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CRISPR Display: A modular method for locus-specific targeting of long noncoding RNAs and synthetic RNA devices *in vivo*

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Abstract

Noncoding RNAs (ncRNAs) comprise an important class of regulatory molecules that mediate a vast array of biological processes. This broad functional capacity has also facilitated the design of artificial ncRNAs with novel functions. To further investigate and harness these capabilities, we developed CRISPR-Display (“CRISP-Disp”), a targeted localization method that uses *Sp. Cas9* to deploy large RNA cargos to DNA loci. We demonstrate that exogenous RNA domains can be functionally appended onto the CRISPR scaffold at multiple insertion points, allowing the construction of Cas9 complexes with protein-binding cassettes, artificial aptamers, pools of random sequences, and RNAs up to 4.8 kilobases in length, including natural lncRNAs. Unlike most existing CRISPR methods, CRISP-Disp allows simultaneous multiplexing of distinct functions at multiple targets, limited only by the number of available functional RNA motifs. We anticipate that this technology will provide a powerful method with which to ectopically localize functional RNAs and ribonucleoprotein (RNP) complexes at specified genomic loci.

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Author Contributions

D.M.S. designed and performed experiments; F.E.H. designed and performed microscopy experiments (Figs. 4c–d and Supplementary Figs. 13–14); S.T.Y. assisted with computational analysis (Fig. 3d and Supplementary Figs. 5, 10); J.L.R. directed research. D.M.S. and J.L.R. wrote the manuscript.

Competing financial interests

The authors declare competing financial interests.

Introduction

Noncoding RNAs (ncRNAs) are central components of diverse and fundamental processes in all kingdoms of life¹. In eukaryotes, many well-established ncRNAs, and a number of newly identified mammalian long ncRNAs, are thought to help initiate or maintain regulatory processes within the nucleus^{1–3}. However, mechanistic dissection of these putative nuclear regulators *in vivo* is often technically limited. For example, established knock-in and knockout strategies^{4, 5} lack the throughput required for high-resolution structure/function analysis, and cannot easily separate roles performed by an RNA transcript from those performed by a functional DNA element or by a cryptically encoded peptide^{5, 6}.

Therefore, an experimental method that post-transcriptionally relocates a ncRNA transcript to an ectopic site would be an important tool for the study of natural ncRNA function. In addition, this method could provide a powerful engine for synthetic biology. Many RNA domains—both natural and artificial—have been adapted as components in synthetic regulators, reporters and scaffolds^{7–13}. Targeting such RNA devices to specific DNA loci would enable a wide range of novel synthetic biological methods.

Towards these goals, we aimed to develop a general ncRNA ectopic localization system. We reasoned that this could be achieved using an artificial protein “conduit” that is programmed to bind a ncRNA and target it to specific DNA loci. (Supplementary Fig. 1a). A potentially powerful source for such a conduit is the *S. pyogenes* Cas9 nuclease (*Sp. Cas9*), an extremely high-affinity, programmable DNA-binding protein isolated from a type II CRISPR-associated system^{14–16}. In a multitude of CRISPR-based biotechnology applications^{17–22}, the guide that programs Cas9 DNA-targeting is presented in a so-called sgRNA, wherein the two natural Cas9 RNA cofactors (crRNA and tracrRNA)^{14, 15} are fused via an engineered loop (Supplementary Fig. 1b). Yet, despite recent work dissecting the determinants of Cas9 RNA recognition^{23–26}, it remained unclear if and where large, structured RNA domains could be implanted within CRISPR complexes while maintaining RNA-directed localization.

Here, we demonstrate that a nuclease-deficient *Sp. dCas9* mutant, “dCas9” (Refs. 17, 27–29)) can be co-opted to deploy a large RNA cargo to targeted DNA loci by directly linking that cargo to the sgRNA. We term this strategy, in which RNA domains are displayed on dCas9, CRISPR-Display, or “CRISP-Disp.” With the appropriate expression system and insertion point, CRISP-Disp does not appear inherently limited by the size or sequence composition of its RNA cargo. This allows dCas9 complexes to be functionalized with structured RNA domains, natural lncRNAs several kb in length, artificial RNA modules and pools of random sequences. Furthermore, these RNA-based functions can be simultaneously multiplexed using a shared pool of dCas9. This work provides initial insights into the general utility of CRISP-Disp for both the study of natural ncRNAs and the construction of novel RNA-based devices.

Results

Adapting CRISPR-Cas9 as an RNA Display Device

To assess potential ncRNA localization methods *in vivo*, we first implemented a highly sensitive reporter system in HEK293FT cells. In this system, the efficacy of an ectopically targeted regulator (protein or RNP) is reported by luciferase and fluorescent protein expression. RNA-targeting strategies were surveyed using both transiently transfected and stably integrated reporters (Supplementary Fig. 1c–f, Supplementary Table 1)^{27, 30}.

In our initial strategy, the RNA-deploying “conduits” targeted to these reporters were based upon transcription activator-like effectors (TALEs)³⁰. However, we were unable to coax TALEs into recruiting ncRNAs to a chromatin-integrated locus (Supplementary Figs. 2–3). We ascribed this problem to the separable DNA- and RNA-binding activities of our conduit, which could be independently saturated without forming DNA•TALE•RNA ternary complexes (Supplementary Data).

To circumvent this issue, we turned to the *S. pyogenes* CRISPR-(d)Cas9 system, which intrinsically couples its DNA- and RNA-binding activities (Supplementary Fig. 1b–f)^{15, 16}. However, it was unclear *a priori* where insertions within the dCas9-bound RNAs would be tolerated, and how large they could be. To examine this, we devised five model cofactor RNAs (TOP1–4; INT) in which structured, 81–250 nt “accessory domains” were appended to the termini of, or internally within the sgRNA or tracrRNA¹⁵. Each accessory domain carried a cassette of stem-loops recognized by the PP7 phage coat protein³¹ (Fig. 1a; Supplementary Fig. 4a–b). At 357 nt, the largest of these constructs adds three-fold more sequence than does the longest modified sgRNA previously reported^{20, 25, 26}.

We subjected these RNA chimeras to two variations of our CRISPR transcription activator assay (Fig. 1b). In “direct activation” assays, they were coexpressed with dCas9 fused to the VP64 transcription activator (dCas~VP). In this format, reporter gene activation indicates that the sgRNA variant competently binds and targets dCas9. In “bridged activation” assays, constructs were coexpressed with dCas9 and PP7~VP, a chimera of PP7 and VP64. Bridged activation should only occur if the accessory domain remains intact in the mature dCas9 complexes.

Using transient reporters, we observed direct activation with all five topologies (Fig. 1c, *top*). However, while the activities of TOP1, TOP2 and INT were reduced less than twofold from that of the minimal sgRNA, TOP3 and TOP4 were ~9–30-fold less proficient. Critically, bridged activation was only observed with TOP1, TOP3 and INT, indicating that these constructs alone retained functional accessory domains in mature dCas9 complexes. These results were corroborated by FACS (Supplementary Fig. 4e) and qualitatively paralleled using integrated reporters (Fig. 1c, *bottom*). We attributed the lack of TOP2 bridged activation to partial degradation of that construct’s accessory domain, as indicated using RNA Immunoprecipitation (RIP) and qRT-PCR (Supplementary Fig. 4f)^{20, 25, 32}.

Targeting long RNAs to endogenous genomic loci

To illustrate the general applicability of CRISP-Disp, we tested whether results from our reporter system could be recapitulated at endogenous loci. We generated pools of sgRNA, TOP1 and INT constructs targeting the human *ASCL1*, *IL1RN*, *NTF3* and *TTN* promoters^{28, 29, 33} (Supplementary Table 2), and surveyed direct and bridged activation of these genes by qRT-PCR. As predicted, activation of each locus paralleled the results obtained using our GLuc reporter, demonstrating CRISP-Disp enables deployment of large RNA domains to genomic loci (Fig. 1d).

To confirm that dCas9 targeting fidelity is not perturbed by the addition of the accessory RNA domains, we performed mRNA seq from reporter cells expressing dCas9~VP and GLuc-targeting sgRNA, TOP1 and INT constructs. As predicted, changes in global gene expression induced by each RNA construct—a proxy for dCas9 off targeting—were essentially indistinguishable, indicating that TOP1- and INT-like accessory domains do not significantly alter dCas9 fidelity (Supplementary Fig. 5).

