Effective CRISPRa-Mediated Control of Gene Expression in Bacteria Must Overcome Strict Target Site Requirements

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Abstract

In bacterial systems, CRISPR-Cas transcriptional activation (CRISPRa) has the potential to dramatically expand our ability to regulate gene expression, but we currently lack a complete understanding of the rules for designing effective guide RNA target sites. We have identified multiple features of bacterial promoters that impose stringent requirements on bacterial CRISPRa target sites. Most importantly, we found that shifting a gRNA target site by 2-4 bases along the DNA target can cause a nearly complete loss in activity. The loss in activity can be rescued by shifting the target site 10-11 bases, corresponding to one full helical turn. Practically, our results suggest that it will be challenging to find a gRNA target site with an appropriate PAM sequence at precisely the right position at arbitrary genes of interest. To overcome this limitation, we demonstrate that a dCas9 variant with expanded PAM specificity allows activation of promoters that cannot be activated by S. pyogenes dCas9. These results provide a roadmap for future engineering efforts to further expand and generalize the scope of bacterial CRISPRa.
Introduction

Developing tools to activate the expression of arbitrary genes has been transformative for biotechnology and biological research\(^1\). In metabolic engineering, regulating the timing and levels of the expression of complex multi-gene pathways is critical for reducing cellular burden and improving production of valuable metabolites\(^2\). To enable these goals, we recently developed a new CRISPR-Cas transcriptional activation (CRISPRa) system that is effective in *E. coli*. Our system can be combined with CRISPRi gene repression to programmably and flexibly target multiple genes for simultaneous activation and repression\(^3\). While our CRISPRa system can be used with heterologous genes, an outstanding challenge is to understand the rules that define effective target sites at arbitrary promoters in the genome.

To programmably target genes, we use nuclease defective Cas9 (dCas9) with a guide RNA (gRNA) that specifies a target site on the DNA. Targeting this complex to a promoter or an open reading frame (ORF) results in gene repression (CRISPRi)\(^4\). To enable simultaneous activation, we use modified guide RNAs, termed scaffold RNAs (scRNAs), that include a 3’ MS2 hairpin to recruit a transcriptional activator fused to the MS2 coat protein (MCP)\(^3\). We can express multiple gRNAs and scRNAs to inhibit and activate genes simultaneously; gRNAs targeted to a promoter or ORF result in CRISPRi and scRNAs targeted to an appropriate site upstream of a minimal promoter result in CRISPRa.

We demonstrate here that the rules for targeting CRISPRa to effective sites in *E. coli* are surprisingly stringent. In prior work, we found that CRISPRa in *E. coli* was effective at target sites located in a narrow 40 base window between 60 and 100 bases upstream of the transcriptional start site (TSS)\(^3\). Here, we show that multiple factors combine to make the requirements for effective sites even more strict. We demonstrate that the basal promoter strength of the target gene and the sequence composition between the target site and the minimal promoter can have dramatic effects on gene activation. Further, by scanning the 40 base window at single base
resolution, we find sharp peaks of activity and broad regions of inactivity that occur in a periodic 10-11 base pattern, corresponding to one helical turn along the DNA target. The observation that only a few precisely-positioned target sites upstream of the TSS are effective for CRISPRa poses a significant challenge, as many genes will likely lack an NGG PAM sequence at exactly the right position necessary for *S. pyogenes* dCas9. These stringent requirements may explain why CRISPRa and other tools for gene activation in bacteria have lagged far behind comparable tools in eukaryotic systems, where such strict target site requirements are absent.

Although the requirements for bacterial CRISPRa target sites pose challenges, our data also demonstrate CRISPRa has the potential to be effective at a broad range of target genes. In addition to $\sigma^{70}$-dependent genes, CRISPRa can activate expression from genes that use the $\sigma^{70}$ family members $\sigma^{38}$, $\sigma^{32}$, and $\sigma^{24}$, which covers a broad range of the *E. coli* genome. We further demonstrate that the strict requirement for a precisely positioned PAM site can be partially overcome using a re-engineered dCas9 protein that targets an expanded set of PAM sequences. Similar results were recently reported for an alternative bacterial CRISPRa system that can target genes regulated by $\sigma^{54}$ promoters, which were not activated by our system. The availability of these complementary systems should further extend the scope of bacterial CRISPRa. More broadly, by systematically defining the rules for effective CRISPRa sites, we have identified new strategies for improving and generalizing synthetic gene regulation in bacteria.

**Results**

**A SoxS double mutant reduces off-target activation of endogenous SoxS-responsive promoters**

Ideally, a synthetic transcriptional activator should only activate its programmed target genes. The activation domain for our CRISPRa system is SoxS, a native *E. coli* transcription factor that directly binds DNA and activates endogenous gene targets as part of a stress response program. We previously demonstrated that point mutations in the SoxS DNA binding site can reduce activation of endogenous SoxS targets while maintaining CRISPRa activity at a
heterologous reporter gene. However, the most effective single point mutants, R93A and S101A, did not completely abolish activity at endogenous targets. To further minimize off-target SoxS activity, we tested a double mutant SoxS(R93A/S101A). This double mutant SoxS retained full CRISPRa activity and showed a reduction in endogenous SoxS-dependent gene expression to levels indistinguishable from background (Figure 1). Thus, SoxS(R93A/S101A) is an effective modular transcriptional effector that can activate gene expression only when recruited to a target gene via the CRISPR-Cas complex.

