprTFs to sequences upstream of the TATA box (by m1, m2, m6, and m7 gRNAs) resulted in higher *yfp* expression compared with the no gRNA control. Error bars indicate the standard error of the mean for three independent biological replicates. Asterisks (*) on each bar indicate statistically significant changes in *yfp* expression relative to the no gRNA control (based on the two-sided Welch’s t test, *p*-value < 0.05).
the TATA box and the transcription start site (using c2 and c6 gRNAs), likely due to interference of crisprTFs with the formation of the transcriptional initiation complex.\textsuperscript{56} No activation was observed with any of the eight tested gRNAs when dCas9, without a fused activator domain, was targeted to pCYC1m (Figure S1, Supporting Information). All the tested gRNAs in this strain repressed \textit{gfp} expression to some extent (Figure S1, Supporting Information), with highest repression observed with c6 and c7. These results demonstrate that dCas9 is able to repress transcription but requires an activation domain (VP64) to activate transcription of a target locus and further supports the hypothesis that dCas9 (or as a fusion to VP64) can act as a repressor by interfering with the formation of the transcriptional initiation complex. Similar results were achieved with the GAL1 promoter (pGAL1) and its variants, where targeting crisprTFs to sequences upstream and downstream of TATA box led to activation and repression of the GFP reporter, respectively (Figure S2, Supporting Information). These results indicate that a single crisprTF can be programmed to act as both an activator and a repressor by targeting it to different positions across endogenous promoters.

The activity of a promoter is determined by combinatorial interactions between transcriptional regulatory factors bound to that promoter. We thus investigated the effects of binding of multiple crisprTFs targeted to the same promoter. To this end, pairwise combinations of gRNAs with non-neutral effects (those that showed either activation or repression in the left panel of Figure 1C) were coexpressed. As shown in the right panel of Figure 1C, coexpression of repressor gRNAs resulted in synergistic repression of the reporter (up to 7× repression was achieved with coexpression of the c5 and c6 pair as well as the c6 and c7 pair). On the other hand, when a repressor gRNA was coexpressed with an activator gRNA (e.g., the c3 and c6 pair), an intermediate level of GFP expression was achieved, indicating an antagonistic interaction between the two gRNAs. Moreover, the effects of repressor gRNAs were dominant over activator gRNAs, suggesting that interruption of the formation of the transcription initiation complex has a stronger effect than activation.\textsuperscript{46} Coexpression of two activator gRNAs (e.g., c3 and c4) did not result in synergistic activation of the reporter, which suggests that the relative positions and interactions of bound activators are important for determining synergistic activation.\textsuperscript{46} Consistent with our results, it has been shown that synergistic activation from synthetic promoters with multiple GAL4 operator sites depends on the distance and helical phase of the operator sites.\textsuperscript{57} Furthermore, in another study it has been shown that not all of the combinations of TALE-activators targeted to the same promoter result in synergistic activation.\textsuperscript{25}

We next sought to investigate the activity of crisprTFs in human cells. To this end, a human-codon-optimized crisprTF cassette was placed on a plasmid under the control of the constitutive cytomegalovirus immediate-early promoter (pCMV). The gRNAs were expressed constitutively from a separate plasmid by the RNA polymerase III-dependent U6 promoter (pU6), as previously described\textsuperscript{47,48} (Figure 2A). After transfection of these plasmids into HEK293T cells, we investigated the regulatory architecture of the minimal adenovirus major late promoter (pMLPm)\textsuperscript{49} by targeting crisprTFs to different positions across this promoter (Figure 3A). YFP was used as the readout for pMLPm promoter activity.