CRISP-Disp with “artificial lncRNA” scaffolds

We next sought to engineer CRISP-Disp for use with even longer ncRNAs, which required replacing the conventional RNA Polymerase III (Pol III) promoter used above with an RNA Polymerase II (Pol II) promoter and terminator. We therefore surveyed a variety of standard and noncanonical Pol II expression systems for the ability to generate nuclear-localized CRISP-Disp RNAs *de novo* (Supplementary Fig. 6–7; Supplementary Discussion). Although such Pol II transcripts were generally less effective than their Pol III-driven counterparts¹⁹, expression from several different backbones partially restored their activity (Supplementary Fig. 6b). Surprisingly, one such backbone (“CMV/3’Box”) also enabled bridged activation with TOP2, which was unviable under Pol III expression (Supplementary Fig. 6c–e).

Next, we next attempted to build dCas9 complexes with model RNAs approaching the size of natural lncRNAs. We expanded our CMV/3’-Box TOP1 and TOP2 constructs with a second P4–P6 domain, bearing stem-loops recognized by the MS2 phage coat protein³¹, to generate a series of “artificial lncRNA” scaffolds with ~650 nt accessory domains (“Double TOP0–1” Fig. 2a, and Supplementary Fig. 4a–d). Although all three of these constructs induced measurable direct activation, Double TOP1–2 were more proficient, with activities nearly rivaling those of their single-domain counterparts (Fig. 2b). Moreover, all three constructs exhibited significant bridged activation. In transient reporter assays, activity monotonically increased upon coexpression with PP7~VP, MS2~VP and both activator proteins, indicating that each RNA construct retained both P4–P6 domains in mature dCas9 complexes (Fig. 2b). A qualitatively similar trend was observed with Double TOP1 using integrated reporters, despite the assay’s limited dynamic range (Supplementary Fig. 8). We furthermore confirmed the integrity of the Double TOP1–2 accessory domains by dCas9 RIP: for each construct, we obtained essentially stoichiometric yields of the sgRNA core and double P4–P6 accessory domains (Fig. 2c).

CRISP-Disp with natural lncRNA domains

A potentially powerful application of CRISP-Disp would be the ectopic localization of natural lncRNAs post-transcriptionally, since reconstitution of lncRNA activity at an ectopic site is unattainable by existing methods^{5, 6}. To demonstrate the plausibility of this approach, we first established that natural long ncRNAs could be incorporated into CRISP-Disp complexes. We generated Pol II-driven TOP1- and INT-like constructs appended with human lncRNA domains: the repressive NoRC-binding pRNA stem-loop³⁴ and *Xist* A-repeat (“RepA”) domains⁴, three enhancer-transcribed RNAs (eRNAs, Ref.³⁵) and the 4799 nt transcription activator *HOTTIP*³⁶ (Supplementary Table 4).

Each construct induced significant direct activation, indicating that each formed functional dCas9-targeting complexes (Fig. 2d). This was further supported by RIP-qPCR: although overall dCas9 complexation appeared to decline monotonically with increasing sgRNA~lncRNA length, in all cases nearly quantitative yields of intact lncRNA domains were recovered, relative to the corresponding sgRNA core (Fig. 2e). We propose that the observed minor variation in construct integrity may be contingent on several factors, including length and RNA structure²⁵.

We next tested if the lncRNA components of CRISP-Disp complexes could themselves regulate our reporter. Encouragingly, most of the lncRNA constructs repressed or activated GLuc expression as would be predicted from existing studies^{4, 34–36}: pRNA and RepA diminished GLuc expression, while two eRNAs (*TRERNA1*, ncRNA-a3) and *HOTTIP* induced moderate activation (Fig. 2f). Although these effects were quite modest, they may have been limited by technical aspects of our assay, or by absence of particular factors in HEK293T cells. Regardless, these mild initial results demonstrate the plausibility of larger-scale lncRNA functional studies with CRISP-Disp.

CRISP-Disp with a diverse array of RNA species

We were particularly intrigued by the proficiency of U6-driven INT (Fig. 1a,c), and reasoned that the sgRNA “engineered loop” might provide a universal insertion point for exogenous RNA domains. To explore this possibility, we first examined the influence of internal insert size. Encouragingly, INT-like constructs bearing 25–137 nt cassettes of PP7 stem-loops each induced robust activation in all assay formats (Fig. 3a), as did a construct bearing a ~250 nt domain equivalent to those of TOP1–4 (Fig. 3b; Supplementary Table 5). Hence, dCas9 can easily accommodate even large, structurally discontinuous inserts within the sgRNA core.

To examine the influence of internal insert sequence, we synthesized a pool of $\sim 1.2 \times 10^6$ unique sgRNA variants displaying internal cassettes of 25 random nucleotides (Supplementary Fig. 9). In aggregate, the activity of this INT-N₂₅Pool rivaled that of the minimal sgRNA (Fig. 3c), implying that many of the variants formed productive CRISP-Disp complexes. To confirm this, we immunoprecipitated dCas9•INT-N₂₅Pool complexes and analyzed the copurified sgRNA sequences by deep sequencing (RIP-Seq, Fig. 3d, Supplementary Fig. 10). Fewer than 0.01% and 0.02% of the sequence variants were significantly enriched or de-enriched, respectively; motif analysis of these variants revealed no apparent sequence constraints influencing sgRNA•dCas9 complexation. Although this

pool represents a small ($\sim 1.1 \times 10^{-9}$), sampling of the total 25mer sequence space, we extrapolate that CRISP-Disp is not intrinsically limited by the sequence of an internal insert.

To explore the potential utility of INT-like inserts, we next generated a series of INT-like constructs displaying an array of functional RNA domains, comprising natural protein-binding motifs^{31, 37, 38} and artificial aptamers that bind proteins^{7, 8} and small molecules¹³ (Supplementary Fig. 11 and Supplementary Table 5). Satisfyingly, all constructs were viable CRISP-Disp substrates (Fig. 3e). Variation in construct efficacy did not appear to result from limiting RNA expression levels (Supplementary Fig. 12), implying that design of a high-efficiency CRISP-Disp construct for a given motif may require some structural or sequence optimization.

Next, we used INT-like constructs to deploy structured RNA domains to endogenous loci. Constructs bearing the S1 streptavidin aptamer, a potentially useful RNA device⁷, and P4-P6[3xPP7], the largest INT-like construct tested, each activated *ASCL1*, *IL1RN*, *NTF3*, and *TTN*^{28, 29, 33} as predicted (Fig. 3f). Furthermore, INT-P4-P6[3xPP7] did not appear to perturb dCas9 targeting fidelity (Supplementary Fig. 5). We therefore infer that INT-like constructs can display RNA large RNA domains to specific loci genome-wide.

CRISP-Disp enables concomitant deployment of multiple functionalities

One potential advantage of CRISP-Disp is that it enables disparate functions to be simultaneously performed at diverse loci in the same cell, using a single toolset. This modularity could be achieved using a set of orthogonal RNA-binding proteins: each protein would be fused to a unique functional domain and targeted by sgRNAs bearing its cognate RNA motif²⁵. To demonstrate the plausibility of this scheme, we first established a preliminary set of such RNA-binding proteins (*A. fulgidus* L7Ae, MS2 and PP7)^{31, 37} for display on dCas9. As predicted, when fused to VP64, each protein was only competent to induce bridged reporter activation when coexpressed with an sgRNA bearing its cognate RNA motif (Fig. 4a).

As a first demonstration of CRISP-Disp modularity, we bound dCas9 to multiple genomic targets, but activated only one (Fig. 4b). We coexpressed complementary pairs of GLuc- and *NTF3*-targeting INT-like sgRNA variants bearing PP7 and MS2 stem-loops (GLuc[3xPP7] with *NTF3*[3xMS2], or GLuc[3xMS2] with *NTF3*[3xPP7]) in integrated reporter cells. In the presence of dCas9~VP, we observed robust activation of both target genes, regardless of the sgRNA pair used (Fig. 4b, *left*), indicating that dCas9 had bound both loci under all conditions. However, in the presence of dCas9 and PP7~VP, only the gene targeted by sgRNAs bearing PP7 stem-loops was activated; the converse was observed upon coexpression of dCas9 and MS2~VP (Fig. 4b, *middle and right*). Hence, CRISP-Disp enables modular control of gene expression, as has been recently demonstrated with TOP1-like constructs²⁵.