A simple distance-based metric for target site selection is not sufficient for effective CRISPRa at endogenous gene targets

To determine if we could predictably activate endogenous genes with CRISPRa, we selected three candidate genes with appropriately positioned PAM sites upstream of the TSS. Previously, we demonstrated that CRISPRa can activate heterologous promoters with target sites positioned within a 40 base window between 60 and 100 bases upstream of the transcriptional start site (TSS)\(^3\). We therefore targeted the CRISPR-Cas complex to the same window upstream of the candidate target genes. First, we targeted the aroK-aroB operon, which expresses enzymes involved in aromatic amino acid biosynthesis, whose programmed overexpression could be useful for bioproduction\(^9\). Targeting the CRISPR-Cas complex to two sites within the optimal 40 base window resulted in up to 5-fold increases in expression, but these effects were highly variable between replicates and most were far smaller. Further, sites inside and outside of the 40 base window gave similar effects (Figure 2A). Next, we targeted cysK, an enzyme involved in cysteine biosynthesis\(^10\). Similar to what we observed with aroK-aroB, targeting a site positioned at -93 from the TSS resulted in expression levels that were highly variable between replicates, ranging from 2.4-fold to 0.8-fold compared to a control expressing an off-target scRNA (Figure 2B). Targeting two other sites within the 40 base window, at -79 and -61, resulted in similarly variable and, in some cases, decreasing levels of expression. Finally, we targeted ldhA, an enzyme involved in
mixed acid fermentation\textsuperscript{11}. We selected 8 sites and observed small increases in gene expression up to 1.5-fold, with no apparent relationship between the position of the target site and \textit{ldhA} expression (Supplementary Figure 1). Together, these results suggest that endogenous genes cannot be activated simply by targeting the CRISPR-Cas complex to sites positioned between 60 and 100 bases upstream of the TSS. The levels of activation we achieved were modest compared to the 50-fold effects seen by RT-qPCR at heterologous promoters\textsuperscript{3}. Even for sites that had modest effects, their positions did not correspond to the predicted optimal sites based on the results from heterologous promoters.

There are several possible explanations for our inability to activate endogenous bacterial genes with CRISPRa. First, we originally demonstrated CRISPRa using a relatively weak synthetic promoter. The basal levels of expression of endogenous genes vary significantly\textsuperscript{12}, and it may be difficult to increase the transcription of genes that are already strongly expressed\textsuperscript{13}. In addition, some endogenous target genes might require an alternative sigma factor. Our original reporter gene is controlled by the $\sigma^{70}$ housekeeping sigma factor, and we do not know if our CRISPRa system is effective at gene targets that use alternative sigma factors. Another possibility is that native transcriptional regulator binding sites near endogenous gene promoters could disrupt CRISPRa. Finally, the optimal distance window metric that we previously identified may have been oversimplified. We initially identified the optimal window from an experiment with target sites spaced 10 bases apart, which may not be sufficient to generalize to any site within the 40 base window. To systematically explore these possibilities, we proceeded to test the efficacy of CRISPRa with a new set of synthetic promoters engineered with variable basal expression levels, alternative sigma factors, variable regulator binding sites, and variable scRNA target site positions.

**CRISPRa is sensitive to promoter strength**

To evaluate whether the intrinsic strength of the promoter affects CRISPRa, we tested activation on a set of fluorescent reporter genes with minimal promoters spanning a 200-fold
range in basal expression level (http://parts.igem.org) (Figure 3A). We observed the most effective gene activation with a moderately weak J23117 promoter. With the weakest promoters, we could not detect any activation, even though their basal expression levels were only 2-fold weaker than the J23117 promoter. With stronger promoters, we observed progressively smaller CRISPRa-mediated activation of gene expression; the basal expression level increased, while the maximal, CRISPRa-induced expression remained roughly constant. These results indicate that the bacterial CRISPRa activity varies considerably with promoter strength, similar to effects observed in eukaryotic systems\textsuperscript{14,15}. Thus, when targeting arbitrary endogenous genes, the level of activation that can be achieved may depend on the basal level of expression of its promoter.

**CRISPRa is effective with alternative sigma factors**

Bacterial transcription is initiated by a sigma factor binding to the minimal promoter and the RNA polymerase holoenzyme\textsuperscript{16}. The SoxS activator binds directly to the $\alpha$ subunit of RNA polymerase\textsuperscript{17}, which suggests that our CRISPRa system could be compatible with genes that are controlled by non-housekeeping sigma factors. To investigate this possibility, we built synthetic promoters regulated by $\sigma^{38}$ (RpoS), $\sigma^{32}$ (RpoH), $\sigma^{24}$ (RpoE), and $\sigma^{54}$ (RpoN) to compare with our original housekeeping $\sigma^{70}$ (RpoD) promoter (Figure 3B)\textsuperscript{18-21}. CRISPRa was able to activate reporter gene expression when we targeted sites between -60 and -100 upstream of $\sigma^{38}$, $\sigma^{32}$, and $\sigma^{24}$-dependent promoters; these $\sigma$ factors are all members of the $\sigma^{70}$ family. CRISPRa was not active on the $\sigma^{54}$ promoter, possibly because $\sigma^{54}$ initiates gene expression using a distinct mechanism that requires additional cis-regulatory elements\textsuperscript{16}. Activation with promoters regulated by alternative sigma factors was significantly lower than the activation we obtained with a $\sigma^{70}$-dependent promoter. Therefore, while activity will depend on the specific characteristics of the target promoter, in principle CRISPRa can activate promoters regulated by non-housekeeping sigma factors such as $\sigma^{38}$, $\sigma^{32}$, and $\sigma^{24}$, and likely other members of the homologous $\sigma^{70}$ family.
A recent paper described an alternative CRISPRa system that is capable of activating $\sigma^{54}$-dependent genes, which is complementary to our system\(^8\).

**CRISPRa is sensitive to intervening sequence composition**

Our initial model for effective CRISPRa target sites considered the distance of the target site from the TSS but ignored the composition of the intervening sequence upstream of the -35 region. To determine if these sequences affect CRISPRa, we constructed a pooled library of synthetic promoters with randomized sequences between the scRNA target site and the -35 region. We observed a broad distribution of gene activation over a 27-fold range, even though each sequence in this library contained the same scRNA target site and the same minimal promoter (Figure 3C). One possible interpretation of this result is that these randomized sequences contain binding sites for endogenous transcriptional regulators; there is evidence that binding sites can emerge with relatively high frequency from random sequences\(^{22}\). These sites could potentially affect the efficacy of CRISPRa by directly blocking access to a scRNA target site, by blocking RNA polymerase binding, or by interfering with the ability of a CRISPRa effector protein to engage with RNA polymerase.

To directly test the hypothesis that a bound transcriptional effector can disrupt CRISPRa, we introduced a binding site for the transcriptional repressor TetR upstream of the -35 region\(^{23}\). The presence of a bound TetR significantly disrupted CRISPRa-mediated gene activation. Further, adding anhydrotetracycline (aTc), which releases TetR from the DNA, restored CRISPRa activity to the levels observed when TetR was not present (Figure 3D). Taken together, these experiments suggest that the composition of the intervening sequence between the CRISPR-Cas complex and the minimal promoter has a significant effect on the efficacy of CRISPRa, possibly due to the presence of protein binding sites. Because endogenous genes contain binding sites for a variety of transcriptional activators and repressors upstream of the minimal promoter\(^{24,25}\),
this effect could be contributing to the inconsistent and variable effects we observed when targeting endogenous genes for CRISPRa (Figure 2).

