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(71) Applicant: THE JACKSON LABORATORY [US/US]; 600 Main Street, Bar Harbor, ME 04609 (US).

(72) Inventors: WANG, Haoyi; Building 956, Unit 1, Room 202, Zhong Guan Cun Nan San Street, Haidian District (CN). CHENG, Albert; 224 New Britain Avenue, #12, Unionville, CT 06085 (US). JILLETTE, Nathaniel; 18 Lakeshore Dr., Apt. A2, Farmington, CT 06032 (US).

(74) Agents: LU, Yu et al.; McCarter & English, LLP, 265 Franklin Street, Boston, MA 02110 (US).

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A THREE-COMPONENT CRISPR/CAS COMPLEX SYSTEM AND USES THEREOF

REFERENCE TO RELATED APPLICATIONS

This is an International Patent Application claiming priority to U.S. Provisional Application Nos. 62/132,644, filed on March 13, 2015, and 62/221,249, filed on September 21, 2015, the entire contents of both applications are incorporated herein by reference in their entirety.

BACKGROUND OF THE INVENTION

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In the CRISPR/Cas system, Cas9 protein and sgRNA (single guide RNA) constitute a sufficient two-component DNA endonuclease whose specificity is provided by target-matching sequence on sgRNA while endonuclease activity resides on the Cas9 protein.

Nuclease-defective or nuclease-deficient Cas9 protein (*e.g.*, dCas9) with mutations on its nuclease domains retains DNA binding activity when complexed with sgRNA. dCas9 protein can tether and localize effector domains or protein tags by means of protein fusions to sites matched by sgRNA, thus constituting an RNA-guided DNA binding enzyme. dCas9 can be fused to transcriptional activation domain (*e.g.*, VP64) or repressor domain (*e.g.*, KRAB), and be guided by sgRNA to activate or repress target genes, respectively. dCas9 can also be fused with fluorescent proteins and achieve live-cell fluorescent labeling of chromosomal regions. However, in such systems, only one Cas9-effector fusion is possible because sgRNA:Cas9 pairing is exclusive. Also, in cases where multiple copies of protein tags or effector fusions are necessary to achieve some biological threshold or signal detection threshold, multimerization of effector or protein tags by direct fusion with dCas9 protein is technically limited, by constraints such as difficulty in delivering the large DNA encoding such fusions, or difficulty in translating or translocating such large proteins into the nucleus due to protein size.

SUMMARY OF THE INVENTION

The invention described herein enables multiplexity and polymerization of effector or protein tags, by providing a three-component CRISPR/Cas complex / system comprising a Cas9 protein (e.g., a wildtype (wt) Cas9, a Cas9 Nickase, or a dCas9 protein), a modified sgRNA as a subject polynucleotide (e.g., "sgRNA-PBS"), and one or more fusion proteins of PUF domain(s) with effector domains or protein tags ("PUF domain-fusion[s]"). sgRNA-PBS can be derived by inserting multiple copies of short PUF (e.g., 8-mer) recognition sequences downstream of the

sgRNA stem loops or upstream of the target-matching region. PUF domains of each PUF domain-effector fusion can be programmed to recognize the 8-mer recognition sequence on the subject polynucleotide, thus bringing the one or more effector domains fused to the PUF domains to specific regions of a target DNA recognized by the target-matching sgRNA.

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The three-component CRISPR/Cas complexes / systems of the invention are advantageous in terms of multiplicity, since different three-component CRISPR/Cas complexes / systems can be simultaneously delivered into a cell or animal, and each can operate at the defined target sites with orthogonality (*i.e.*, without interference with other three-component CRISPR/Cas complexes / systems and their target sites). Since PUF domains can be easily programmed to recognize any 8-mer RNA recognition sequences, this system expands the multiplexibility to a theoretical maximum of 4⁸ (65536) when the RNA recognition sequence is only 8-mer (and potentially much more when the RNA recognition sequence is longer).

The three-component CRISPR/Cas complexes / systems of the invention are also advantageous in terms of polymerizability: the simplicity of the linear 8-mer sequence allows extensive polymerization without hindering Cas9:sgRNA DNA binding activity. Such feature allows multiple molecules of PUF-fusions to be assembled on the modified sgRNA, thus allowing local concentration of effector or protein tags. Such feature is particularly beneficial in applications such as fluorescent imaging or transcriptional regulation, where proximity synergism allows maximal effective regulation or signal-to-noise ratio.

A further advantage of the invention relates to stoichiometric complex formation. Different 8-mer sequences can be orderly inserted onto the sgRNA-PBS construct to allow complex formation with defined stoichiometry and ordering of the PUF-fusions on the sgRNA-PBS.

Thus one aspect of the invention provides a polynucleotide comprising: (1) a DNA-targeting sequence that is complementary to a target polynucleotide sequence; (2) a Cas9-binding sequence; and, (3) one or more copies of a PUF domain-Binding Sequence (PBS), wherein each of said one or more copies of the PBS binds to the same or a different PUF domain; wherein a Cas9 protein (*e.g.*, a wildtype (wt) Cas9, a Cas9 Nickase, or a dCas9 protein) is capable of forming a complex with the polynucleotide by binding to the Cas9-binding sequence.

As used herein, "Cas9 protein" include a wildtype Cas9 protein, a Cas9 nickase in which one of the two catalytic sites for endonuclease activity (RuvC and HNH) is defective or lacks activity, and a dCas9 protein in which both catalytic sites for endonuclease activity are defective

or lack activity. In certain embodiments, the Cas9 protein is a wt Cas9. In certain embodiments, the Cas9 protein lacks nuclease activity or is nuclease deficient. In certain embodiments, the Cas9 protein is a nickase (*e.g.*, for example, the nickase can be a Cas9 Nickase with a mutation at a position corresponding to D10A of *S. pyogenes* Cas9; or the nickase can be a Cas9 Nickase with a mutation at a position corresponding to H840A of *S. pyogenes* Cas9). In certain embodiments, the Cas9 protein is a dCas9 (*e.g.*, a dCas9 with mutations at positions corresponding to D10A and H840A of *S. pyogenes* Cas9). In certain embodiments, the Cas9 protein is not wt Cas9. In certain embodiments, the Cas9 protein is not nickase. In certain embodiments, the Cas9 protein is not dCas9.