As a second demonstration of CRISP-Disp modularity, we sought to simultaneously activate one locus and image another (Fig. 4c). To accomplish this, we implemented a “bridged CRISPR-imaging” approach, in which targeted loci (here, telomeres¹⁸) are visualized using a ternary complex comprising dCas9, MS2~mCherry, and INT-like [3xMS2] constructs

(Supplementary Fig. 13a). Using bridged telomere imaging in HEK293FT cells, we observed numerous (8–55; average of 26.6, in ~97 mCherry+ cells) fluorescent nuclear foci, which required expression of dCas9 and the proper cognate sgRNA (Supplementary Fig. 13b–c). Next, to simultaneously activate one locus and image another, we employed our integrated reporter cells, performing bridged activation (targeted by INT) at the GLuc reporter, and bridged imaging at telomeres (Fig. 4c, *top*). As predicted, each function could be individually or concomitantly controlled by expression with PP7~VP64, MS2~mCherry or both (Fig. 4c, *bottom*).

CRISP-Disp of autonomous RNA domains

Another potential advantage of CRISP-Disp is that it allows autonomously functional RNA domains, such as ribozymes, aptamers and regulatory devices^{7–13}, to be targeted to individual loci. As a preliminary illustration of this approach, we targeted “Spinach2,” an artificial aptamer that binds to and induces fluorescence in a cell-permeable dye¹³, to telomeres¹⁸. When we coexpressed a Spinach2-appended telomere-targeting sgRNA with dCas9 and treated cells with the Spinach2 ligand DFHBI-1T (Ref. 13), we observed numerous (10–20; average of 12, in ~20% of cells) nuclear fluorescent foci (Fig. 4d and Supplementary Figure 14). Critically, no fluorescent foci were observed in the absence of dCas9, or when Spinach2 was targeted to the Gluc reporter. Although the Spinach2 signal was less robust than that observed using conventional CRISPR-imaging¹⁸, or with our own bridged imaging approach, this experiment represent an important proof-of-principle, demonstrating that artificial RNA domains can be harnessed to imbue dCas9 with novel properties.

Discussion

In the short time since its initial characterization^{14, 15}, *Sp.* CRISPR-Cas9 has already been coopted for a host of exquisitely powerful genome modification and regulatory technologies^{17–22}. We envision that CRISPR-Display, which limits the function dCas9 to DNA-targeting and “outsources” all other roles to RNA-based domains, will provide the basis for an even wider array of methods (Fig. 4e).

CRISPR-Display’s ability to ectopically localize diverse RNA cargos to targeted DNA loci has several implications. First, CRISP-Disp allows implementation of cargo-specific functions at each of its target sites, using a common toolkit. In theory, the breadth of distinct functions accessible in a single CRISP-Disp experiment is limited only by the number of orthogonal RNA domains or RNA/binding-protein pairs available (as in Fig. 4a–c). For example one might simultaneously activate and repress transcription, epigenetically mark histones and DNA, induce double-stranded breaks and image discrete sets of target loci. Our preliminary experiments, which targeted discrete—though unrelated—genomic loci for binding, activation and imaging (Fig. 4a–c) provide a small glimpse into the potential power this approach holds.

Second, since CRISP-Disp is not intrinsically limited by RNA length (Figs. 2, 3a–b), it may provide a method for the locus-targeted reconstitution of natural regulatory RNAs, which could dramatically advance the study of lncRNA mechanism^{1–6}. For example, CRISP-DISP

could be used to bring a long noncoding RNA domain to a target locus, and assess if the RNA molecule alone is functional when decoupled from the act of its transcription. The preliminary data presented here (Fig. 2d–f) demonstrate the plausibility of such experiments. However, more thorough, *bona fide* lncRNA reconstitution will likely require consideration of many variables (*e.g.*, the chromatin state at the targeted locus), and may depend on the particular lncRNA under consideration.

Third, CRISP-Disp expands the scope of Cas9-based methods by making available the broad functional repertoire of artificial RNA devices. While our initial illustration of this principal focused on imaging (Fig. 4d), other applications built from aptamers, ribozymes, sensors, processors and scaffolds^{7–13} are possible. For example, CRISP-Disp with RNA scaffolds might allow enzymatic activities to be uniquely targeted to discrete subnuclear sites¹²; RNA processors might enable complex regulatory functions to be gated in response to external stimuli or small molecules^{9–11, 19}.

Finally, CRISP-Disp may provide a platform for the isolation of novel functional RNA domains. That the sgRNA scaffold can accommodate an expansive breadth of internally inserted sequences (Fig. 3c,d), suggests that it could be used as the backbone for SELEX. Such selections might yield, for example, aptamers that sequester or recruit endogenous protein complexes to target loci, potentially allowing host proteins to be re-targeted genome-wide.

The present studies establish a preliminary framework for the implementation of CRISPR-Display, including “best practices” for construct design (Supplementary Data). Moreover, the proof-of-principle experiments presented here hint to the larger scope of novel applications CRISPR-Display enables (Fig. 4e). We anticipate that this application list is, at best, preliminary.

ONLINE METHODS

Plasmid synthesis

Mammalian expression and reporter constructs were generated using standard restriction enzyme-based and ligation-independent cloning methods. Components were acquired as follows: The T7 promoter-targeting TALE³⁰ was the generous gift of Feng Zhang (Broad Institute). *Gaussia* and *Cypridina* luciferases were derived from pGLuc-Basic and pCLuc-Basic, respectively (New England Biolabs). dCas9 (*S. pyogenes* D10A/H841A Cas9) was isolated from Addgene plasmid 47754, the EF1a promoter from Addgene plasmid 11154, mCerulean from Addgene plasmid 23244, Venus from Addgene 15753 and the human Ubiquitin C promoter (hUBCPro) used to drive expression of L7Ae~VP, PP7~VP (Supplementary Fig. 2a, *bottom*) and MS2~mCherry (Supplementary Fig. 13a, *right*) from Addgene plasmid 17627. All other components were synthesized *de novo* from small synthetic oligonucleotides or from gBlocks (Integrated DNA Technologies).

The backbone for Lentiviral reporter constructs was derived from pLenti6.3/TO/V5-DEST (Life Technologies), from which the Tet-reponsive promoter and Gateway cloning sites were removed. The backbone for the T7 TALE and MS2~VP constructs was derived from

pcDNA3.1(+) (Life Technologies) in which the Neomycin expression cassette was removed. All other constructs were cloned into pNEB193 (New England Biolabs).

L7Ae, MS2 and PP7 were codon-optimized for expression in human cells and synthesized as gBlocks (Integrated DNA Technologies). The PP7 construct consists of two tandem copies of the non-aggregating FG mutant³¹ joined by a flexible seven amino acid linker with the sequence GSTSGSG (Supplementary Fig. 2a, *bottom*). Similarly, the MS2 construct consists of two tandem copies of the non-aggregating V75E/A81G mutant⁴⁰ joined by the same linker. L7Ae was designed according to a published sequence³⁷.

INT-like constructs (Figs. 3,4 and Supplementary Table 5) were cloned as follows. We first cloned an INT general-purpose cloning vector, “sgINTgpc,” containing the following pertinent sequence:

```
GATCTAGATACGACTCACTATGTTTAAGAGCTATGCTGCGAATACGAGAAGTCTTCTTTTTTGA
AGACAATCGTATTCGCAGCATAGCAAGTTTAAATAAGGCTAGTCCGTTATCAACTTGAAAAAGT
GGCACCGAGTCGGTGCTTTTTTTT
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...Wherein italicized nucleotides denote the GLuc-targeting protospacer sequence (Supplementary Table 2), underlined nucleotides denote an extended sgRNA stem1 (Ref. 18) and bold nucleotides denote two outward-facing *BbsI* restriction sites. This cassette is under expression of a human U6 promoter (*not shown*). Inserts cloned into this backbone had the general format: 5′-CGAG-[Insert]-CTCGT-3′, wherein underlined nucleotides denote the sticky ends used for cloning; the additional C following the insert restores base-pairing at the end of stem1. These inserts were generated by PCR and restriction digestion with *BbsI*, or by annealing synthetic, 5′-phosphorylated oligonucleotides (following the protocol used for the N₂₅ pool, *below*). Inserts were ligated into *BbsI*-digested, gel-purified sgINTgpc using the Quick Ligation Kit (New England Biolabs).

All sgRNAs and derivatives were initially cloned bearing a GLuc-targeting protospacer (Supplementary Table 2). *ASCL1*-, *ILIRN*-, *NTF3*-, *TTN*- and telomere-targeting constructs (Figs 1d, 3f, 4b–d; Supplementary Table 2) were derived from these parental constructs using an inverse-PCR method, using a forward primer that anneals downstream of the protospacer and a reverse primer that anneals to the 3′-end of the U6 promoter. Namely, PCR products were amplified with primers of the general format:

```
Forward: TAGTAGAAGACAAXXXXXXXXXXXXGTTTAAGAGCTATGCTGCGAATACG
Reverse: TAGTAGAAGACAAYYYYYYYYGGTGTTCGTCCTTCCAC
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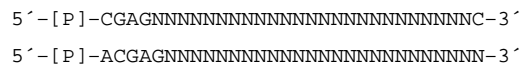
...Wherein bold nucleotides denote *BbsI* restriction sites; X’s denote nucleotides 9–21 of the new protospacer sequence; Y’s denote the reverse complement of nucleotides 1–9 of the new protospacer; underlined nucleotides are reverse complementary to one another. PCR products were purified using the QIAgen PCR cleanup kit, digested with *BbsI* and *DpnI*,

purified again and quantified by UV-vis spectroscopy. Products (25 ng, in 11 μ L final) were self-ligated using the Quick Ligation Kit (New England Biolabs).