**CRISPRa is sharply dependent on single base shifts in target site position**

Our original hypothesis that optimal target sites are located -60 to -100 bases upstream of the TSS was based on an experiment with scRNA sites spaced every 10 bases\(^3\). To further test this hypothesis, we targeted the CRISPRa complex to a window from -61 to -113 at single base resolution. We used a reporter gene with 5 scRNA sites located at -61, -71, -81, -91, and -101 relative to the TSS, and we inserted 1-12 bases upstream of the -35 site to generate a set of reporter genes that allowed the CRISPRa complex to target every possible distance in the optimal targeting window. Using this reporter gene set, we found that shifting the target site by 1-3 bases caused significant decreases in activation (Figure 4A). Shifting the target site further by 4-9 bases decreased expression to levels nearly indistinguishable from background. At 10-11 base shifts, corresponding to one full turn of a DNA helix, gene expression increased again. This periodic positional dependence of CRISPRa extended over the entire -60 to -100 window, with the strongest peaks centered at -81 and -91 and smaller peaks centered at -102 and -70. There is no recovery of activity when the site at -101 is shifted to -111, outside of the -60 to -100 window. This sharp periodic relationship suggests that the criteria for effective target sites are quite stringent, and that both distance and relative periodicity to the TSS are critical factors.

Notably, the distance to the TSS is not the sole determining factor for CRISPRa-mediated expression level. Sites that overlap at the same distance, such as the original -81 site and the -71 site shifted by 10, do not give the same gene expression output (Figure 4A). These discrepancies could arise from intrinsic differences in the activity of the 20 base scRNA target sequence (Supplementary Figure 2) or from the effect of different intervening sequence composition between the scRNA target site and the minimal promoter (Figure 3).
Because we demonstrated that sequence composition can have unexpected effects on CRISPRa (Figure 3), we tested whether the periodicity of CRISPRa was similar in different sequence contexts. We obtained a comparable periodic phase dependence when different nucleotide sequences were used to shift the scRNA target site, and when the bases were inserted at a different location in the promoter (Supplementary Figure 3A). Similar results were also obtained when we performed the base shift experiment with a reporter that had a different 5’ upstream sequence (Supplementary Figure 3B) or where the minimal BBa_J23117 promoter was replaced by endogenous aroK promoter (Supplementary Figure 3C). Further, the sharp positioning dependence was observed when targeting the template or non-template strand of the reporter (Supplementary Figure 3D). Finally, one possible confounding effect could arise if the basal expression level of the reporter gene changes when bases are inserted, which can affect the efficacy of CRISPRa (Figure 3A). However, we observed that basal expression from the original reporter and the +5 base shifted reporter were indistinguishable (Supplementary Figure 3E). Together, these experiments confirm that bacterial CRISPRa is sensitive to periodicity in multiple different sequence contexts.

In the experiments described above, comparisons between single base shifted scRNA sites were performed with different reporter gene constructs, each with a differing number of inserted bases. To test the positional dependence of CRISPRa at single base resolution in a single reporter construct, we designed an alternative reporter gene with 6 adjacent scRNA target sites between -81 and -86. We again observed sharp drops in gene expression when targeting sites one or more bases away from the optimal site at -81 (Supplementary Figure 3F).

The finding that CRISPRa displays the same ~10 base periodicity as the DNA helix suggests that the angular phase of the CRISPRa complex relative to the minimal promoter is critical for effective activation. Our bacterial CRISPRa system requires a direct interaction between the SoxS activation domain and RNA polymerase\(^3\), and this interaction appears to be highly sensitive to both the distance and relative phase of the target site to the minimal promoter.
Effective target sites must be located not only at the proper distance, but also at one of the narrow peaks of activation within the optimal distance range. These stringent requirements suggest that targeting endogenous genes will be extremely challenging. There is ~1 PAM site every 10 bases in the regions upstream of endogenous promoters in *E. coli* (Supplementary Figure 4), and the likelihood that a PAM site will be located at the appropriate phase within a 10 base window is low (Supplementary Figure 4C).

Modifying the CRISPRa complex structure does not expand the range of effective target sites

If rotating the CRISPRa complex out of phase along the DNA prevents SoxS from interacting with RNA polymerase, then a longer amino acid linker to SoxS might allow effective CRISPRa at more scRNA sites. To test this possibility, we extended the linker between MCP and SoxS from 5 amino acids (aa) to 10 or 20 aa, but even with these longer linkers we observed the same sharp dependence on the target site position as with the original 5 aa linker (Figure 4B). We obtained similar results using a linker with a different amino acid composition (Supplementary Figure 5A).

Another potential approach to expand the range of effective CRISPRa sites would be to change the spatial position of the MCP-SoxS protein by altering the position of the MS2 hairpin that binds MCP. We therefore tested multiple alternative scRNA designs that present the MS2 hairpin at different locations. Extending the MS2 stem by 2, 5, 10, and 20 bp resulted in progressively lower CRISPRa activity, but no change in the position of the target sites that were most effective (Supplementary Figure 5B). Similarly, no changes were observed with alternative scRNA designs with one or two MS2 hairpins presented from different locations within the scRNA structure (Supplementary Figure 5C).

Finally, we assessed whether any alternative activation domains could produce a different phase dependent behavior. Previously, these constructs all produced weaker activation than SoxS$^3$, perhaps because they have each distinct optimal target site positions. We tested MCP
fused to TetD, αNTD, lambda cII, and RpoZβ, and dCas9 fused to RpoZβ; however, none of these constructs produced gene activation at any site that was not already effective with SoxS (Supplementary Figure 6).

The sharp phase dependence of CRISPRa may be a general feature of transcriptional regulation in *E. coli*. The native SoxS protein and other transcription factors such as CAP and LacI have restrictive positioning requirements that correspond to DNA periodicity27–29; we confirmed this result with an endogenous SoxS reporter (Supplementary Figure 7). It remains surprising that no structural modifications of the CRISPRa complex produced any changes in the phase dependence. If SoxS is simply tethered to the CRISPRa complex by a flexible linker, we would have expected the peak of effective CRISPRa sites to broaden with longer linkers. The failure of this prediction suggests that our understanding of the CRISPR-Cas complex and its interactions with bacterial transcriptional machinery is fundamentally incomplete, or that the linker tethering SoxS to the CRISPRa complex is not truly flexible. Practically, it means that we still lack a way to expand the range of effective CRISPRa target sites.