Consistent with the results obtained in \textit{S. cerevisiae}, crisprTFs activated gene expression when targeted to sequences upstream of the pMLPm TATA box (using m1, m2, m6, or m7 gRNA) or downstream of the transcription start site (m8 gRNA) (Figure 2B and C). Since the basal expression level of the pMLPm promoter is low, it was challenging to detect significant repression from this promoter. Thus, to demonstrate that crisprTFs can function as transcriptional repressors in mammalian cells, we placed mKATE under the control of a constitutive mammalian promoter, phosphoglycerate kinase 1 (pPGK1),\textsuperscript{50} and targeted crisprTFs to this promoter (Figure 3A). pPGK1 is a strong, constitutive, TATA-less promoter that contains a CCAAT box\textsuperscript{51} and five GC-boxes.\textsuperscript{52} These sites are the binding sites for the endogenous human transcription factors CBP and SP1, respectively. Targeting dCas9 alone, dCas9 fused to VP64 domain, or dCas9 fused to KRAB domain to the CCAAT box or the GC-boxes resulted in significant repression of the reporter gene (Figure 3B), presumably by preventing endogenous transcription factors from binding to specific DNA recognition elements\textsuperscript{51,52} within the pPGK1 promoter.

We further sought to explore the tunability of crisprTFs in the context of synthetic promoters. In order to do so, we engineered multiple artificial binding sites (operators), pCYC1m separated by 20 base-pair sequences, upstream of the pCYC1m promoter in \textit{S. cerevisiae} (Figure 4A). Expression of a gRNA, which targeted these arrayed operator sites resulted in synergistic activation of the \textit{bfp} reporter (Figure 4B). Higher levels of \textit{bfp} expression (up to 70-fold activation with 12× gRNA operator sites) were achieved by increasing the number of gRNA binding sites upstream of the engineered pCYC1m. This level of activation in yeast is comparable to the activation reported for commonly used endogenous yeast promoters (e.g.,

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\caption{CrisprTF-mediated repression of the constitutive pPGK1 promoter in HEK293T cells. (A) Map of the pPGK1 promoter illustrating the relative positions of known regulatory elements. (B) CrisprTF-based targeted repression of the constitutive pPGK1 promoter. Constructs expressing different dCas9-based proteins (dCas9, dCas9-VP64, and dCas9-KRAB) were cotransfected with plasmids containing pPGK1_mKATE and constructs expressing no gRNAs or gRNAs targeting the CCAAT box or the GC-box gRNA. Significant repression of the pPGK1 promoter relative to the no gRNA control was observed with all of the three different dCas9 constructs (dCas9, dCas9-VP64, and dCas9-KRAB). Error bars indicate the standard error of the mean for three independent biological replicates.}
\end{figure}
pGAL153 and pCUP154) and synthetic promoters that are orthogonal with respect to each other, we tested three gRNAs (cm1, cm2, and cm5) that were designed to match the modified target sites. However, this modified promoter responded to a new set of gRNAs (cm1, cm2, and cm5) that were designed to match the modified target sites. The wild-type pCYC1m promoter did not respond to the new set of gRNAs (cm1—cm6) (Figure 6B, left panel). These experiments demonstrate that as little as a single base-pair mismatch is sufficient to direct the crisprTF to one locus while preventing activity at another locus.

To further demonstrate the potential of crisprTFs toward constructing synthetic promoters and gRNAs that are orthogonal with respect to each other, we tested three randomly designed gRNAs (a1, a2, and a3 gRNAs) for their ability to activate each other’s target sequences. As shown in Figure 6C, each of the gRNAs exhibited high activity at their cognate target sequences but low activity at noncognate sequences. These results suggest that one can construct synthetic promoters and gRNAs that are orthogonal with respect to each other and to the host genome, especially within eukaryotes with smaller genomes, such as yeasts.