All plasmid sequences were confirmed by Sanger sequencing (GeneWiz) prior to use.

Cloning the N₂₅ Pool

Pool oligonucleotides (Integrated DNA Technologies) were as follows:



...Wherein 5'–[P] denotes a 5' Phosphate, and N denotes an equimolar mixture of all four nucleotides. Oligonucleotides were each resuspended in annealing buffer (10 mM Tris, pH 7.0, 50 mM NaCl) to 100 μ M. 10 μ L of each oligo were mixed in a 0.2 mL PCR tube; this mixture was heated to 95°C for 10 minutes and slowly annealed to 25°C over the course of two hours in a thermocycler. The reaction was snap-cooled on ice and diluted 100-fold with ice-cold annealing buffer. 1 μ L of this diluted duplex mix was ligated into 25 ng of *BbsI*-cut sgINTgpc, in 12 μ L final volume, using the Quick Ligation Kit (New England Biolabs). The entire reaction was transformed into 120 μ L of XL10-Gold ultracompetent cells (Agilent), plated onto 12 LB Ampicillin plates and grown overnight at 37°C. Seven bacterial colonies were picked for Sanger sequencing (Supplementary Fig. 9), and the remainder were pooled by scraping the plates into 100 mL of liquid LB(Amp). Bacteria were pelleted by ultracentrifugation, and the plasmid pool was harvested in a single plasmid maxi-prep (QIAgen) (Supplementary Figs. 9–10).

Cell Culture, stable and transient transgene expression

HEK 293FT cells (ATCC) were maintained on gelatinized plates in high glucose Dulbecco's modified Eagle's medium (DMEM, Gibco), supplemented with 10% FBS, 1 \times penicillin/streptomycin and 2 mM L-Glutamine (Gibco). Cells were grown at 37°C and 5% CO₂ in a humidified incubator.

Lentiviral particles were generated using standard second generation packaging plasmids, in 293T cells. Integrated reporter cells were generated as follows: 250,000 HEK 293FT cells were plated per well of a gelatinized six-well dish and incubated overnight. Growth media was thereafter removed; cells were washed once in warmed PBS, and supplied with 1.7 mL fresh warmed media supplemented with 200 μ L CLuc reporter lentivirus and 8 μ g/mL polybrene. After 24 hr this process was repeated with a second dose of CLuc virus. Cells were subsequently passaged onto 10 cm gelatinized plates and selected with 2 μ g/mL puromycin. CLuc reporter cells were then plated onto gelatinized six-well dishes and transduced with GLuc reporter lentivirus following the same transduction protocol. GLuc-transduced cells were not selected with hygromycin prior to use. A lentiviral variant of our EF1 α -dCas9 construct (Supplementary Fig. 1c) was also used for aptamer-based imaging (Fig. 4d) following the same transduction protocol, without antibiotic selection. To enrich for cells that expressed low levels of dCas9 (Ref. 18), we transiently transfected GLuc

reporter, U6-INT and PP7~VP plasmids, as in analytical luciferase assays (*see below*), and collected GLuc⁺ cells by FACS.

Transient transfections were performed using Lipofectamine 2000 (Life Technologies), following the manufacturer's protocol. For luciferase assays, 125,000 cells in 0.6 mL media were plated per well of gelatinized 12-well dishes and incubated overnight. Transfection mixes contained 33 ng of each luciferase reporter plasmid (where appropriate), 59 ng of dCas9 or dCas9~VP plasmid, 66 ng of PP7~VP, L7Ae~VP or MS2~VP (where appropriate), 11.6 ng of U6-driven or 542 ng of Pol II-driven sgRNA variants. For experiments using TOP3 and TOP4 (Fig. 1a,c, Supplementary Fig. 6b), 11.6 ng of a separate U6-driven gRNA plasmid was also included. For FACS, (Supplementary Figs. 1e, 2b, 4e and 6d) transfection mixes also contained 10 ng of an mCherry cotransfection control. In all cases, the total transfected plasmid mass was brought to 750 ng per well using pNEB193 (New England Biolabs) in 18 μ L final volume, with 2.25 μ L Lipofectamine 2000.

For RNA immunoprecipitation (RIP) and RIP-Seq experiments, 2.1 million cells in 10 mL growth media were plated onto gelatinized 10 cm dishes and grown overnight. Transfection mixes were as described above, but all masses and volumes were scaled 15.7-fold to account for the increase in growth area and cell number. RIP transfection mixes included each luciferase reporter to independently monitor CRISP-Disp function.

To test CRISP-Disp function at endogenous loci (Figs. 1d, 3f), cells were plated in gelatinized 12-well dishes as in standard luciferase assays. Transfection mixes were similar to those described²⁸, and contained 500 ng dCas9 or dCas9~VP plasmid, 500 ng GLuc-Targeting sgRNA construct or 500 ng of a mix containing equal masses (125 ng each) of four *ASCL1*-, *IL1RN*-, *NTF3*- or *TTN*-targeting constructs^{28, 29, 33} (Supplementary Table 2). Where appropriate, 556 ng of PP7~VP plasmid was also included. All mixes were brought to 1556 ng per well using pNEB193, in 38 μ L final volume, with 4.7 μ L Lipofectamine 2000.

For multiplexing experiments (Fig 4b) cells were plated in gelatinized 12-well dishes as above. Transfection mixes contained 250 ng dCas9 or dCas9~VP, 250 ng GLuc-targeting sgRNA variant, 250 ng of a mix containing equal masses (62.5 ng each) of four *NTF3*-targeting constructs (Supplementary Table 2), and 278 ng of PP7~VP or MS2~VP, where appropriate. In all cases the total transfected mass was brought to 1028 ng using pNEB193, in 30 μ L volume, with 3.1 μ L Lipofectamine 2000.

In bridged imaging experiments (Fig. 4c, Supplementary Fig. 13), 80,000 cells in 1 mL growth media were plated per well of untreated Nunc Lab-Tek glass two-chamber slides (Thermo Scientific). Twenty-four hours thereafter, growth media was changed, and cells were transfected with 440 ng dCas9 and 235 ng of each modified sgRNA. Where appropriate, 440 ng of PP7~VP64 and/or 100 ng of MS2~mCherry were included. The total transfected mass was brought to 1500 ng with pNEB193, in 11.4 μ L, with 4.5 μ L Lipofectamine 2000, according to the manufacturer's protocol. For aptamer-based imaging (Fig. 4d), 80,000 dCas9-transduced (“+dCas9,” *see above*) or untransduced (“-dCas9”) cells in 1 mL growth media were plated per well of Nunc Lab-Tek glass two-chamber slides that

had been treated as follows. Wells were coated with 100 µg/mL poly-L-lysine (Millipore) overnight at 4°C. The next day, wells were washed twice with ddH₂O, UV sterilized for five minutes in a biosafety cabinet, coated with 100 µg/mL rat collagen-I (Corning) and 50 µg/mL laminin (Life Technologies) for two hours at 37°C, and dried prior to plating cells. Transfections were performed 24 hours thereafter, with 600 ng (telomere- or GLuc-targeting) INT-spinach2 construct, 600 ng of pNEB193 and 4.5 ng of an mCherry cotransfection control, in a total volume of 11.4 µL, with 3.8 µL Lipofectamine 2000, according to the manufacturer's protocol. All live-cell imaging experiments were performed 48–72 hours post-transfection (*see below*).

Luciferase and FACS Assays

Luciferase assays were performed using the BioLux *Gaussia* and *Cypridina* Luciferase Assay kits (New England Biolabs), following the manufacturer's protocols. Growth media (200 µL) was harvested three days after transfection and, if not used immediately, was stored in the dark at 4°C in parafilm-sealed 96-well dishes. 20 µL of each experimental sample was manually pipetted into black-walled 96-well plates (Corning) and assayed using a FLUOstar OPTIMA Luminometer equipped with automatic injectors (BMG Labtech). *Gaussia* and *Cypridina* assays were performed in parallel. For each, a single empirically determined gain was applied to all samples within an experimental series. Each sample was injected with 50 µL of luciferase assay buffer and mixed for two seconds prior to data acquisition. Signal was integrated over 20 seconds using an open (unfiltered) top-down optic.