An expanded PAM dCas9 variant expands the scope of targetable CRISPRa sites

Because there is a limited number of genes with an appropriate NGG PAM site at precisely the optimal position upstream of the promoter (Supplementary Figure 4C), we attempted to expand the scope of targetable PAM sites for CRISPRa. We used a recently characterized dCas9 variant, dxCas9(3.7), that has improved activity at a variety of non-NGG PAM sites including NGN, GAA, GAT, and CAA7. We generated reporter plasmids by replacing AGG PAM sites with alternative PAM sequences and delivered a CRISPRa system with dxCas9(3.7) to target these reporters. dxCas9(3.7) maintained the ability to target the AGG PAM and showed significantly increased levels of activation at alternative PAM sites compared to dCas9 (Figure 5A). Activation levels varied with different PAM sites and correlated well with dxCas9(3.7) activity previously reported in human cells (Supplementary Figure 8A)7. dxCas9(3.7) showed similar distance and
phase dependent target site preferences as dCas9 (Supplementary Figure 8B & C), but its expanded PAM scope makes it more likely that an arbitrary gene will have a targetable PAM site at an effective position. Bioinformatic analysis of the sequences between transcriptional units in *E. coli* revealed that there are on average 6.4 times more dxCas9(3.7)-compatible PAM sites than NGG PAM sites (Supplementary Figure 8D). Accounting for the fact that dCas9 has some activity at non-NGG sites⁷ (Figure 5A), there are still on average ~2.2-fold more dxCas9(3.7)-compatible PAM sites than dCas9-compatible PAM sites (Supplementary Figure 8D).

To demonstrate the utility of dxCas9(3.7) for CRISPRa at sites inaccessible to dCas9, we constructed a reporter plasmid that contains an AGG PAM site at the original position with maximum CRISPRa activity and an AGT PAM 5 bases downstream. Using this reporter, we observe that both dCas9 and dxCas9(3.7) are effective for CRISPRa at the optimally-positioned NGG PAM site, but neither is capable of activating the AGT PAM site, which is 5 bases out of phase from the optimal site (Figure 5B). We then inserted 5 bases into the reporter to shift the AGT PAM site into the peak activation range. With this reporter, neither dCas9 nor dxCas9(3.7) can activate the NGG PAM site, which is now out of phase. dxCas9(3.7) was now able to effectively activate the AGT PAM site, and dCas9 was ineffective at this site (Figure 5B). This result confirms that dxCas9(3.7) is able to activate optimally-positioned target sites that are inaccessible to dCas9. We expect that this behavior will be effective at many σ⁷₀-family promoters (Figure 3B), and a recent report demonstrated a similar behavior of dxCas9(3.7) at σ⁵₄-dependent promoters⁸.

**Defined rules for effective CRISPRa enable endogenous gene activation**

Our systematic characterization of the requirements for effective CRISPRa in *E. coli* demonstrates that candidate genes must have a targetable PAM site located at one of the sharp peaks of activity upstream of the TSS. In hindsight, the scRNA sites at endogenous genes that we initially targeted in Figure 2 did not meet this criterion. To determine if the revised rules would
enable activation of endogenous \textit{E. coli} genes, we surveyed the genome for candidate genes with appropriately positioned, dxCas9(3.7)-compatible PAM sites (Supplementary Methods) (Supplementary Figure 4C). We selected candidates with multiple potentially effective PAM sites and further narrowed the pool based on two additional criteria: (1) genes should not be too highly expressed (Figure 3A) and (2) genes should be regulated by σ\textsubscript{70}, which was the sigma factor that produced the strongest activation (Figure 3B). Ideally, we would also exclude genes with tightly bound transcriptional regulators in the promoter region (Figure 3D), but this information is not readily available. We chose six genes that could be tested using fluorescent reporter strains from the \textit{E. coli} promoter collection\textsuperscript{30} and targeted two PAM sites for each gene.

We first examined the \textit{yajG} gene, which had two plausible target sites, one of which was only compatible with dxCas9(3.7). We also included an additional site predicted to be out of phase and ineffective for CRISPRa. We observed significant, ~4-6-fold gene activation for the two sites located at the predicted peak of activity at -80/-81, and no activation at the out of phase site at -87 (Figure 6A). The site at -81 is inaccessible to dCas9, and we only observed activation with dxCas9(3.7). We proceeded to test an additional five genes with partial success. We observed significant activation at three genes, although the effects were closer to ~2-fold (Figure 6B). We validated these results by performing RT-qPCR on the endogenous \textit{yajG} and \textit{poxB} loci; the better performing site for each gene resulted in increases in RNA levels (Supplementary Figure 9). For the remaining two candidate genes, there was no detectable activation. Similarly, one of the \textit{ldhA} sites that we targeted in initial experiments (Supplementary Figure 1) was at a predicted optimal site at -91 and failed to give substantial activation.

Although any success at endogenous gene activation is encouraging, significant challenges remain for predictable CRISPRa in bacteria. Our results suggest that even with a precise distance metric for effective target sites, some genes will not be predictably activated. There are several possible explanations: (1) tightly bound negative regulators could interfere with CRISPRa (Figure 3D), and (2) small errors in transcription start site annotation could lead to
inaccurate predictions for effective sites, given that 1-2 base shifts can have dramatic effects on CRISPRa (Figure 4), and (3) intrinsic differences in the activity of the 20 base scRNA target sequence (Supplementary Figure 2).

Discussion

Bacterial CRISPRa is sensitive to a number of factors, including (i) the strength of the target promoter, (ii) the sigma factor regulating the promoter, (iii) the sequence composition immediately upstream of the minimal promoter, (iv) the composition of the scRNA target sequence, (v) the position of the scRNA target site with respect to the TSS at single base resolution. Some of these factors, such as promoter strength and scRNA target sequence composition, are also relevant in eukaryotic systems\textsuperscript{13,15,31,32}. Other factors are plausible given our understanding of bacterial transcription. Sigma factor levels are regulated to control gene expression in response to cell state and external signals\textsuperscript{16}, so it is reasonable that we observed variable levels of activation from promoters with alternative sigma factors. Many bacterial genes are controlled by negative regulators\textsuperscript{33}, and different sequences upstream of the minimal promoter could be recruiting repressors.