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In certain embodiments, a "modified Cas9 protein" refers to a Cas9 that is not a wt Cas9 protein, such as a dCas9 or Cas9 nickase.

In certain embodiments, the dCas9 protein is nuclease-deficient but retains DNA-binding ability when complexed with the polynucleotide.

In certain embodiments, the DNA-targeting sequence base-pairs with the target polynucleotide sequence when the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) is complexed with the polynucleotide.

In certain embodiments, the target polynucleotide sequence comprises or is adjacent to a transcription regulatory element. For example, the transcription regulatory element may comprise one or more of: core promoter, proximal promoter element, enhancer, silencer, insulator, and locus control region.

In certain embodiments, the target polynucleotide sequence comprises or is adjacent to a telomere sequence, a centromere, or a repetitive genomic sequence.

In certain embodiments, the target polynucleotide sequence comprises or is adjacent to a genomic marker sequence (or a genomic locus of interest).

In certain embodiments, the target polynucleotide sequence is immediately 3' to a PAM (protospacer adjacent motif) sequence of the complementary strand, which can be 5'-CCN-3' wherein N is any DNA nucleotide.

In certain embodiments, the DNA-targeting sequence is complementary to the target polynucleotide sequence over about 12-22 nucleotides (nts), about 14-20 nts, about 16-20 nts, about 18-20 nts, or about 12, 14, 16, 18, or 20 nts (preferably, the complementary region comprises a continuous stretch of 12-22 nts, preferably at the 3' end of the DNA-binding sequence). For example, the DNA-binding sequence can be 50, 60, 70, 80, 90, or 95-100%

complementary to the target polynucleotide sequence.

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In certain embodiments, the DNA-binding sequence has a 5' end nucleotide G.

In certain embodiments, the polynucleotide further comprises a linker sequence linking the DNA-targeting sequence to the Cas9-binding sequence.

In certain embodiments, the Cas9-binding sequence forms a hairpin structure.

In certain embodiments, the Cas9-binding sequence is about 37-47 nt, or about 42 nt.

In certain embodiments, the Cas9 nickase protein lacks endonuclease activity due to point mutations at one endonuclease catalytic sites (RuvC and HNH) of wild type Cas9. The point mutations can be D10A or H840A.

In certain embodiments, the dCas9 protein lacks endonuclease activity due to point mutations at both endonuclease catalytic sites (RuvC and HNH) of wild type Cas9. The point mutations can be D10A and H840A.

In certain embodiments, each of the one or more copies of the PBS has about 8 nucleotides.

In certain embodiments, the polynucleotide comprises 1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 35, 40, 45, 46, 47, 48, 49, or 50 copies, or 1-50, 2-45, 3-40, 5-35, 5-10, 10-20 copies of identical or different PBS.

In certain embodiments, the polynucleotide comprises a PBS of the sequence 5'-UGUAUGUA-3' that can be bound by the PUF domain PUF(3-2).

In certain embodiments, the polynucleotide comprises a PBS of the sequence 5'-UUGAUAUA-3' that can be bound by the PUF domain PUF(6-2/7-2).

Another aspect of the invention provides a vector encoding any one of the subject polynucleotide.

In certain embodiments, transcription of the polynucleotide is under the control of a constitutive promoter, or an inducible promoter.

In certain embodiments, the vector is active in a cell from a mammal (a human; a non-human primate; a non-human mammal; a rodent such as a mouse, a rat, a hamster, a Guinea pig; a livestock mammal such as a pig, a sheep, a goat, a horse, a camel, cattle; or a pet mammal such as a cat or a dog); a bird, a fish, an insect, a worm, a yeast, or a bacterium.

In a related aspect, the invention provides a plurality of any one of the subject vectors, wherein two of the vectors differ in the encoded polynucleotides in their respective DNA-targeting sequences, Cas9-binding sequences, and/or the copy number, identity, or relative order

of the PBS.

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Another aspect of the invention provides a complex comprising any one of the subject polynucleotide, and the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein).

In certain embodiments, the complex further comprises one or more PUF domain(s) bound to said one or more PBS(s).

In certain embodiments, each of the PUF domains is fused to an effector domain.

In certain embodiments, the effector domain is independently a transcription repressor, a transcription activator, a fluorescent protein, an enzyme, or a chromatin remodeling protein (HDAC/HAT).

In certain embodiments, at least two of the PUF domains are fused to different effector domains.

In certain embodiments, the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), the PUF domain, and/or the effector domain further comprises a nuclear localization sequence (NLS).

In certain embodiments, the complex is bound to the target polynucleotide sequence through the DNA-targeting sequence.

Another aspect of the invention provides a host cell comprising any one of the subject vector, or the plurality of the subject vectors.

In certain embodiments, the host cell further comprises a second vector encoding the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein).

In certain embodiments, the second vector further encodes an effector domain fused to the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein).

In certain embodiments, expression of the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) is under the control of a constitutive promoter or an inducible promoter.

In certain embodiments, the host cell further comprises a third vector encoding said one or more PUF domains, each fused to an effector domain.

In certain embodiments, expression of the one or more PUF domains is independently under the control of a constitutive promoter or an inducible promoter.

In certain embodiments, the effector domain is a transcription repressor, a transcription activator, a fluorescent protein, an enzyme, or a chromatin remodeling protein (HDAC/HAT).

In certain embodiments, the second vector further encodes a nuclear localization signal fused to the Cas9 protein (e.g., wt, nickase, or dCas9 protein) or the effector domain, and/or the

third vector further encodes a nuclear localization signal fused to the PUF domain or the effector domain.

In certain embodiments, the second vector is the same as the vector, and/or wherein the third vector is the same as the vector or the second vector.

In certain embodiments, the host cell is in a live animal.

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In certain embodiments, the host cell is a cultured cell.