For each sample, experimental raw luciferase signals were background-subtracted, and the ratio of Luciferase values, (GLuc/CLuc), was calculated. Biological replicates (at least three per experiment) were used to calculate a mean value, <GLuc/CLuc>. Fold activation was then calculated relative to a control sample in which dCas9~VP was expressed in the absence of an sgRNA construct:

$$\text{Fold Activation} = \frac{\left\langle \frac{GLuc}{CLuc} \right\rangle (\text{Experimental sample})}{\left\langle \frac{GLuc}{CLuc} \right\rangle (dCas9 \sim VP \text{ alone})}$$

Statistical significance testing likewise used this dCas9~VP control as the basis of comparison.

For FACS assays, cells were propagated and transfected in gelatinized 12-well dishes, as described for luciferase assays, and analyzed three days after transfection. Cells were harvested by trypsinization, quenched by the addition of chilled growth media, diluted threefold in chilled staining media (Hank's Balanced Salt Solution (HBSS, Gibco), supplemented with 2% Donor Bovine Serum (DBS, Atlanta Biologicals)), and pelleted at 200 g in a swinging bucket rotor. Cells were resuspended in chilled staining media and analyzed on a BD LSR II Flow Cytometer (BD Sciences), equipped with HcRed, CFP and YFP filters. Voltages, compensations and gates were empirically determined using unstained and single color controls, according to standard methods. 100,000 mCherry⁺ cells were recorded from each sample.

RNA Immunoprecipitation (RIP)

Cells were propagated on gelatinized 10-centimeter dishes, transfected as described above, and harvested three days after transfection. Thereafter, RIP was performed essentially as described previously⁴¹. Growth media was aspirated, and cells were washed twice with 10 mL room temperature PBS (Gibco). Cells were crosslinked by incubation in 0.1% (v/v) formaldehyde in PBS for 10 minutes at room temperature, under very gentle agitation. Crosslinking was quenched by the addition of Glycine to 133 mM and gentle agitation for an additional five minutes at room temperature, after which the liquid phase was aspirated. Crosslinked cells were washed twice with room temperature PBS, harvested by scraping, allotted into samples of 1×10^7 cells (typically three samples per 10 cm dish), and pelleted at 200 g in a swinging bucket rotor. PBS was aspirated and cell pellets were flash-frozen in liquid nitrogen and stored at -80°C until use.

Cell pellets were thawed on ice, gently resuspended into 1 mL of ice-cold RIPA(+) buffer (standard RIPA supplemented with 0.1 U/ μL RNaseOUT (Life Technologies), $1 \times$ EDTA-free Proteinase Inhibitor Cocktail (Thermo Scientific) and 0.5 mM DTT), and lysed for 10 minutes at 4°C with end-over-end agitation. Samples were then sheared using a Branson Digital Sonifier 250 (Emerson Industrial Automation) at 10% amplitude for three 30-second intervals (0.7 seconds on + 1.3 seconds off), with 30-second resting steps between intervals. Samples were held in ice-cold metal thermal blocks throughout sonication. Sheared samples were then clarified by ultracentrifugation and diluted with 1 mL each of ice-cold Native Lysis Buffer(+) (25 mM Tris, pH 7.4, 150 mM KCl, 5 mM EDTA, 0.5% (v/v) NP-40, supplemented with inhibitors and DTT, as above), filtered through a $0.45 \mu\text{m}$ syringe-mounted filter, and flash-frozen in liquid nitrogen before use.

Clarified lysates were thawed on ice and pre-cleared by incubation with buffer-equilibrated magnetic Protein G beads (Life Technologies) for 30 minutes at 4°C , with end-over-end rotation. 100 μL aliquots were removed and frozen, to serve as “input” normalization controls. Cleared lysates corresponding to 5×10^6 cells were then incubated with 6 μg rabbit anti-FLAG (SIGMA) or Rabbit normal IgG (Cell Signaling Technology), for two hours at 4°C with end-over-end rotation. Buffer-equilibrated magnetic Protein G beads were then added and the samples were again rotated end-over-end for one hour at 4°C . Beads were collected and twice washed with Native Lysis Buffer(+) for 10 minutes at 4°C , with end-over-end rotation. Immunoprecipitated RNA was thereafter isolated as described below.

RNA Isolation, Quantitative RT-PCR and mRNA Seq

Whole cell RNA (Figs. 1d, 3f, 4b and Supplementary Figs. 5 and 12) and RNA from subcellular fractions (Supplementary Fig. 7) were isolated by extraction with Trizol and Trizol-LS Reagent (Life Technologies), respectively, following the manufacturer’s protocols. RNA was precipitated with isopropanol using GlycoBlue (Life Technologies) as a carrier, and subsequently purified using RNEasy spin columns (QIAGEN), following the manufacturer’s “RNA Cleanup” protocol, with on-column DNase treatment.

RNA from RIP and RIP-Seq experiments (Figs. 2c,e, 3d and Supplementary Figs. 2d, 4f, and 6e) was isolated as follows. Following RIP (*see above*), protein G beads were suspended

in 56 μL nuclease-free water, and processed alongside input samples (56 μL ; 5.6% of the total). All samples were brought to 100 μL with 3 \times Reverse Crosslinking Buffer (final concentrations: 1 \times PBS, 2% N-Lauroyl Sarcosine, 10 mM EDTA, 5 mM DTT, 0.4 U/ μL RNaseOUT and 2 mg/mL proteinase K (Ambion)). Formaldehyde crosslinks were reversed by incubation in a thermocycler at 42°C for one hour, and then 55°C for one hour. RNA was thereafter purified using four volumes (400 μL) Agencourt RNAClean XP Beads (Beckman Coulter), following the manufacturer's protocol, and eluted into 30 μL nuclease-free water. Residual DNA was removed by treatment with 5 U RNase-free DNAs (RQ1, Promega) in 50 μL , following the manufacturer's protocol. RNA was subsequently purified using four volumes (200 μL) Agencourt RNAClean XP beads, eluted into 20 μL nuclease-free water, and stored at -20°C until use.

cDNA was synthesized using SuperScript III Reverse Transcriptase (Life Technologies), according to the manufacturer's protocol, priming from anchored oligo-dT₂₁, random hexamers (Life Technologies) or a gene specific primer (Integrated DNA Technologies), where appropriate. Target RNA abundance was quantified by qRT-PCR on a 7900HT Fast Real-Time PCR System (Applied Biosystems), using Rox-normalized FastStart Universal SYBR Green Master Mix (Roche) and gene-specific primers (Supplementary Table 3), in quadruplicate. Non-reverse-transcribed RNA was used as a negative control. "Clipped" data were processed using Realtimer PCR Miner⁴², to calculate C_T and efficiency values (Supplementary Table 3). Bulk gene expression measurements (Figs. 1c, 3f, 4b and Supplementary Fig. 12) were normalized to a GAPDH internal control (Supplementary Table 3); RIP measurements were normalized to input RNA levels. In subcellular fractionation experiments (Supplementary Fig. 7), the yield of RNA in each compartment was quantified relative to the unfractionated input level, as in RIP experiments. Data analysis was performed using standard methods.

For global gene expression analysis (Supplementary Fig. 5), Poly(A)⁺ mRNA seq libraries were prepared using the TruSeq RNA sample preparation kit, v2 (Illumina) as described⁴¹. Indexed libraries were pooled and subjected to 50 cycles of paired end sequencing, followed by 25 cycles of indexing, on two lanes of an Illumina HiSeq 2500 (FAS Center for Systems Biology, Harvard). For characterization of gene expression, sequencing reads were mapped to a custom gene set comprising UCSC known human genes (hg19), appended with dCas9, GLuc, CLuc and sgRNA constructs, using TopHat2 with default options⁴³. Differential analysis of gene expression was assessed using Cuffdiff2 with default options⁴⁴. Genes plotted in (Supplementary Fig. 5) were restricted to the top 75% of expressed genes, based on FPKM values.