During the course of the peer-review for this work, similar systems for transcriptional control in eukaryotic cells were described. Gilbert et al. demonstrated that CRISPR-mediated gene repression and activation can be achieved in both yeast and mammalian cells by using fusions of dCas9 with repressor and activator domains respectively. Furthermore, Maeder et al. and Perez-Pinera et al. showed synergistic CRISPR/Cas-based gene activation in human cells with multiple gRNAs. In this paper, we additionally show that both activation and repression can be achieved in human cells with multiple gRNA operators.
repression functions can be achieved with only one transcription factor, by targeting dCas9_VP64 fusions to different regulatory sequences along a promoter. Our finding that one can activate or repress the expression of a gene of interest by directing a single protein to different positions of a promoter is advantageous for the efficient design of synthetic transcriptional networks or rewiring natural ones. This property obviates the need for using separate orthogonal Cas9 protein fusions as activators and repressors. Furthermore, we demonstrate that more sophisticated regulatory motifs, such as small-molecule responsive modules, can be built for crisprTFs, thus enabling external control of crisprTF-based transcriptional circuits. Such synthetic modules can be interfaced with other regulatory elements to achieve more complex regulation for synthetic biology.

Our results show that dCas9 can be used as a customizable RNA-guided DNA-binding platform for the regulation of gene expression at natural and synthetic promoters in eukaryotic cells. The ease of design and expression of customized gRNAs in comparison to ZFs and TALEs make CRISPR-based transcription factors appealing as synthetic TFs for modulating endogenous gene expression as well as for synthetic biology. The ability to customize the target site of dCas9 via the expression of short gRNAs obviates the need to engineer multiple orthogonal DBDs in order to construct complex transcriptional circuits. This could potentially reduce the overall metabolic burden on cells and enable the integration of more complex synthetic computation and logic within living cells.62,63 More complex regulatory and logic circuits, such as cascades and complex digital logics gates can be built by layering crisprTFs. The possibility of integrating multiple inputs at a single promoter expands the regulatory potential and provides us with increased flexibility that can be leveraged while designing synthetic transcriptional networks or rewiring endogenous pathways.

Furthermore, since both activation and repression functions can be achieved with crisprTFs, the crisprTF platform may be advantageous compared to noncoding RNA-based gene regulatory platforms where only repression can be achieved. In a way, crisprTFs combine the multiplexability of RNA-based regulatory approaches with the flexibility and rich functionality repertoire of protein-based gene regulatory approaches: Cas9 can be functionalized with regulatory domains of interest (e.g., activation, repression, or epigenetic effector) and then be targeted to multiple loci using different gRNAs.

Future work is needed to define the range of effector domains that can be used with dCas9 for a variety of regulatory functions, including transcriptional regulation and epigenetic modifications. In addition, the identification, characterization, and optimization of Cas9 homologues or evolved variants may enable enhanced activity and specificity of this system. Moreover, the ability to synthesize random libraries of gRNAs opens the possibility for high-throughput perturbations of transcriptional networks and screening for desirable phenotypes. Ultimately, we envision that crisprTFs will enable the regulation and perturbation of natural transcriptional networks as well as the construction of complex synthetic circuits at an unprecedented speed and scale.

## METHODS

### Strain and Plasmid Construction.

*Saccharomyces cerevisiae.* dCas9 (endonuclease-deficient Cas9, with D10A and H841A mutations relative to the wild-type sequence of *S. pyogenes* Cas939) with an N-terminal SV40 nuclear localization signal (NLS) was codon-optimized for expression in *S. cerevisiae* and cloned into a pRS314 backbone under control of the pTPGI promoter.23 The RNA-guided transcription factors (crisprTFs) were built by fusing four repeats of the minimal dCas9 domain of the herpes simplex viral protein 16 (VP16) to the C-terminal of dCas9 (dCas9_VP64). The crisprTF-expressing plasmid was then integrated into the TRP1 locus of *S. cerevisiae* W303.

The reporter plasmids were built by cloning yeast-enhanced gfp under the control of the wild-type or modified pCYC1m promoter into pRS406 using one-step Gibson assembly. The reporters for the multiple-gRNA-binding-site experiment (Figure 4A) were built by cloning the corresponding number of binding sites upstream of the pCYC1m promoter driving production of EBFP2. All reporters were integrated into the BLA1 locus of the integrated crisprTF plasmid.