Another aspect of the invention provides a method of assembling the complex of the invention at the target polynucleotide sequence, the method comprising contacting or bringing to the vicinity of the target polynucleotide sequence: (1) any one of the subject polynucleotide, or any one of the subject vector, or the subject plurality of vectors; (2) the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), or any one of the subject second vector; and, (3) one or more of the PUF domains, each fused to an effector domain, or any one of the subject third vector.

In certain embodiments, the complex is assembled inside a cell, the target polynucleotide sequence is a part of the genomic DNA of the cell, and wherein the subject vector, the subject second vector, and the subject third vector are introduced into the cell.

In certain embodiments, the target polynucleotide sequence is at or near a genomic locus rich in heterochromatin, and wherein the effector domain is a detectable marker (e.g., a) fluorescent protein).

In certain embodiments, the target polynucleotide sequence is at or near a transcription regulatory element of a target gene, and wherein the effector domain is a transcription modulator (*e.g.*, activator, suppressor).

In certain embodiments, transcription of the target gene affects cell fate determination, cell differentiation, metabolic flux, or a biologically or biochemically determinable outcome.

Another aspect of the invention provides a method of modulating transcription of a plurality of target genes in a cell, the method comprising: introducing into the cell the subject plurality of the vectors, a coding sequence for a Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), and a coding sequence for one or more PUF domains, wherein each of said target genes comprises a target polynucleotide sequence that permits (1) the assembly, at the target polynucleotide sequence, of a tripartite complex of a polynucleotide encoded by one of said plurality of the vector, the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), and a PUF domain; and (2) transcription modulation of the target gene comprising the target polynucleotide sequence. In certain embodiments, the Cas9 protein is a dCas9 protein.

In certain embodiments, the transcription of at least one target gene is enhanced / stimulated, while the transcription of at least another target gene is inhibited.

In a related aspect, the invention also provides a method of epigenotic modulation (*e.g.*, modulating the epigenetic states of chromatin not directly related to transcriptional activity), at a plurality of target genes in a cell, the method comprising: introducing into the cell the subject plurality of the vectors, a coding sequence for a wt Cas9 protein or a Cas9 nickase, and a coding sequence for one or more PUF domain fusions, wherein each of the target genes comprises a target polynucleotide sequence that permits (1) the assembly, at the target polynucleotide sequence, of a tripartite complex of a polynucleotide encoded by one of the plurality of the vector, the wt / nickase Cas9 protein, and a PUF domain fusion; and (2) epigenotic modulation of the target gene comprising the target polynucleotide sequence. The method can be useful, for example, to change epigenetic state (*e.g.*, opening up the chromatin) at the same time to gain access / stability of Cas9 binding to closed chromatin sites (*e.g.*, to increase cut and genome editing at those sites).

Another aspect of the invention provides a kit comprising: (1) a subject polynucleotide, or a subject vector; (2) a subject second vector encoding the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein); and (3) a subject third vector encoding one or more PUF domains, each fused to an effector domain.

In certain embodiments, the kit further comprises transformation, transfection, or infection reagents to facilitate the introduction of said vectors into a cell.

It should be understood that any embodiments described herein, including those only described in the Example section or only under one aspect of the invention, can be combined with any one or more other embodiments, unless specifically disclaimed or otherwise improper.

BRIEF DESCRIPTION OF THE DRAWINGS

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FIGs. 1A-1D show that insertion of PUF domain-binding sequences (PBS) to sgRNA 3'-end did not substantially impact dCas9/sgRNA function, and that independent recruitment and multimerization of activators can be achieved using the subject 3-component CRISPR/Cas complex / system. FIG. 1A is a schematic drawing showing the subject 3-component CRISPR/Cas complex / system, which improves the conventional two-hybrid dCas9 fusion design by splitting it into a three-hybrid system, in which sgRNA-PBS bridges the DNA binding activity of dCas9/sgRNA with the effector function provided by a PUF fusion. The middle panels represent the structure of a representative PUF domain, showing the 8 repeats in the C to

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N direction and the corresponding interaction with the 8-mer target RNA in the 5' to 3' direction. PUF RNA recognition code table shows exemplary di-residues and the corresponding RNA base recognized. In the lower panel, a table of notation adopted for simplicity to describe the 4 PUF isotypes and the corresponding pumilio binding sites (PBS) and their sequences. FIG. 1B, upper panel, is a schematic for the experiment to test the ability of dCas9-VP64 to bind and activate a tdTomato transgene after inserting varying number of PBS at the 3' end of the sgRNA, e.g., experimental set up for testing the effect of sgRNA-PBS (with 0, 5, 15, 25, or 47 PBS) on the ability of the dCas9::VP64 construct to activate a TetO::tdTomato transgene. The lower panel is column plot showing the mean fold changes (±S.E.M.) in tdTomato fluorescence (relative to the dCas9-VP64/sgCtl-0×PBSa control), as measured by fluorescence activated cell sorting (FACS), of cells transfected with the different constructs indicated in the legend below the plot. The legend describes the sgRNA used in three parameters: sgRNA match refers to the DNA target recognized by the sgRNA; #PBS and PBS Type indicate the number and the types of PBS, respectively, appended to the end of the sgRNA. In FIG. 1C, upper panel, is a schematic describing the experiment to test activation of a TetO::tdTomato transgene by the subject activator with different numbers of appended PBS. The lower panel is a column plot showing the fold changes (±S.E.M.) of tdTomato fluorescence (relative to control dCas9/PUFb-VP64/sgCtl-0×PBSb) of cells transfected with the different constructs indicated in the legend blow the plot. The legend describes the PUF isotype (PUF-VP64) used and the sgRNA-PBS used in terms of the number and type of PBS as well as the DNA target recognized by sgRNA indicated by shaded boxes. In FIG. 1D, upper panel, is a schematic illustrating the experiment to test the independency of the subject activator isotypes in activating a TetO::tdTomato transgene. The lower panel is a column plot showing the mean fold changes (±S.E.M.) of tdTomato fluorescence (relative to the respective controls dCas9/PUFx-VP64/sgCtl-5×PBSx for PUF/PBS isotype x) of cells transfected with the different constructs indicated in the legend below the plot. The legends indicate the PUF isotype used (PUF-VP64), the PBS isotype (5×PBS; "-" indicates sgRNA without PBS) and DNA target indicated by shaded boxes (sgRNA Match). All plots show results of three replicate measurements.