The most unexpected property that we observed with bacterial CRISPRa was its sharp, periodic dependence on target site position. This behavior is quite distinct from CRISPRa in eukaryotes, where a broad range of sites upstream of the TSS are effective\textsuperscript{34}, possibly because eukaryotic activators typically recruit transcription factors and chromatin modifying machinery rather than directly recruiting RNA polymerase. There is precedent for bacterial transcriptional activators that are sensitive to target site periodicity\textsuperscript{27–29}, but the dramatic changes in activity with only single base shifts is surprising. Moreover, it is puzzling that we were unable to predictably alter or broaden the range of sites that are effective. Our models for how activators interact with bacterial transcription machinery may be incomplete. It will likely be productive to continue screening for activity at out-of-phase target sites using additional systematic modifications to the
CRISPRa complex structure, alternative CRISPR-Cas systems, and additional candidate transcriptional activation domains.

Despite the challenges described above for identifying effective CRISPRa sites in *E. coli*, our systematic characterization provides a framework for immediate practical applications and a path for future improvements. We now have a clear understanding of the criteria needed to design synthetic promoters that can be regulated by CRISPRa, which will enable the construction of complex, tunable synthetic multi-gene circuits. To extend the scope of CRISPRa to endogenous target genes, expanded PAM variants like dxCas9(3.7)\(^7\), or orthologous dCas9 proteins with alternate PAM specificities\(^{35,36}\) will open more DNA sites for targeting, increasing the likelihood of finding a targetable site at an optimal position relative to the TSS. These strategies lay the groundwork for more widespread use of bacterial CRISPRa in basic research and practical applications including functional genomics screens, metabolic engineering, and synthetic microbial communities.
Methods

Bacterial Strain Construction and Manipulation

Plasmids were cloned using standard molecular biology protocols. Bacterial strains with sfGFP or mRFP1 reporter strains are described in Supplementary Table 1. The CRISPRa system used for each figure panel is described in Supplementary Table 2. Guide RNA target sequences are described in Supplementary Table 3. Plasmid containing the reporter genes and the CRISPR components are described in Supplementary Table 4. S. pyogenes dCas9 (Sp-dCas9) or dxCas9(3.7) were expressed from the endogenous Sp.pCas9 promoter in a p15A vector. MCP-SoxS containing wild-type and mutant SoxS were expressed using the BBa_J23107 promoter (http://parts.igem.org) in the same plasmid with dCas9. The scRNAs were expressed using the BBa_J23119 promoter, either in the same plasmid with the dCas9 protein and the activation domain or in a separate CoIE1 plasmid. The scRNA.b1 or scRNA.b2 designs, where the endogenous tracr terminator hairpin upstream of MS2 was removed3, were used in all experiments except otherwise noted. The zwfp-lacZ and fumCp-lacZ reporter plasmids were generated in a previous study3. mRFP1 and sfGFP reporters were expressed from the weak BBa_J23117 minimal promoter (http://parts.igem.org) in a low-copy pSC101** vector. Variant versions of reporter genes are described in the Supplementary Methods. Plasmid libraries containing N26 sequences between the scRNA target site and BBa_J23117 minimal promoter were constructed by PCR amplification using mixed bases oligos (IDT). The dxCas9(3.7)-VPR plasmid was a gift from David Liu (Addgene #108383)7.

Flow Cytometry

Single colonies from LB plates were inoculated in 500 μL EZ-RDM (Teknova) supplemented with appropriate antibiotics and grown in 96-deep-well plates at 37 °C and shaking. Cultures were grown overnight at 37 °C and shaking and then diluted in 1:50 in DPBS and analyzed on a
MACSQuant VYB flow cytometer (Miltenyi Biotec) using a previously described strategy to gate for single cells\textsuperscript{3}. A side scatter threshold trigger (SSC-H) was applied to enrich for single cells. A narrow gate along the diagonal line on the SSC-H vs SSC-A plot was selected to exclude the events where multiple cells were grouped together. Within the selected population, events that appeared on the edges of the FSC-A vs. SSC-A plot and the fluorescence histogram were excluded.

**Plate Reader Experiments**

Single colonies from LB plates were inoculated in 500 μL EZ-RDM (Teknova) supplemented with appropriate antibiotics and grown in 96-deep-well plates at 37 °C and shaking overnight. For experiments with the *E. coli* promoter collection\textsuperscript{30} the activation domain was placed under the control of a tet-inducible promoter. Attempts to use constitutive CRISPRa were unsuccessful due to plasmid instability, possibly because of toxicity arising from increased expression of the target genes. Single colonies from LB plates were inoculated in 500 μL EZ-RDM supplemented with appropriate antibiotics and 400 nM anhydrotetracycline (aTc) and grown in 96-deep-well plates at 37 °C and shaking overnight. 150 μL of the overnight culture were transferred into a flat, clear-bottomed black 96-well plate and the OD\textsubscript{600} and fluorescence were measured in a Biotek Synergy HTX plate reader. For mRFP1 detection, the excitation wavelength was 540 nm and emission wavelength was 600 nm. For sfGFP detection, the excitation wavelength was 485 nm and emission wavelength was 528 nm.

**Quantitative RT-PCR**

Single colonies from LB plates were inoculated in 5 mL LB containing appropriate antibiotics and grown overnight at 37 °C and shaking. Overnight cultures were diluted 1:100 into 5 mL EZ-RDM supplemented with appropriate antibiotics and grown at 37 °C and shaking until an OD\textsubscript{600} of 0.5 (using 150 μL of culture in a 96 well plate) was reached. For the experiments targeting *yajG* and
poxB, the activation domain was placed under the control of a tet-inducible promoter (see above) and cultures in EZ-RDM were supplemented with 400 nM aTc. Cultures were pelleted and total RNA was extracted using the Aurum Total RNA Mini Kit (Bio-rad). Reverse transcription reactions were performed from 1 μg RNA in 20 μL reactions using iScript reverse transcriptase (Bio-Rad). qPCR reactions were prepared in triplicate in a final volume of 10 μL using SsoAdvanced Universal SYBR Green Supermix (Bio-Rad), 0.5-5 ng of cDNA and 400 nM primers. The reaction was performed in a CFX Connect (Bio-Rad) with a 58 °C annealing temperature and 30 s extension time. A list of the qPCR primer sequences is provided in Supplementary Table 5. Expression levels for each gene were calculated by normalizing to the 16S rRNA gene and relative to a negative control carrying an off target-scRNA using the ΔΔCT method.37