To build gRNA-expressing plasmids, empty gRNA expressing vectors were first made by cloning the pPR1 promoter (an RNA-polymerase-III-dependent promoter54), the gRNA handle (flanked by HindIII and XhoI sites), and the RPR terminator by targeting dCas9_VP64 fusions to different regulatory sequences along a promoter. A TetR operator site (1xTetO) was placed in the pRPR1 promoter to make an aTc-responsive pRPR1_TetO promoter. Addition of aTc releases TetR-mediated repression on the pRPR1_TetO promoter and results in a1_gRNA expression. (A) Schematic of the aTc-inducible pRPR1_TetO promoter. Expression of dCas9_VP64 is driven by the galactose-inducible pGAL1 promoter. A TetR operator site (1xTetO) was placed in the pRPR1 promoter to make an aTc-responsive pRPR1_TetO promoter. Expression of dCas9_VP64 is driven by the galactose-inducible pGAL1 promoter. A TetR operator site (1xTetO) was placed in the pRPR1 promoter to make an aTc-responsive pRPR1_TetO promoter. Addition of aTc releases TetR-mediated repression on the pRPR1_TetO promoter and results in a1_gRNA expression. (B) aTc-dependent bfp expression from a synthetic 6x(a1_op)_pCYC1m promoter. S. cerevisiae cells containing the circuit shown in (A) were grown in galactose media with either 250 ng/mL aTc or no aTc. Error bars indicate the standard error of mean for three biological replicates.
Figure 6. Constructing orthogonal crisprTF-responsive promoters. (A) A schematic view of gRNAs targeting the wild-type and modified pCYC1m promoters in yeast. Only the c1–c8 gRNAs have perfect homology to the target sequences in pCYC1m. On the other hand, c1, c4, c5, and cm1–cm6 gRNAs have perfect homology to the sequences in the pCYC1m(modified) promoter. Mismatches between pCYC1m and pCYC1m(modified) are marked by asterisks (*). (B) pCYC1m only responds to gRNAs that are perfectly matching gRNAs (c1–c8 gRNAs) and not to those that contain mismatches (cm1–cm6 gRNAs). The pCYC1m(modified) promoter responds to the cm1–cm6 gRNAs. Those gRNAs that bind to the sequences upstream of the TATA boxes activate gfp expression and those that target sequences downstream of the TATA boxes repress gfp expression. The c1 and c5 gRNAs have similarly neutral effects on both the wild-type and modified promoters. Error bars indicate standard error of the mean for three independent biological replicates. Asterisks (*) on each bar indicate statistically significant activation or repression relative to no gRNA controls (based on the two-sided Welch’s t test, p-value < 0.05). (C) Heat map illustrating the orthogonality of crisprTFs in human cells. Plasmids encoding three orthogonal gRNAs (a1, a2, and a3 gRNAs) were cotransfected into HEK293T cells along with one of the three reporter plasmids (each encoding 4x operator sites for a given gRNA) upstream of pMLPm promoter driving yfp expression. Only cognate interactions between gRNAs and target binding sites resulted in significant activation. The standard error of the mean of YFP fluorescence for three independent biological replicates is indicated in each cell of the heat-map plot.
into the Sacl and KpnI sites of either the pRS423 or pRS425 plasmid using one-step Gibson assembly. The specificity determinant sequence (SDS) for each gRNA was then cloned into the HindIII site of these vectors by one-step Gibson assembly. Sequences of the constructs used in this study are listed in Table S1, Supporting Information.