FIGs. 2A and 2B relate to the assembly of the subject 3-component CRISPR/Cas complex / system comprising VP64 and P65-HSF1. FIG. 2A is a schematic of the experiment testing the assembly of PUF(3-2)::VP64 and PUF(6-2/7-2)::P65-HSF1 via recruitment by sgRNA containing both PBS32 and PBS6272. The activity was measured by the tdTomato fluorescent reporter activity. FIG. 2B is a column chart showing the relative mean tdTomato

fluorescence resulting from transfecting the activator protein(s) with non-targeting (sgControl) and Tet-targeting (sgTetO) sgRNAs with 4×[PBS32-PBS6272] heterodimer sites.

FIG. 2C shows comparison of the subject 3-component system activator using VP64 (PUFa::VP64; red columns) versus p65HSF1 (PUFa::p65HSF1; blue columns) as the activation domain in conjunction with Control sgRNA with 5×PBSa or TetO-targeting sgRNA with 0, 1, 5, 15, or 25 copies of PBSa. Columns show mean fold change (with S.E.M.; n=3) of tdTomato fluorescence relative to experiments using control sgRNA (sgCtl). The legend indicates the number of PBSa (#PBSa) on the sgRNA-PBS as well as the DNA match indicated by the shaded boxes.

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FIGs. 3A-3D show that the subject system allows for multimerization of activator to achieve robust endogenous gene activation. FIG. 3A, upper panel: a gene model showing the relative match positions (Strokes labeled 1~4) of sgRNA-PBS used to activate OCT4 gene. Lower panel: Mean fold changes (with 95% C.I.) measured by qRT-PCR (compared to the Control sample) for activation of OCT4 expression using dCas9/PUFa-p65HSF1 3-component system activator module, or dCas9-p65HSF1 activator with the indicated cocktail of OCT4 targeting sgRNA-5×PBSa or control sgRNAs-5×PBSa. The shaded boxes in the legend indicate the use of single sgRNA-5×PBSa with a control (Ctl) sequence, the individual OCT4-targeting sgRNA-5×PBSa corresponding to numbered strokes in the gene model, or a cocktail of the 4 OCT4-targeting sgRNA-5×PBSa. FIG. 3B, upper panel: a gene model showing the relative match positions (Strokes labeled 1~4) of sgRNA-PBS used to activate SOX2 gene. Mean fold changes (with 95% C.I.) measured by qRT-PCR (compared to the Ctl sample) for activation of SOX2 expression using dCas9/PUFa-p65HSF1 activator or dCas9-p65HSF1 activator with the indicated cocktail of SOX2 targeting sgRNA-5×PBSa or control sgRNA-5×PBSa. The shaded boxes in the legend indicate the use of single sgRNA-5×PBSa with a control (Ctl) sequence, the individual SOX2-targeting sgRNA-5×PBSa corresponding to the numbered strokes in the gene model, or a cocktail of 4 SOX2-targeting sgRNA-5×PBSa. FIG. 3C shows Mean fold changes (with 95% C.I.) of OCT4 expression with the indicated single or cocktails of OCT4-targeting sgRNA-PBSa with 1, 5, 15, or 25 copies of PBSa. FIG. 3D shows Mean fold changes (with 95% C.I.) of SOX2 expression with the indicated single or cocktails of SOX2-targeting sgRNA-PBSa with 1, 5, 15, or 25 copies of PBSa.

FIGs. 4A and 4B show that the subject 3-component CRISPR/Cas complex / system allows simultaneous activation and repression of two different target reporter genes. FIG. 4A is a schematic showing an experiment to simultaneously activate a TetO::tdTomato transgene with

dCas9/sgTetO-PBS32/PUF(3-2)::VP64 and repress a SV40::EGFP transgene with dCas9/sgSV40-PBS6272/KRAB::PUF(6-2/7-2). FIG. 4B is a column chart showing relative mean EGFP and tdTomato fluorescence for the samples transfected with the constructs indicated in the table.

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FIGs. 4C and 4D further demonstrate that the subject 3-component CRISPR/Cas complex / system can activate and repress different genes simultaneously. FIG. 4C, left panel: schematic diagram illustrating the experiment to achieve simultaneous activation and repression of TetO::tdTomato and SV40::EGFP by PUFc-p65HSF1 and KRAB-PUFa, respectively. Right panel: Top column plot shows mean fold changes (with S.E.M.) of tdTomato fluorescence; Bottom column plot shows mean fold changes (with S.E.M.) of EGFP fluorescence of cells transfected with constructs indicated in the central legend. The central legend indicates the inclusion by shading the transfection of PUFc-p65HSF1 and KRAB-PUFa, as well as the DNA match to either Ctl, TetO or SV40P1 of the sgRNA-PBSc and sgRNA-PBSa by the black shaded boxes. FIG. 4D, left panel: schematic diagram illustrating the experiment to simultaneously activate and repress OCT4 and SOX2, respectively by PUFb-p65HSF1 and BFPKRAB-PUFa. Right panel: Top column plot shows mean fold changes (with 95% C.I.) of gene expression of OCT4; Bottom column plot shows mean fold changes (with 95% C.I.) of gene expression of SOX2 of cells transfected with constructed indicated in the central legend. The central legend indicates the DNA match for the sgRNA-5×PBSb and sgRNA-5×PBSa to control (Ctl), OCT4 promoters (OCT4pp) or SOX2 promoters (SOX2pp) by the black shaded boxes. The PUFbp65HSF1 + BFPKRAB-PUFa row indicates the inclusion of the activator-repressor models in samples with the yellow-highlighted boxes. These experiments used cocktails of 4 sgRNA-5×PBS for both *OCT4* and *SOX2* genes.