Error Propagation and Reproducibility

For Luciferase and qRT-PCR assays, experimental uncertainties were propagated as described previously⁴⁵. Given S , the sum or difference of values A , B , uncertainly was calculated using the formula:

$$\sigma_S = \sqrt{(\sigma_A)^2 + (\sigma_B)^2}$$

...wherein σ_A and σ_B are the measurement errors of A and B, respectively. For P, the product or quotient of values A and B, uncertainty was calculated using the formula:

$$\sigma_P = P \times \sqrt{\left(\frac{\sigma_A}{A}\right)^2 + \left(\frac{\sigma_B}{B}\right)^2}$$

The uncertainty of other functions, $F(x)$, was calculated using the first derivative approximation:

$$\sigma_{f(x)} = \sigma_x \times f'(x)$$

Sample sizes were determined in accordance with standard practices used in similar experiments in the literature; no sample-size estimates were performed to ensure adequate power to detect a prespecified effect size. Experiments were neither randomized nor blinded to experimental conditions. No samples were excluded from analysis.

Subcellular Fractionation

Cytoplasmic and nuclear fractions (Supplementary Fig. 7) were isolated as described^{46, 47}. Briefly, cells were grown and transfected in gelatinized 10-cm dishes, as described for RIP experiments, above. Three days after transfection, cells were harvested by trypsinization, quenched with growth media, pelleted and washed thrice with ice-cold PBS. Cells were gently resuspended in five packed cell pellet volumes ("cv's") of ice-cold Cyto Extract Buffer(+) (20 mM Tris, pH 7.6, 0.1 mM EDTA, 2 mM MgCl₂, supplemented with 0.5 U/ μ L RNaseOUT and 1 \times EDTA-free Proteinase Inhibitor Cocktail), and swollen by incubation at room temperature for two minutes, and on ice for ten minutes more. Cells were then lysed by addition of CHAPS to 0.6% final, gentle pipetting, and two passages through a syringe equipped with a 20G needle. Lysate was clarified by centrifugation at 500g in a tabletop microcentrifuge at 4°C; 70% of the resulting supernatant was retrieved as the cytoplasmic fraction. The pellet, corresponding to nuclei and cell debris, was washed twice by gentle resuspension into five cv's of Nuclear Wash Buffer(+) (Cyto Extract Buffer, supplemented to 0.6% CHAPS and with inhibitors, as above), followed by centrifugation at 500g. Washed nuclei were gently resuspended into two cv's of Nuclei Resuspension Buffer(+) (10 mM Tris, pH 7.5, 150 mM NaCl, 0.15% (v/v) NP-40, supplemented with inhibitors, as above) layered onto a cushion of five cv's Sucrose Buffer(+) (10 mM Tris, pH 7.5, 150 mM NaCl, 24% (w/v) Sucrose, plus inhibitors), and pelleted at 14,000 rpm in a tabletop microcentrifuge at 4°C. The resulting pelleted nuclei were resuspended into two cv's of ice-cold PBS and pelleted at 500g, before harvesting RNA. We confirmed the success of our fractionations by two methods: western blotting and qRT-PCR (Supplementary Fig. 7b-c). In western blots, aliquots of whole cell lysate, the cytoplasmic fraction and PBS-suspended nuclei were probed using antibodies against (α/β)-Tubulin and Fibrillarin (Cell Signaling Technology). For qPCR, extracted RNA (*see above*) was quantified using primers against the cytoplasmic ncRNA *SNHG5* and the nuclear ncRNA *XIST* (Supplementary Table 3).

N₂₅ RNA Library Preparation, Sequencing and Analysis

For the N₂₅ RIP-Seq experiment (Fig. 3d), cell growth, transfection, RIP and RNA preparation were performed as described above, in triplicate. Seven deep sequencing libraries were prepared: one from the starting plasmid pool, three from replicates of the input RNA, and three from replicates of the immunoprecipitated RNA. The plasmid pool library was generated directly via PCR, using 5 ng of plasmid template in a 50 µL reaction, amplified through 19 cycles of PCR with Pfu Ultra II HS polymerase (Agilent), according to the manufacturer's protocol. Gene-specific PCR primers that bracketed the N₂₅ insertion site, appended with standard Illumina adapters and indexes, were used (Supplementary Table 6, and Supplementary Fig. 9). For each input and RIP library, 10 ng RNA was reverse-transcribed in 20 µL as described above, using a gene specific primer (Supplementary Table 6, and Supplementary Fig. 9). Each cDNA reaction was used in its entirety as PCR template, using the same primer design as was used for the plasmid pool, but with different Illumina indexes. The pools were amplified in 200 µL, through 26 cycles of PCR with Pfu Ultra II HS polymerase (Agilent), according to the manufacturer's protocol. The resulting deep sequencing libraries were purified twice with 1.0 volume of Agencourt AMPure XP Beads (Beckman Coulter), according the manufacturer's protocol, and eluted in EB Buffer (QIAGEN). The plasmid pool library contained traces of high molecular weight contaminants (*not shown*) that were removed by "reverse selection:" the sample was treated with 0.65 volumes of AMPure XP Beads, and the unbound fraction was retained. The integrity and concentration of each final library was measured using a "DNA High Sensitivity" assay on an Agilent 2100 model Bioanalyzer (Supplementary Fig. 9).

Libraries were denatured in 50 mM NaOH, diluted in buffer HT1 (Illumina) and combined to yield a 20 pM pool, according to standard protocols. This pool was doped with TailorMix Indexed PhiX Control Library (SeqMatic), at a ratio of 7:3 N₂₅Pool:PhiX, and sequenced on two lanes of an Illumina HiSeq 2500 (FAS Center for Systems Biology, Harvard) for 150 cycles, followed by 25 cycles of indexing.

Random insert sequences were extracted from raw sequencing reads by removing the constant sequences abutting each side of the insertion point. The number of occurrences of each random sequence within each individual sample was then tabulated. Sequence counts were used to calculate enrichment using DESeq2 (Ref.⁴⁸).

Live Cell Imaging

Images in (Supplementary Fig. 1d) were collected on an Axio Observer D1 system (Zeiss), equipped with eYFP and eCFP filters.

Live fluorescence images (Fig. 4c–d, and Supplementary Figs. 13–14) were taken with an LSM 700 Inverted Confocal Microscope (Harvard Center for Biological Imaging), with an aperture setting of 1 A.U., using the DAPI filter for Hoechst 33342, the CFP filter for mCerulean, the mCherry filter for mCherry and the FITC filter for DFHBI-1T, where appropriate. In bridged imaging experiments (Fig. 4c; Supplementary Fig. 13), cells were imaged two days post-transfection, in their growth media. Images are max-merges of 37–47 Z-stacks, taken with a step size of 0.33 µm, at 63× magnification. For aptamer-based

imaging (Fig. 4d and Supplementary Fig. 14), growth media was replaced with imaging media (Fluorobrite DMEM (Life Technologies), 25 mM HEPES, 5 mM MgSO₄, 1 µg/ml Hoechst 33342 (Life Technologies), and 20 µM DFHBI-1T (Lucerna)) for 30 minutes at 37°C. Images in (Fig. 4c–d) are max-merges of 20–30 Z-stacks, taken with a step size 0.35 µm, at 63× magnification.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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D.M.S. and J.L.R. have filed for patents concerning the design and expression of extended sgRNAs to expand the repertoire of dCas9 function.