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Author Contributions

J.F., C.D., C.K., B.I.T., J.M.C., and J.G.Z. designed experiments and analyzed data. J.F., C.D., C.K., V.P.C., and B.I.T. performed experiments. J.F., C.D., J.M.C. and J.G.Z. wrote the manuscript.
Figures

Figure 1: A SoxS double mutant does not activate endogenous SoxS targets

A) Reporter system for measuring the CRISPRa activity and endogenous SoxS-dependent gene expression of wild-type or mutant SoxS constructs. CRISPRa activity was determined in a strain harboring a genomically-integrated sfGFP reporter (CD06, Supplementary Table 1). The endogenous SoxS-dependent gene expression was determined by monitoring lacZ expression from reporter plasmids where lacZ was driven by SoxS-regulated promoters zwfp and fumCp. GFP fluorescence was measured by flow cytometry and lacZ activity was measured using a β-galactosidase assay. B) SoxS(R93A/S101A) maintains CRISPRa activity and does not activate expression from the endogenous expression from the zwfp and fumCp reporters. Fluorescence and lacZ activity values were baseline-subtracted using a strain that does not express a scRNA. Both GFP levels and lacZ activities were normalized to the values observed in the strain with wild-type SoxS. Dots indicate the average values between three biological replicates. Error bars indicate the standard deviation.
Figure 2: A simple distance metric does not predict CRISPRa activity.

**A** CRISPRa on the *aroK*-aroB operon. Two scRNA target sites within the 40 base window where CRISPRa is effective (-100 to -60) in heterologous reporter genes (A1-A2) and two sites further upstream (A3-A4) were chosen for the *aroKp1* promoter. **B** CRISPRa on the *cysK* gene. Three scRNA target sites within the 40 base window where CRISPRa is effective in heterologous reporter genes (C1-C3) and two sites further downstream (C4-C5) were chosen for the *cysKp2* promoter. The C4 and C5 sites resulted in repression; targeting these sites close to the core promoter may interfere with RNA polymerase binding. Gene expression was measured using RT-qPCR. Fold activation represents expression levels relative to a strain expressing an off-target scRNA (hAAVS1). Black dots indicate the values of individual biological replicates. Error bars indicate the standard error of the mean between three biological replicates.
Figure 3: CRISPRa is sensitive to promoter identity and local sequence

A

B

C

D

Promoter | Sequence | α factor
---|---|---
sodCp | 5’CCGGTCTGATGATTGTTTCACTTATCTGCTGG-3’ | α^38\(^{(RpoS)}\)
glnAp2 | 5’GACACGAGGCGAAGGCAGATGATTTT-3’ | α^33\(^{(RpoN)}\)
rdgBp | 5’TTGAAAAGGCGAAGCAGAAGCC-3’ | α^32\(^{(RpoH)}\)
yeEp | 5’GAATACTGACGTTTATGCTGACCTCA-3’ | α^37\(^{(RpoK)}\)
J32117 | 5’TTCGGAGCTGCGTCCCCATTCCGTCGGATGAC-3’ | α^30\(^{(RpoD)}\)

Baseline promoter activity

RFP fluorescence/OD (a.u.)

Baseline expression

Fold activation

Distance to TSS (bp)

Fold activation

RFP fluorescence/OD (a.u.)

Single Colonies from Library
**Figure 3: CRISPRa is sensitive to promoter identity and local sequence.**

**A)** CRISPRa is sensitive to promoter strength. Reporter genes were constructed by replacing Bba_J23117 from J3-J23117-mRFP1 reporter (Supplementary Methods) with other minimal promoters from the Anderson series (J231NN). J3-J231NN promoters contain a scRNA target site at the position where CRISPRa is most effective (-81 from the TSS on the non-template strand)\(^3\). The plot at the bottom left shows the Fluorescence/OD\(_{600}\) values of strains expressing an on-target or off-target scRNA. The plot on the right shows the fold activation measured at each promoter against their baseline expression in the presence of an off-target scRNA (J206).

**B)** CRISPRa can activate promoters regulated by \(\sigma^{38}\)(RpoS), \(\sigma^{32}\)(RpoH), and \(\sigma^{24}\)(RpoE) sigma factors when targeting sites positioned -60 to -100 bases upstream of the TSS. The J1-J23117-mRFP1 reporter with scRNA sites situated every 10 bp was used to construct the reporter plasmids\(^3\). The minimal promoter from the reporter plasmid was replaced by the sodCp (RpoS), glnAp2 (RpoN), rdgBp (RpoH), or yieEp (RpoE) minimal promoters. The plot on the top right shows the baseline Fluorescence/OD\(_{600}\) of each promoter when expressing the CRISPRa components with off-target scRNA (hAAVS1). The bottom plot shows the fold activation versus the position of the scRNA target.

**C)** CRISPRa activity differs significantly among promoters with varying sequence composition between the scRNA target and the -35 region. A pooled reporter library was constructed by replacing 26 bases of sequence with random bases between the scRNA target site and the -35 region on the J3-J23117-sfGFP reporter. The cloned plasmid library was co-transformed with a plasmid containing the CRISPRa components. Each green bar represents the Fluorescence/OD\(_{600}\) value of an overnight culture from a single colony of the library. The blue bar represents the Fluorescence/OD\(_{600}\) value of a strain expressing the original J3-J23117-sfGFP reporter and the CRISPRa components with a scRNA targeting J306. The grey bar represents a negative control expressing the J3-J23117-sfGFP reporter plasmid and the CRISPRa components with an off-target scRNA (J206).