FIG. 5A-5C show that the subject 3-Component CRISPR/Cas Complex / System can be used to recruit histone acetyltransferase (HAT) domain of CREB-binding protein (CBP) at enhancers to activate target gene expression. FIG. 5A is a schematics of enhancer activation experiment using dCas9-CBPHAT direct fusion or 3-component module dCas9/CBPHAT-PUFa or dCas9/PUFa-CBPHAT to target Proximal Promoter (PP), Proximal Enhancer (PE) or Distal Enhancer (DE) of *OCT4*. The 4 guides targeting each of these regions are shown with the number above the red strokes indicating the locations of match. FIG. 5B shows Mean fold changes (with 95% C.I.) of *OCT4* expression (relative to the corresponding sgCtl targeting experiments) of cells transfected with plasmids expressing dCas9-CBPHAT, dCas9/CBPHAT-PUFa or dCas9/PUFa-CBPHAT and cocktail of 4 sgRNA-5×PBSa targeting each of PP, PE or

Distal Enhancer DE. FIG. 5C shows Mean fold changes (with 95% C.I.) of *OCT4* expression (relative to the sgCtl experiment) after transfection of dCas9/CBPHAT-PUFa and single or cocktails of sgRNAs targeting PP, PE, DE of *OCT4*. The legend indicates the inclusion of the individual guides targeting each of the region or a cocktail of guides with the shaded boxes.

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FIGs. 6A-6G show that the subject 3-component CRISPR/Cas complex / system allows multimerization of fluorescent proteins and simultaneous labeling of telomeres and centromeres (Scale bars: 5µm). FIG. 6A is a schematic showing the use of dCas9/sgTelomere-PBS32/Clover::PUF(3-2) (or PUFa) to label telomeric repeats with green fluorescence. FIG. 6B shows confocal fluorescent microscopy images showing labeling of telomeres by Clover-PUFa and sgTelomere equipped with, from left to right, increasing number (0, 5, 15, 25) of PBSa. FIG. 6C shows anti-TRF2 immunostaining confirmation of labeling of telomeres by dCas9/Clover-PUFa/sgTelomere-25×PBSa. FIG. 6D shows quantification of the number of fluorescent foci in HEK293T cells transfected with dCas9/PUFa::Clover and a telomeretargeting sgRNA with 0,5,15 or 25 PBSa sites. (n=20; Mann-Whitney statistics: ***=p<0.0005, ****=p<0.0001). FIG. 6E shows quantification of signal-to-noise ratio as a proportion of total signal at foci over the total nuclear signal by the subject 3-component system with 5,15, or 25×PBSa on the sgRNA targeting telomeres. (n=20; Mann-Whitney statistics: ****=p<0.0001). FIG. 6F shows anti-CREST confirmation of labeling of centromeres by Clover-PUFc/sgCentromere-20×PBSc. FIG. 6G is a representative confocal fluorescent microscopy image showing the co-labeling of centromeres and telomeres by Clover-PUFc/sgCentromere-20×PBSc and mRuby2-PUFa/sgTelomere-25×PBSa, respectively.

FIG. 7 is a representative confocal microscopy image of the MUC4 labeling, showing that the subject 3-component CRISPR/Cas complex / system allows labeling of non-repeat region with 7 sgRNA-15×PBS32 targeting MUC4 locus.

FIGs. 8A-8C is a cartoon illustration highlighting some features of the subject 3-component CRISPR/Cas complex / system. FIG. 8A illustrates multiplexing: sgRNA with different PBS isotypes can recruit the effectors tethered by the cognate PUF isotypes, providing the mechanism for multiplexing dCas9 for localizing different effector functions or proteins tags at separate chromosomal loci. FIG. 8B illustrates multimerization: the short and linear feature of PBS allow sgRNA to be equipped with many copies of PBS, thus allowing recruitment of many molecules of PUF-fusions at target loci. FIG. 8C illustrates complex formation: sgRNA equipped with different combinations, orders and numbers of PBS can potentially act as a scaffold to direct assembly of protein complexes with desired stoichiometry and configurations.

DETAILED DESCRIPTION OF THE INVENTION

1. Overview

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The invention described herein provides a polynucleotide comprising three functional sequences, for binding to a target polynucleotide sequence (*e.g.*, the DNA-targeting sequence); for binding to either a wildtype (wt) Cas9 protein, or a modified Cas9 protein (*e.g.*, Cas9 nickase or dCas9) with reduced or deficient nuclease activity (*e.g.*, Cas9-binding sequence); and for binding to one or more PUF domain(s), each fused to a functional or effector domain. The polynucleotide of the invention, together with the wt or modified Cas9 protein and the one or more PUF domain fusion proteins, may form a 3-component complex (the subject 3-component CRISPR/Cas complex / system) at a specific target DNA sequence to effect one or more biological effects at the specific target DNA sequence.

The invention also provides a vector encoding such a polynucleotide, and a complex formed by the polynucleotide, the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), and at least one of the PUF domain fusion proteins. The invention further provides host cells comprising the vector or the polynucleotide.

The subject 3-component CRISPR/Cas complex / system can bring about a variety of biological functions at the target DNA sequence, including but are not limited to: enhanced homologous recombination to increase efficiency of knock-in, simultaneous transcription activation and/or repression at multiple genomic loci; detection of specific sequences at genomic loci by fluorescent imaging or other detectable signal; and affecting cell fate determination, cell differentiation, metabolic flux, or a biologically or biochemically determinable outcome, *etc*.

The invention further provides kits and reagents for carrying out the methods of the invention.

Thus in one aspect, the invention provides a polynucleotide comprising: (1) a DNA-targeting sequence that is complementary to a target polynucleotide sequence; (2) a Cas9-binding sequence; and, (3) one or more copies of a PUF domain-Binding Sequence (PBS), wherein each of the one or more copies of the PBS binds to the same or a different PUF domain; wherein a Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) is capable of forming a complex with the polynucleotide by binding to the Cas9-binding sequence. In certain embodiments, the dCas9 protein has reduced nuclease activity, or lacks nuclease activity (*e.g.*, is nuclease-deficient), but retains DNA-binding ability when complexed with the subject polynucleotide. In certain embodiments, (1) - (3) are arranged from 5' to 3', in that order. In other embodiments,

one or more of the PBS may be 5' to the DNA-targeting sequence, and/or 5' to the Cas9-binding sequence.