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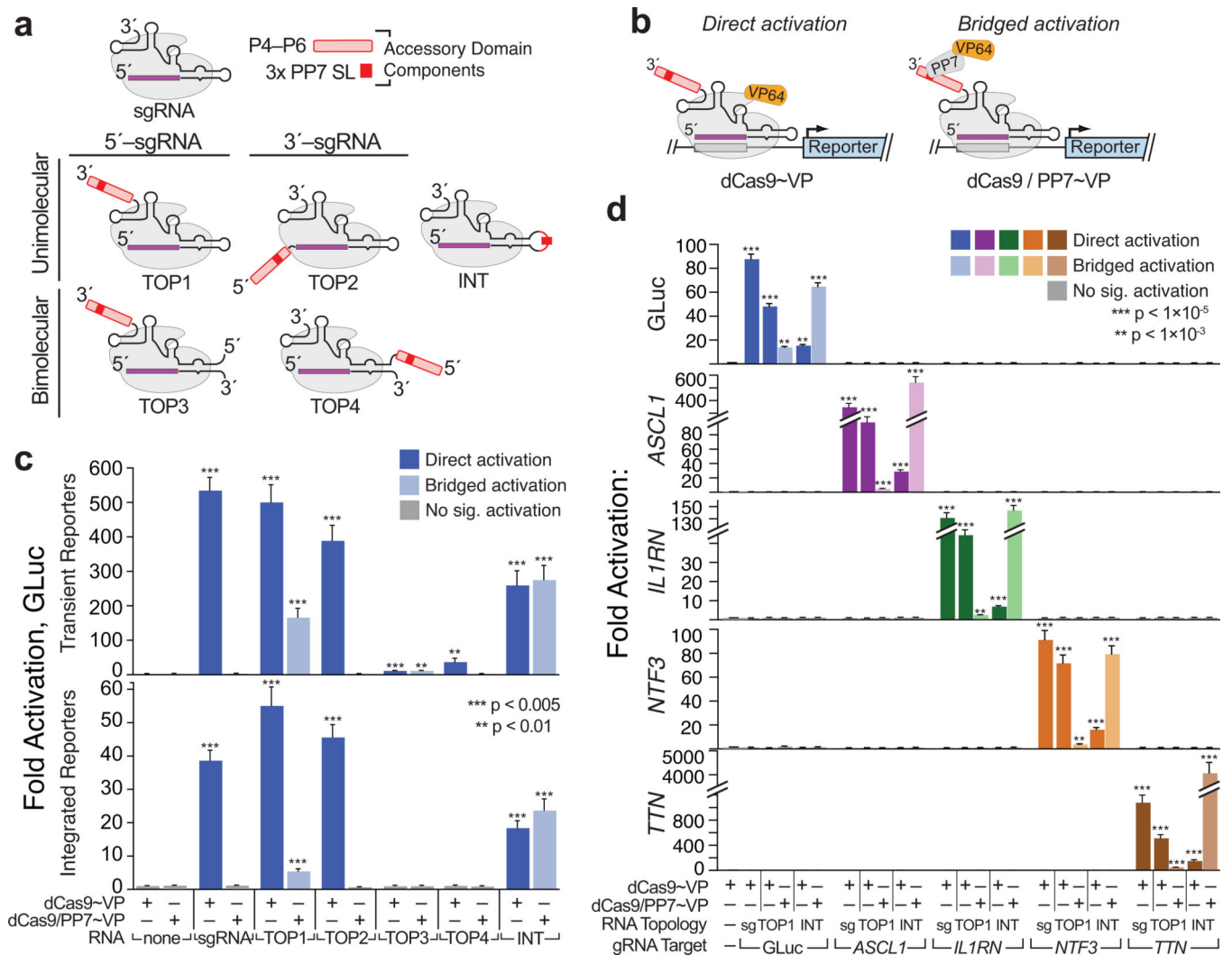


Figure 1. Large, structured RNA domains can be functionally appended onto the sgRNA scaffold at multiple points

(a) Design of “TOP” topology constructs. Accessory RNA domains are detailed in (Supplementary Fig. 4a–b); expression constructs in (Supplementary Fig. 1c). (b) Schematics summarizing direct activation (*left*) and bridged activation (*right*) assays. (c) Luciferase reporter assays of the five topology constructs. (d) Targeting minimal (“sg”) and expanded (“TOP1” and “INT”) sgRNAs to endogenous loci. *ASCL1*, *IL1RN*, *NTF3* and *TTN* were each targeted using mixed pools of four gRNAs^{28, 29, 33} (Supplementary Table 2). GLuc activation was measured by luciferase assays; endogenous gene activation was measured using qRT-PCR (Supplementary Table 3). Values are means \pm standard deviation, (n=3) Luciferase assays; (n=4) qRT-PCR. Student’s one-tailed t-test relative to cells expressing dCas9~VP alone.

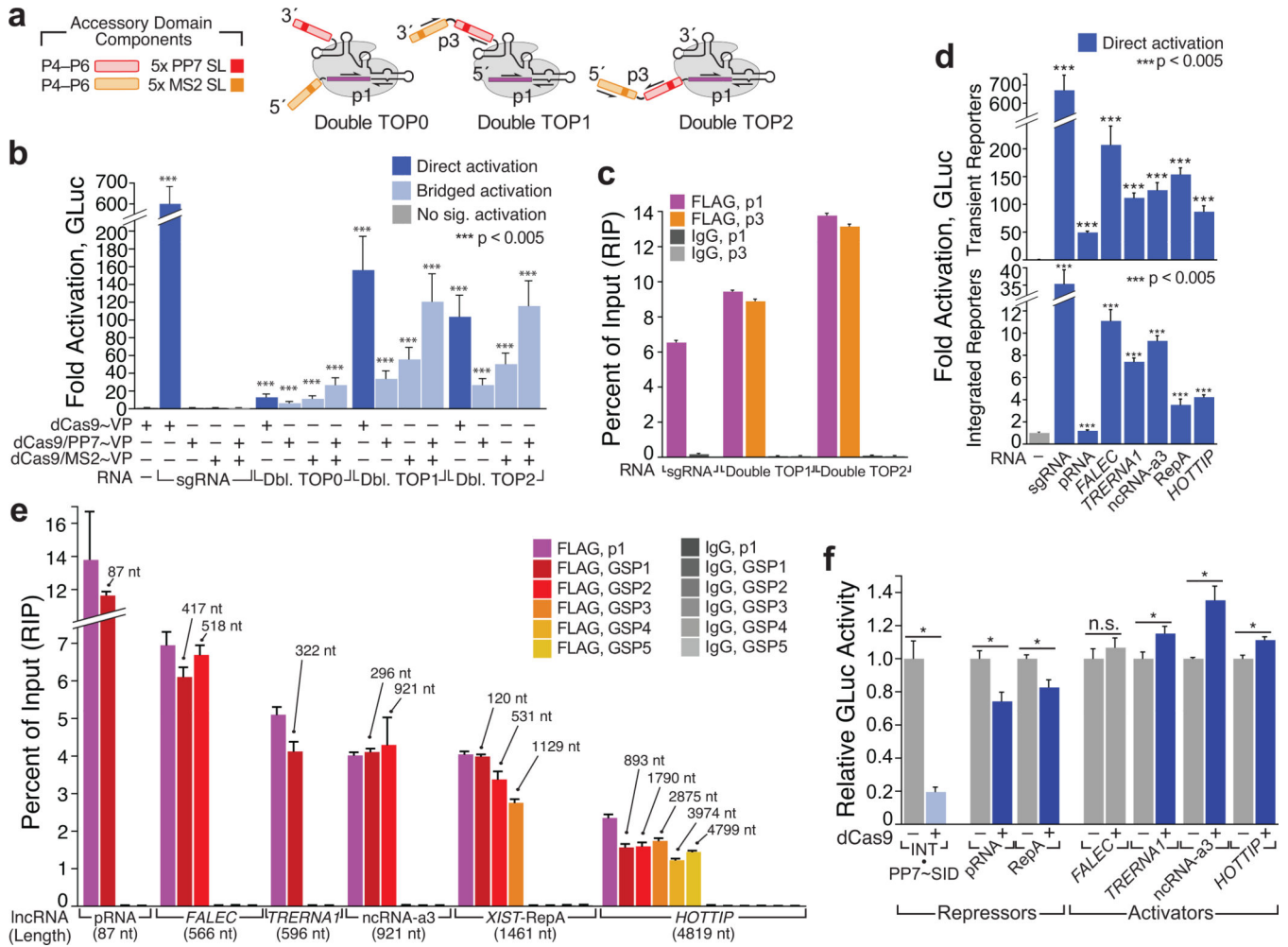


Figure 2. CRISPR-Display with artificial and natural lncRNAs

(a) Design of “Double TOP” artificial lncRNA constructs. Accessory domains are detailed in (Supplementary Fig. 4a–d). The sgRNA core “p1” and domain-spanning “p3” qPCR primer pairs are indicated (Supplementary Table 3). (b) Direct and bridged activation assays using Double TOP (“Dbl TOP”) constructs, expressed from the CMV/3’ Box backbone (Supplementary Fig. 6a). Transient reporter assays are shown. (c) RIP/qRT–PCR of dCas9•Double TOP1 and dCas9•Double TOP2. (d) sgRNAs appended with a battery of natural lncRNA domains (Supplementary Table 4) form functional complexes with dCas9~VP. Direct activation assays are shown. The minimal “pRNA” stem-loop³⁴ was displayed internally; all other domains were appended on the sgRNA 3’ terminus. RNAs were expressed using the CMV/MASC system (Supplementary Fig. 6a). (e) lncRNA accessory domains remain intact in CRISP-Disp complexes. Immunopurified RNA was analyzed using qPCR primers targeting the sgRNA core (p1) and intervals along each lncRNA domain (GSP1–GSP5, Supplementary Table 3). Above each primer set, the maximum distance from the sgRNA core is indicated. (f) Transient reporter assays with CRISP-Disp lncRNA constructs. Values are normalized relative to those of control cells expressing each lncRNA alone. For comparison, bridged repression with U6-driven INT, complexed with dCas9 and PP7~SID (Ref.³⁹) is shown (*left, light blue*). Values are means ±