**D)** CRISPRa was partially inhibited by the presence of a TetR transcriptional repressor binding between the scRNA target and the
minimal promoter. Releasing TetR from its target DNA with 1 μM aTc fully restored CRISPRa activity to its original level. The reporter plasmid was constructed by placing a tet operator (tetO) site upstream of the -35 region in the J3-J23117-mRFP1 reporter (Supplementary Information). Cultures where CRISPRa was targeted to the J306 site or to an off-target site (J206) were grown overnight in media +/- 1 μM aTc. Fluorescence/OD<sub>600</sub> values were measured using a plate reader. Fold activation indicates the Fluorescence/OD<sub>600</sub> of strains relative to an off-target control. Colored dots and bars indicate the average values between three biological replicates. Black dots indicate the values of individual biological replicates. Error bars indicate the standard deviation between biological replicates.
Figure 4: CRISPRa is sensitive to the precise position of the scRNA target.

A) CRISPRa displays periodic positioning dependence with peak activities every 10-11 bases between -60 to -100 from the TSS. Reporter genes were constructed by inserting 0-12 bases upstream of the -35 region of the J1-J23117-mRFP1 reporter. Five scRNA sites (J102, J104, J106, J108, J110) with positions -61, -71, -81, -91, -101 from the TSS on the non-template strand of the original promoter were targeted. In this way, the complete -61 to -113 region can be covered at single base resolution. The color coding indicates data for the same target site shifted across a 12 base window. The panel on the right shows the baseline expression of reporters with shifted bases when an off-target scRNA was used (J206). The grey area represents the range of the
baselines among the reporter series. For comparison, previous CRISPRa data for the J102, J104, J106, J108, J110 target positions at 10 base resolution are shown on the schematic above the plot. B) Extending the linker length between MCP and SoxS does not change the position dependence of CRISPRa. The J1-J23117-mRFP1 reporter plasmid series with base shifts were delivered together with CRISPRa components for targeting J106. The MCP-SoxS(R93A) effector contained 5aa, 10aa, and 20aa linkers. For comparison, previous CRISPRa data with 5aa, 10aa, or 20aa linker between MCP and SoxS targeting at the -81 and -91 positions are shown on the schematic above the plot. Fluorescence/OD$_{600}$ was measured using a plate reader. Dots indicate the average values between three biological replicates. Error bars indicate the standard deviation.
Figure 5: dxCas9(3.7) expands the range of targetable sites by recognizing alternative PAMs.

A) CRISPRa with dxCas9(3.7) displayed activity on non-NGG PAM sites with AGA, AGC, AGT, CGA, CGC, CGT, GGA, GGC, GGT, TGA, TGC, TGT, GAA, GAT, CAA sequences. CRISPRa activity with dxCas9(3.7) on non-NGG PAM sites was generally lower (6-fold to 89-fold activation relative to a control without a scRNA) compared to the AGG PAM site (188-fold activation). Sp-dCas9 also displayed moderate CRISPRa activity at non-NGG PAM sites with AGA, GGA, GGT, TGA sequences, consistent with published reports. Reporter plasmids were
constructed by replacing the AGG PAM site for the J306 target in the J3-J23117-mRFP1 reporter with alternative PAM sequences that have been previously reported to be recognized by dxCas9(3.7) in human cells. The (-) sign indicates a control expressing the original reporter with the AGG PAM and the CRISPRa components with Sp-dCas9, the activation domain and no scRNA. B) dxCas9(3.7) can activate promoters that cannot be activated by Sp-dCas9. When the scRNA target at the optimal position (-81 to the TSS) has an AGG PAM site, both Sp-dCas9 and dxCas9(3.7) increased gene expression by 50-fold. When the scRNA target at the optimal position has an AGT PAM site, only dxCas9(3.7) displayed a 7-fold increase in gene expression while Sp-dCas9 was inactive. The reporter gene has a target with an AGG PAM (M1) and a target with an AGT PAM (M2) upstream of a BBa_J23117 minimal promoter. In reporter gene A, the AGG target was located -81 to the TSS on the non-template strand and the AGT target was located -76 to the TSS on the non-template strand. In reporter gene B, 5 bases were inserted upstream of the -35 region, shifting the locations of the AGG target and AGT target to -86 and -81, respectively. The (-) sign indicates a negative control strain that contains the reporter plasmid and a plasmid expressing Sp-dCas9, the activation domain and an off-target scRNA (J206). Bars in Panel A represent the average of the median fluorescence among three biological replicates. Bars in Panel B represent the average RFP Fluorescence/OD600 among three biological replicates. Black dots represent the individual biological replicates and error bars indicate the standard deviation.
Figure 6: Predictive rules enable endogenous gene activation at some targets

A) CRISPRa using dCas9 and dxCas9(3.7) was targeted to the reporter plasmid containing the yajG promoter from the E. coli promoter collection. Three scRNA target sites were selected; two sites were located at the positions where CRISPRa was most effective (Y1-Y2), and one was located out of phase (Y3). A negative control (OT) expressing an off-target scRNA (J306) was included. B) CRISPRa was targeted to yajG and five additional promoters from the E. coli promoter collection (Supplementary Methods). Two scRNA sites located at the positions where CRISPRa was most effective were targeted for each gene using dxCas9(3.7). Fluorescence/OD$_{600}$ was measured using a plate reader. Fold activation indicates the Fluorescence/OD$_{600}$ of strains relative to an off-target control (J306). Bars represent the average among three biological replicates. Black dots represent individual biological replicates (panel A), and the Fluorescence/OD600 of individual biological replicates divided by the average Fluorescence/OD600 of the off-target control (panel B). Error bars indicate the standard deviation.
References

1. Brophy, J. A. & Voigt, C. A. Principles of genetic circuit design. Nat Methods 11, 508–520 (2014).

2. Nielsen, J. & Keasling, J. D. Engineering Cellular Metabolism. Cell 164, 1185–1197 (2016).

3. Dong, C., Fontana, J., Patel, A., Carothers, J. M. & Zalatan, J. G. Synthetic CRISPR-Cas gene activators for transcriptional reprogramming in bacteria. Nat Commun 9, 2489 (2018).

4. Qi, L. S. et al. Repurposing CRISPR as an RNA-Guided Platform for Sequence-Specific Control of Gene Expression. Cell 152, 1173–1183 (2013).

5. Wang, H., Russa, M. & Qi, L. S. CRISPR/Cas9 in Genome Editing and Beyond. Annu Rev Biochem 85, 1–38 (2015).

6. Keseler, I. M. et al. The EcoCyc database: reflecting new knowledge about Escherichia coli K-12. Nucleic Acids Res 45, D543–D550 (2017).

7. Hu, J. H. et al. Evolved Cas9 variants with broad PAM compatibility and high DNA specificity. Nature 556, 57–63 (2018).