The target polynucleotide sequence can be any DNA sequence. In certain embodiments, the target polynucleotide sequence comprises, or is adjacent to, one or more transcription regulatory element(s). In certain embodiments, the transcription regulatory element(s) comprises one or more of: a core promoter, a proximal promoter element, an enhancer, a silencer, an insulator, and a locus control region. In another embodiment, the target polynucleotide sequence comprises, or is adjacent to, a centromere sequence, a telomere sequence, or a repetitive genomic sequence. The telomere sequence may be characterized by having 5-15 kb tracks of TTAGGG repeats. In yet another embodiment, the target polynucleotide sequence comprises, or is adjacent to, a genomic marker sequence or any genomic locus of interest.

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In certain embodiments, the target polynucleotide sequence is immediately 3' to a PAM (protospacer adjacent motif) sequence of the complementary strand. For example, in certain embodiments, the PAM sequence of the complementary strand is 5'-CCN-3', wherein N is any DNA nucleotide.

In other embodiments, the PAM sequence of the complementary strand matches the specific Cas9 protein or homologs or orthologs to be used.

As is known in the art, for Cas9 to successfully bind to DNA, the target sequence in the genomic DNA must be complementary to the guide RNA sequence and must be immediately followed by the correct protospacer adjacent motif or PAM sequence. The PAM sequence is present in the DNA target sequence but not in the guide RNA sequence. Any DNA sequence with the correct target sequence followed by the PAM sequence will be bound by Cas9.

The PAM sequence varies by the species of the bacteria from which the Cas9 was derived. The most widely used Type II CRISPR system is derived from *S. pyogenes* and the PAM sequence is 5'-NGG-3' located on the immediate 3' end of the guide RNA recognition sequence (or 5'-CCN-3' on the complementary strand). The PAM sequences of other Type II CRISPR systems from different bacterial species are listed in the Table below.

Streptococcus pyogenes (SP)	NGG
Neisseria meningitidis (NM)	NNNGATT
Streptococcus thermophilus (ST)	NNAGAA
Treponema denticola (TD)	NAAAAC

In certain embodiments, the DNA-targeting sequence base-pairs with the target polynucleotide sequence when the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) is complexed with the polynucleotide.

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It should be noted that the DNA-targeting sequence may or may not be 100% complementary to the target polynucleotide sequence. In certain embodiments, the DNA-targeting sequence is complementary to the target polynucleotide sequence over about 8-25 nucleotides (nts), about 12-22 nucleotides, about 14-20 nts, about 16-20 nts, about 18-20 nts, or about 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, or 25 nts. In certain embodiments, the complementary region comprises a continuous stretch of about 12-22 nts, preferably at the 3' end of the DNA-targeting sequence. In certain embodiments, the 5' end of the DNA-targeting sequence has up to 8 nucleotide mismatches with the target polynucleotide sequence. In certain embodiments, the DNA-binding sequence is about 50, 55, 60, 65, 70, 75, 80, 85, 90, 95, or 100% complementary to the target polynucleotide sequence.

In a related embodiment, there is no more than 15-nucleotide match at the 3' end of the DNA-targeting sequence compared to the complementary target polynucleotide sequence, and the Cas9 protein in the complex is a wt Cas9 protein which, under the circumstance, binds but does not cut a target DNA.

In certain embodiments, the DNA-binding sequence has a 5' end nucleotide G.

In certain embodiments, the polynucleotide further comprises a linker sequence linking the DNA-targeting sequence to the Cas9-binding sequence.

In certain embodiments, the Cas9-binding sequence forms a hairpin structure. In certain embodiments, the Cas9-binding sequence is about 30-100 nt, about 35-50 nt, about 37-47 nt, or about 42 nt in length.

An exemplary Cas9-binding sequence is GTTTTAGAGCTAGAAATAGCAAGTTAA AATAAGGCTA. Another exemplary Cas9-binding sequence is GTTTAAGAGCTATGC TG GAAACAGCATAGCAAGTTTAAATAAGGCTA.

The modified Cas9 protein (nickase or dCas9) may have reduced nuclease activity, or lacks nuclease activity at one or both endonuclease catalytic sites. In certain embodiments, the dCas9 protein lacks endonuclease activity due to point mutations at both endonuclease catalytic sites (RuvC and HNH) of wild type Cas9. For example, the point mutations may be D10A and H840A, respectively, in the *S. pyogenes* Cas9, or in the corresponding residues in species other than *S. pyogenes*. In certain embodiments, the modified Cas9 protein lacks endonuclease catalytic activity at one but not both sites of wt Cas9, and is able to create a nick on a dsDNA target (Cas9 nickase).

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In certain embodiments, each of the one or more copies of the PBS has about 8 nucleotides. One exemplary PBS may have a sequence of 5'-UGUAUGUA-3', which can be bound by the PUF domain PUF(3-2). Another exemplary PBS may have a sequence of 5'-UUGAUAUA-3', which can be bound by the PUF domain PUF(6-2/7-2). Additional PBS and the corresponding PUF domains are described below.

The polynucleotide of the invention may have more than one copies of the PBS. In certain embodiments, the polynucleotide comprises 1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 35, 40, 45, 46, 47, 48, 49, or 50 copies of PBS, such as 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, or 15 copies of PBS. In certain embodiments, the range of the PBS copy number is L to H, wherein L is any one of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 20, 25, 30, 35, or 40, and wherein H is any one of 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 80, 90, or 100, so long as H is greater than L. Each PBS may be the same or different.

In certain embodiments, the polynucleotide comprises about 5-15 copies of PBS, or about 5-14 copies, about 5-13 copies, about 5-12 copies, about 5-11 copies, about 5-10 copies, or about 5-9 copies of PBS.

In certain embodiments, the amount of the sgRNA-PBS and/or the amount of the PUF fusions transfected or expressed is adjusted to maximize PBS/PUF binding. For example, this can be achieved by increasing the expression of PUF-activator by a stronger promoter or using an inducible promoter, such as a Dox-inducible promoter.