**HEK293T Cells.** To construct the mammalian dCas9::VP64 expressing plasmid, we first introduced D10A and H841A mutations into hCas937 (Addgene, Plasmid #41815). Then, three repeats of SV40 NLS (3xNLS) were fused to the C-terminus of the mutated hCas9 using a PCR-based assembly protocol. Using a multipart Gibson assembly protocol, the immediate-early promoter of cytomegalovirus (pCMV), dCas9_3xNLS, VP64, and SV40 polyaT terminator were cloned into the NotI site of the pGS-Luc plasmid (Promega). To monitor successfully transfected cells by flow cytometry, we replaced the original luciferase gene in pGS-Luc with mkATE (Evrogen). The resulting pPGK1::mkATE cassette served as a constitutive fluorescent protein control that was used to gate for the presence of the crisprTF-expressing plasmid with flow cytometry.

The gRNA expression plasmids were constructed by cloning the 138 bp human U6 promoter (an RNA-polymerase-III-dependent promoter48), along with the gRNA handle and terminator into a plasmid containing pPGK1-eBFP2 flanked by the SV40 polyaT terminator (a gift from Lior Nissim). A Sacl site was placed at the 3′-end of the U6 promoter to enable the cloning of different specificity determining sequences for each gRNA. The reporters were assembled into the gRNA-expressing plasmid through a one-step Gibson assembly reaction, where the upstream polyadenylation signal and transcriptional pause site from pG5-Luc, along with a 41 bp, minimal adenovirus type 2 major late promoter (pMLPm), mYFP, and HSV polyA signal were cloned into the AatIII site of the gRNA-expressing plasmids.

For the synthetic promoter experiments, additional gRNA operator sites were cloned in the Nhel site upstream of the pMLPm promoter (see Supporting Information). For the repression experiments, dCas9::KRAB was constructed by cloning a 366 bp KRAB domain to the C-terminus of dCas9. GCCACC was used as the Kozak sequence for the expression of dCas9::VP64, mYFP, eBFP2, and mkATE.

Unless directly targeted by gRNAs for repression assays, the mkATE fluorescent protein on the crisprTF-expression plasmid and the eBFP2 fluorescent protein on the reporter/gRNA plasmid served as our gating controls for flow cytometry analysis.

**Fluorescence Assays.** To assess expression of the reporter constructs, yeast cells expressing different gRNAs (or no gRNA as control) were grown overnight (900 rpm, 30°C) in 96-deep-well plates in yeast minimal media supplemented with glucose with appropriate selection (three independent cultures for each sample). Ten microliters of these cultures were then transferred into fresh media supplemented with galactose +250 ng/mL anhydrotetracycline (aTc) and grown for 20 h (900 rpm, 30°C) before analysis by flow cytometry.

For the human cell culture experiments, HEK293T kidney epithelial cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% (v/v) fetal bovine serum (FBS), 1% glutamine, and 1% penicillin/streptomycin. Cells were grown under 5% CO2 at 37 °C. HEK293T cells were transfected with Fugene-HD transfection reagent (Promega) and assayed for gene expression with flow cytometry at 48 h post transfection.

An LSR Fortessa II flow cytometer equipped with 405 nm, 488 nm, and 561 nm lasers was used for all the experiments. GFP/YFP, BFP, and mKATE levels were detected using 488/FITC, 405/Pacific-Blue, and 561/TX-red laser/filter sets, respectively. All samples were uniformly gated by forward and side scatter. Additional gating for the presence of red and blue fluorophores was applied to the HEK293T samples to ensure only cells successfully transfected with both the crisprTF and the reporter/gRNA plasmids are analyzed. For each gated sample, the mean fluorescence per cell was calculated. Three independent biological samples were used to calculate the mean and standard error of the mean for each data point.

**ASSOCIATED CONTENT**

**Supporting Information**

This information is available free of charge via the Internet at http://pubs.acs.org.

**AUTHOR INFORMATION**

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**Author Contributions**

F.F. built and tested the yeast crisprTF system. S.D.P. built and tested the human crisprTF system. F.F., S.D.P., and T.K.L. designed experiments, analyzed data, and wrote the paper.

**Notes**

The authors declare no competing financial interest.

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