8. Liu, Y., Wan, X. & Wang, B. Engineered CRISPRa enables programmable eukaryote-like gene activation in bacteria. Nat Commun 10, 3693 (2019).

9. Rodriguez, A. et al. Engineering Escherichia coli to overproduce aromatic amino acids and derived compounds. Microb Cell Fact 13, 126 (2014).

10. Byrne, C., Monroe, R., Ward, K. & Kredich, N. DNA sequences of the cysK regions of Salmonella typhimurium and Escherichia coli and linkage of the cysK regions to ptsH. J Bacteriol 170, 3150–3157 (1988).

11. Jiang, G., Nikolova, S. & Clark, D. P. Regulation of the IdhA gene, encoding the fermentative lactate dehydrogenase of Escherichia coli. Microbiology+ 147, 2437–2446 (2001).

12. Silander, O. K. et al. A Genome-Wide Analysis of Promoter-Mediated Phenotypic Noise in Escherichia coli. Plos Genet 8, e1002443 (2012).

13. Chavez, A. et al. Highly efficient Cas9-mediated transcriptional programming. Nat Methods
12, 326–328 (2015).

14. Chavez, A. et al. Comparison of Cas9 activators in multiple species. *Nat Methods* **13**, 563–567 (2016).

15. Konermann, S. et al. Genome-scale transcriptional activation by an engineered CRISPR-Cas9 complex. *Nature* **517**, 583–588 (2015).

16. Gruber, T. M. & Gross, C. A. Multiple Sigma Subunits and the Partitioning of Bacterial Transcription Space. *Annu Rev Microbiol* **57**, 441–466 (2003).

17. Shah, I. M. & Wolf, R. E. Novel Protein–Protein Interaction Between Escherichia coli SoxS and the DNA Binding Determinant of the RNA Polymerase α Subunit: SoxS Functions as a Co-sigma Factor and Redeploys RNA Polymerase from UP-element-containing Promoters to SoxS-dependent Promoters during Oxidative Stress. *J Mol Biol* **343**, 513–532 (2004).

18. Gort, A., Ferber, D. M. & Imlay, J. A. The regulation and role of the periplasmic copper, zinc superoxide dismutase of Escherichia coli. *Mol Microbiol* **32**, 179–191 (1999).

19. Nonaka, G., Blankschien, M., Herman, C., Gross, C. A. & Rhodius, V. A. Regulon and promoter analysis of the E. coli heat-shock factor, σ32, reveals a multifaceted cellular response to heat stress. *Gene Dev* **20**, 1776–1789 (2006).

20. Rhodius, V. A., Suh, W., Nonaka, G., West, J. & Gross, C. A. Conserved and Variable Functions of the σE Stress Response in Related Genomes. *Plos Biol* **4**, e2 (2006).

21. Tian, Z., Li, Q., Buck, M., Kolb, A. & Wang, Y. The CRP–cAMP complex and downregulation of the glnAp2 promoter provides a novel regulatory linkage between carbon metabolism and nitrogen assimilation in Escherichia coli. *Mol Microbiol* **41**, 911–924 (2001).

22. Yona, A. H., Alm, E. J. & Gore, J. Random sequences rapidly evolve into de novo promoters. *Nat Commun* **9**, 1530 (2018).

23. Lee, T. et al. BglBrick vectors and datasheets: A synthetic biology platform for gene expression. *J Biol Eng* **5**, 12 (2011).

24. Mendoza-Vargas, A. et al. Genome-Wide Identification of Transcription Start Sites,
Promoters and Transcription Factor Binding Sites in E. coli. *Plos One* 4, e7526 (2009).

25. Lloyd, G., Landini, P. & Busby, S. Activation and repression of transcription initiation in bacteria. *Essays Biochem* 37, 17–31 (2001).

26. Bikard, D. *et al.* Programmable repression and activation of bacterial gene expression using an engineered CRISPR-Cas system. *Nucleic Acids Res* 41, 7429–7437 (2013).

27. Wood, T. I. *et al.* Interdependence of the position and orientation of SoxS binding sites in the transcriptional activation of the class I subset of Escherichia coli superoxide-inducible promoters. *Mol Microbiol* 34, 414–430 (1999).

28. Zhou, Y., Kolb, A., Busby, S. & Wang, Y.-P. Spacing requirements for Class I transcription activation in bacteria are set by promoter elements. *Nucleic Acids Res* 42, 9209–9216 (2014).

29. Müller, J., Barker, A., Oehler, S. & Müller-Hill, B. Dimeric lac repressors exhibit phase-dependent co-operativity. *J Mol Biol* 284, 851–857 (1998).

30. Zaslaver, A. *et al.* A comprehensive library of fluorescent transcriptional reporters for Escherichia coli. *Nature Methods* 3, 623–628 (2006).

31. Kiani, S. *et al.* CRISPR transcriptional repression devices and layered circuits in mammalian cells. *Nat Methods* 11, 723–726 (2014).

32. Gander, M. W., Vrana, J. D., Voje, W. E., Carothers, J. M. & Klavins, E. Digital logic circuits in yeast with CRISPR-dCas9 NOR gates. *Nat Commun* 8, 15459 (2017).

33. Rojo, F. Repression of transcription initiation in bacteria. *Journal of bacteriology* 181, 2987–91 (1999).

34. Gilbert, L. A. *et al.* Genome-Scale CRISPR-Mediated Control of Gene Repression and Activation. *Cell* 159, 647–661 (2014).

35. Leenay, R. T. & Beisel, C. L. Deciphering, Communicating, and Engineering the CRISPR PAM. *J Mol Biol* 429, 177–191 (2017).

36. Shmakov, S. *et al.* Diversity and evolution of class 2 CRISPR-Cas systems. *Nat Rev Microbiol* 15, 169–182 (2017).
37. Livak, K. J. & Schmittgen, T. D. Analysis of Relative Gene Expression Data Using Real-Time Quantitative PCR and the 2−ΔΔC T Method. *Methods* **25**, 402–408 (2001).

38. Griffith, K. L. & Wolf, R. E. A Comprehensive Alanine Scanning Mutagenesis of the Escherichia coli Transcriptional Activator SoxS: Identifying Amino Acids Important for DNA Binding and Transcription Activation. *J Mol Biol* **322**, 237–257 (2002).