In certain embodiments, the spacing between PBS sites and/or spacer sequences are optimized to improve system efficiency. For example, spacing optimization can be subject to particular PUF fusions, and can be different between PUF fusions that work as individual proteins and those PUF fusions that may need to be positioned close enough to function (e.g., protein complexes).

Another aspect of the invention provides a vector encoding any one of the subject polynucleotide. In certain embodiments, transcription of the polynucleotide is under the control of a constitutive promoter, or an inducible promoter. In certain embodiments, the vector is active in a cell from a mammal (a human; a non-human primate; a non-human mammal; a rodent such as a mouse, a rat, a hamster, a Guinea pig; a livestock mammal such as a pig, a sheep, a goat, a horse, a camel, cattle; or a pet mammal such as a cat or a dog); a bird, a fish, an insect, a worm, a yeast, or a bacterium.

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In certain embodiments, the vector is a plasmid, a viral vector (such as adenoviral, retroviral, or lentiviral vector, or AAV vector), or a transposon (such as piggyBac transposon). The vector can be transiently transfected into a host cell, or be integrated into a host genome by infection or transposition.

A related aspect of the invention provides a plurality or a library of any one of the vectors of the invention, wherein two of the vectors differ in the encoded polynucleotides in their respective DNA-targeting sequences, Cas9-binding sequences, and/or the copy number, identity (sequence, binding specificity, *etc.*), or relative order of the PBS.

Another aspect of the invention provides a complex comprising any one of the polynucleotide of the invention, and the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein). In certain embodiments, the complex comprises any one of the polynucleotide of the invention, and the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein). In certain embodiments, the complex does not comprise the wt Cas9 protein. In certain embodiments, the complex comprises the wt Cas9.

In certain embodiments, the complex may further comprise one or more PUF domain or fusion thereof bound to the one or more PBS(s). In certain embodiments, each of the PUF domain is fused to an effector domain. Each effector domain can be independently (but is not limited to): a transcription repressor, a transcription activator, a fluorescent protein, an enzyme, or a chromatin remodeling protein (HDAC/HAT). In certain embodiments, at least two of the PUF domains are fused to different effector domains.

In certain embodiments, the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), the PUF domain, and/or the effector domain further comprises a nuclear localization signal (NLS).

In certain embodiments, the complex is bound to the target polynucleotide sequence through the DNA-targeting sequence of the polynucleotide.

Another aspect of the invention provides a host cell comprising any one of the subject vector, or the plurality of vectors.

In certain embodiments, the host cell further comprises a second vector encoding the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein). In certain embodiments, the second vector further encodes an effector domain fused to the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein). The expression of the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) can be under the control of a constitutive promoter or an inducible promoter.

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In certain embodiments, the host cell may further comprise a third vector encoding the one or more PUF domains, each fused to an effector domain. The expression of the one or more PUF domains can be independently under the control of a constitutive promoter or an inducible promoter.

The effector domain can have any of many functions or biological effects. Merely to illustrate, the effector domain can be a protein involved in homologous recombination, a transcription repressor, a transcription activator, a fluorescent protein, an enzyme, or a chromatin remodeling protein (HDAC/HAT), *etc*.

In certain embodiments, the second vector may further encode a nuclear localization signal (NLS) fused to the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) or the effector domain, and/or the third vector may further encode a nuclear localization signal (NLS) fused to the PUF domain or the effector domain.

In certain embodiments, sequences that can be encoded by different vectors may be on the same vector. For example, in certain embodiments, the second vector may be the same as the vector, and/or the third vector may be the same as the vector or the second vector.

The host cell may be in a live animal, or may be a cultured cell.

In certain embodiments, the host cell may constitutively or inducibly express one or more components of the subject 3-component system (e.g., dCas9, PUF fusions).

Yet another aspect of the invention provides a method of assembling the complex of the invention at the target polynucleotide sequence, the method comprising contacting or bringing to the vicinity of the target polynucleotide sequence: (1) any one of the subject polynucleotide, or any one of the subject vector, or the plurality of vectors; (2) the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), or any one of the subject second vector encoding the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein); and, (3) one or more of the PUF domains, each fused to an effector domain, or any one of the third vector encoding the PUF domain fusions.

In certain embodiments, the complex is assembled inside a cell, the target polynucleotide sequence is a part of the genomic DNA of the cell, and wherein the subject vector, second

vector, and third vector are introduced into the cell.

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In certain embodiments, the target polynucleotide sequence is at or near a genomic locus rich in heterochromatin, and wherein the effector domain is a detectable marker (*e.g.*, a fluorescent protein). In another embodiment, the target polynucleotide sequence is at or near a transcription regulatory element of a target gene, and wherein the effector domain is a transcription modulator (*e.g.*, activator, suppressor). The transcription of the target gene, for example, may affect cell fate determination, cell differentiation, metabolic flux, or a biologically or biochemically determinable outcome.

A related aspect of the invention provides a method of modulating transcription of a plurality of target genes in a cell, the method comprising: introducing into the cell the subject plurality of the vectors, a coding sequence for a dCas9 protein, and a coding sequence for one or more PUF domain fusions, wherein each of the target genes comprises a target polynucleotide sequence that permits (1) the assembly, at the target polynucleotide sequence, of a tripartite complex of a polynucleotide encoded by one of the plurality of the vector, the dCas9 protein, and a PUF domain fusion; and (2) transcription modulation of the target gene comprising the target polynucleotide sequence.

In a related aspect, the invention also provides a method of epigenetic modulation (*e.g.*, modulating the epigenetic states of chromatin not directly related to transcriptional activity), at a plurality of target genes in a cell, the method comprising: introducing into the cell the subject plurality of the vectors, a coding sequence for a wt Cas9 protein or Cas9 nickase, and a coding sequence for one or more PUF domain fusions, wherein each of the target genes comprises a target polynucleotide sequence that permits (1) the assembly, at the target polynucleotide sequence, of a tripartite complex of a polynucleotide encoded by one of the plurality of the vector, the wt Cas9 protein or the Cas9 nickase, and a PUF domain fusion; and (2) epigenetic modulation of the target gene comprising the target polynucleotide sequence. The method can be useful, for example, to change epigenetic state (*e.g.*, opening up the chromatin) at the same time to gain access / stability of Cas9 binding to closed chromatin sites (*e.g.*, to increase cut and genome editing at those sites).