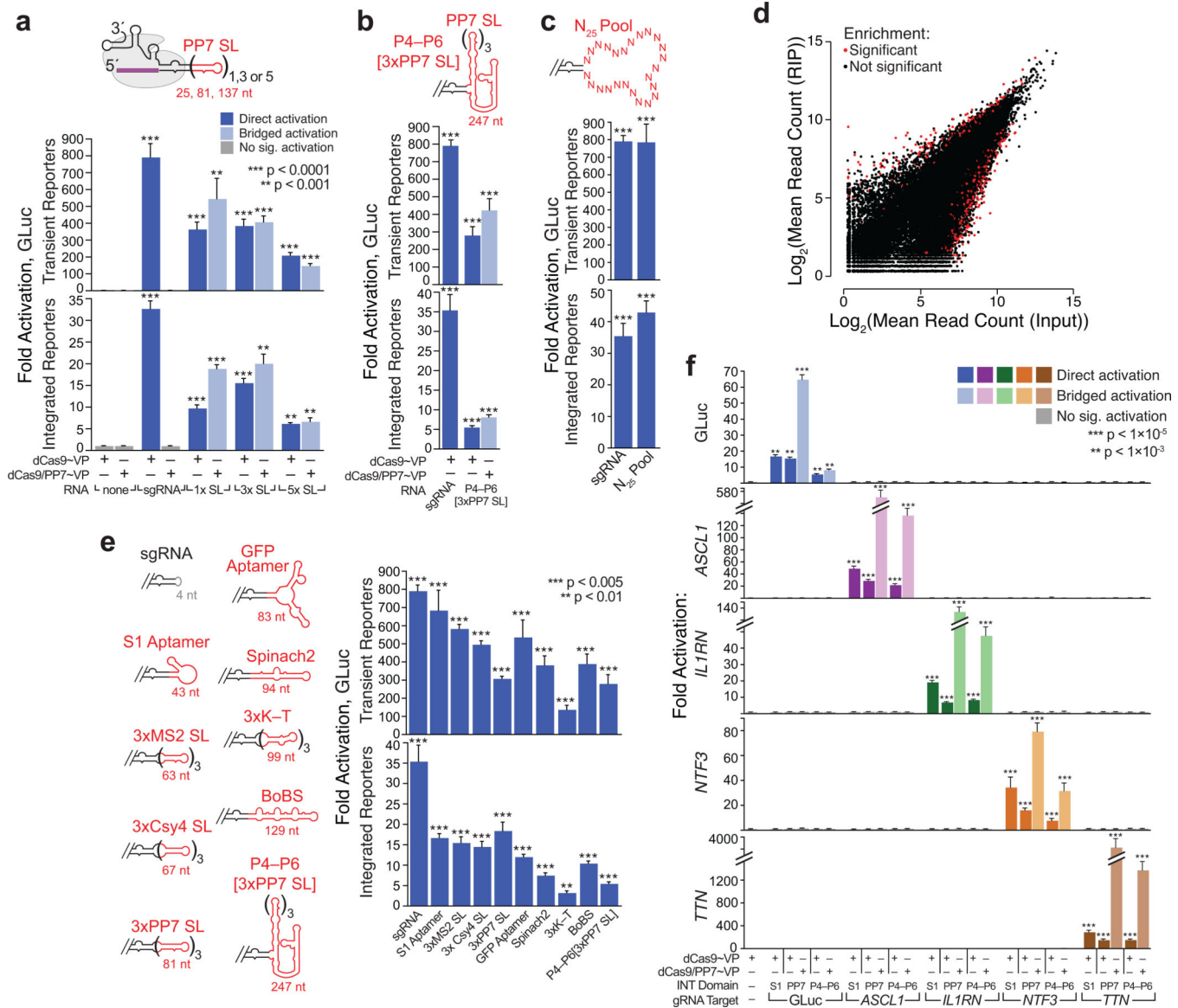
standard deviation, (n=3) Luciferase assays; (n=4) qPCR. Student's one-tailed t-test relative to cells expressing (b, d) dCas9~VP alone or (f) lncRNA alone. *, $p < 0.05$.

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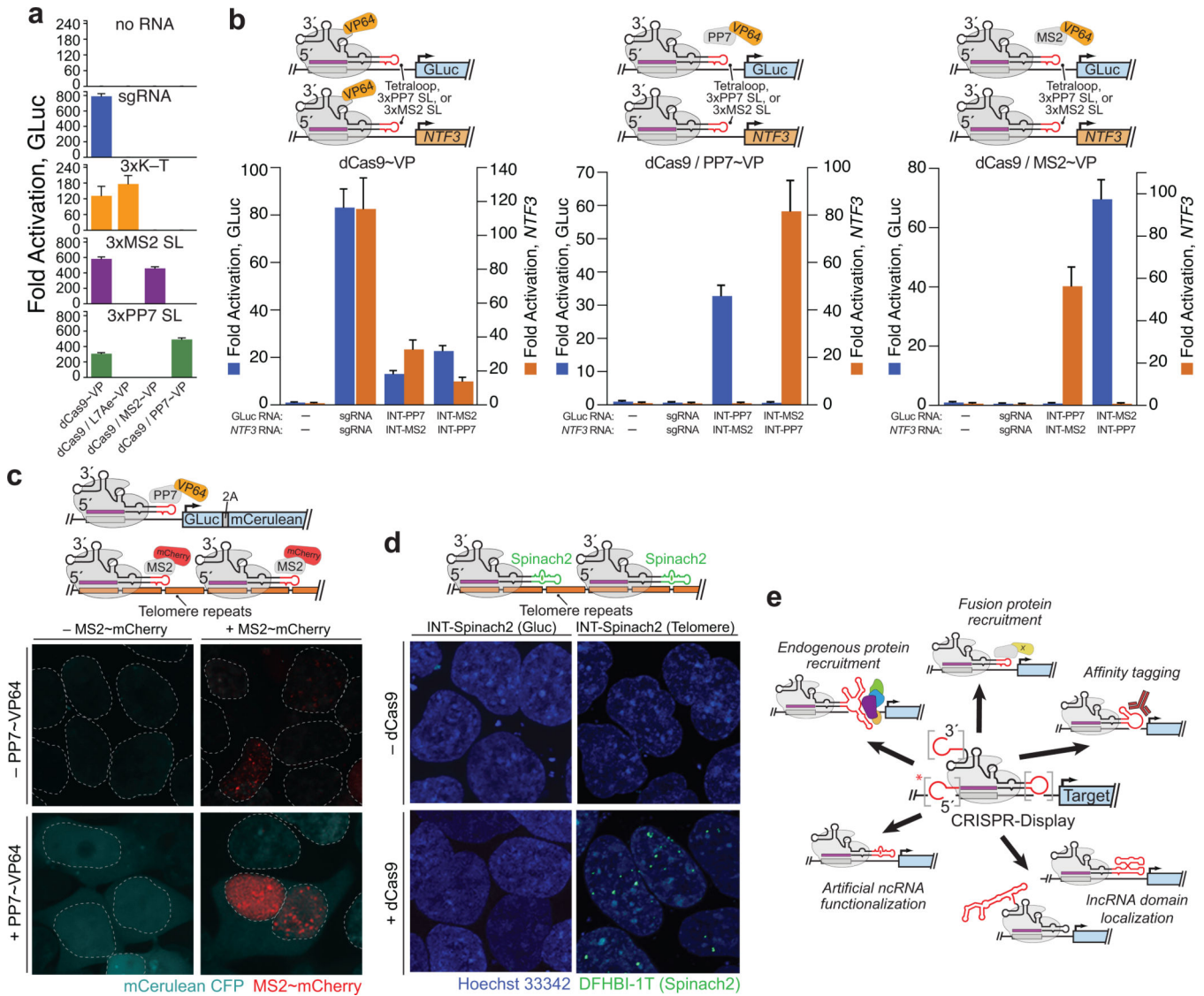
endogenous loci. Constructs were targeted to *ASCL1*, *IL1RN*, *NTF3* and *TTN*, as in (Fig. 1d). Data for the original INT (“PP7”) are included for comparison. Values are means \pm standard deviation, (n=3) Luciferase assays; (n=4) qPCR. Student’s one-tailed t-test relative to cells expressing dCas9~VP alone. RNA constructs were expressed from the human U6 promoter.

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Novel, multiplexable functions made possible by CRISPR-Display. Accessory domains on the sgRNA 5' end (*asterisk*) are not tolerated by most expression systems (Supplementary Fig. 6). Values in **(a, b)** are means \pm standard deviation. (n=3) Luciferase assays; (n=4) qPCR. Confocal fluorescence images **(c, d)**, at 63 \times magnification.

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