In certain embodiments, the transcription of at least one target gene is enhanced / stimulated, while the transcription of at least another target gene is inhibited.

The invention further provides a kit comprising: (1) a subject polynucleotide, or a vector encoding the same; (2) a second vector encoding the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein); and (3) a third vector encoding one or more PUF domain(s), each fused to an effector

domain. The kit may further comprise transformation, transfection, or infection reagents to facilitate the introduction of the vectors into a cell.

With the invention generally described above, various features of the invention will be further elaborated below. It should be understood that features of the invention, even when described in the context of separate embodiments, or even separate embodiments under different aspects of the invention, may be provided in combination in a single embodiment. Conversely, various features of the invention described in the context of a single embodiment, may also be provided separately or in any suitable subcombination. All combinations of the embodiments pertaining to the invention are specifically embraced by the present invention and are disclosed herein just as if each and every combination was individually and explicitly disclosed. In addition, all sub-combinations of the various embodiments and elements thereof are also specifically embraced by the present invention and are disclosed herein just as if each and every such sub-combination was individually and explicitly disclosed herein.

15 2. The Polynucleotide of the Invention

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The polynucleotide of the invention comprises three sequence segments: i) a first segment comprising a nucleotide sequence that is complementary to a target sequence; ii) a second segment that interacts with a Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein with reduced nuclease activity or lacks nuclease activity) (*e.g.*, the Cas9-binding sequence); and iii) one or more copies of a PUF domain-Binding Sequence (PBS).

In certain embodiments, the target sequence is an RNA. In certain embodiments, the target sequence is a DNA. In the description herein, the first segment is generally referred to as the "DNA-targeting sequence" when the target sequence is a DNA (such as a genomic DNA). In related embodiments in which the target sequence is an RNA, the description herein below applies generally as well except that the reference to "DNA-targeting sequence" is replaced with "RNA-targeting sequence," in order to avoid redundancy. That is, the first segment comprises a nucleotide sequence complementary to the target polynucleotide sequence (DNA or RNA).

In certain embodiments, the three segments i) - iii) are arranged, in that order, from 5' to 3'.

In certain embodiments, the polynucleotide of the invention can be a single RNA molecule (single RNA polynucleotide), which may include a "single-guide RNA," or "sgRNA." In another embodiment, the polynucleotide of the invention can comprise two RNA molecules (e.g., joined together via hybridization at the Cas9-binding sequence, see below). Thus the

subject polynucleotide is inclusive, referring both to two-molecule polynucleotide and to single-molecule polynucleotide (*e.g.*, sgRNAs).

a. DNA-Targeting Sequence

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The DNA-targeting sequence is functionally similar or equivalent to the crRNA or guide RNA or gRNA of the CRISPR/Cas complex / system. However, in the context of the instant invention, the DNA-targeting sequence may not originate from any particular crRNA or gRNA, but can be arbitrarily designed based on the sequence of the target polynucleotide sequence.

The DNA-targeting sequence comprises a nucleotide sequence that is complementary to a specific sequence within a target DNA (or the complementary strand of the target DNA). In other words, the DNA-targeting sequence interacts with a target polynucleotide sequence of the target DNA in a sequence-specific manner via hybridization (*i.e.*, base pairing). As such, the nucleotide sequence of the DNA-targeting sequence may vary, and it determines the location within the target DNA that the subject polynucleotide and the target DNA will interact. The DNA-targeting sequence can be modified or designed (*e.g.*, by genetic engineering) to hybridize to any desired sequence within the target DNA. In certain embodiments, the target polynucleotide sequence is immediately 3' to a PAM (protospacer adjacent motif) sequence of the complementary strand, which can be 5'-CCN-3', wherein N is any DNA nucleotide. That is, in this embodiment, the complementary strand of the target polynucleotide sequence is immediately 5' to a PAM sequence that is 5'-NGG-3', wherein N is any DNA nucleotide. In related embodiments, the PAM sequence of the complementary strand matches the wt or dCas9. See above for the PAM sequences from species other than *S. pyogenes*.

The DNA-targeting sequence can have a length of from about 12 nucleotides to about 100 nucleotides. For example, the DNA-targeting sequence can have a length of from about 12 nucleotides (nt) to about 80 nt, from about 12 nt to about 50 nt, from about 12 nt to about 40 nt, from about 12 nt to about 30 nt, from about 12 nt to about 25 nt, from about 12 nt to about 20 nt, or from about 12 nt to about 20 nt, from about 19 nt to about 20 nt, from about 19 nt to about 30 nt, from about 19 nt to about 35 nt, from about 19 nt to about 40 nt, from about 19 nt to about 45 nt, from about 19 nt to about 50 nt, from about 19 nt to about 60 nt, from about 19 nt to about 70 nt, from about 19 nt to about 25 nt, from about 20 nt to about 35 nt, from about 20 nt to about 30 nt, from about 20 nt to about 35 nt, from about 20 nt to about 30 nt, from about 20 nt to about 35 nt, from about 20 nt to about 45 nt, from about 20 nt to about 30 nt, from about 20 nt to about 30 nt, from about 20 nt to about 30 nt, from about 20 nt to about 50 nt from about 20 nt to about 50 nt from about 20 nt from

about 80 nt, from about 20 nt to about 90 nt, or from about 20 nt to about 100 nt.

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The nucleotide sequence of the DNA-targeting sequence that is complementary to a target polynucleotide sequence of the target DNA can have a length of at least about 12 nt. For example, the DNA-targeting sequence that is complementary to a target polynucleotide sequence of the target DNA can have a length at least about 12 nt, at least about 15 nt, at least about 18 nt, at least about 19 nt, at least about 20 nt, at least about 25 nt, at least about 30 nt, at least about 35 nt or at least about 40 nt. For example, the DNA-targeting sequence that is complementary to a target polynucleotide sequence of a target DNA can have a length of from about 12 nucleotides (nt) to about 80 nt, from about 12 nt to about 50 nt, from about 12 nt to about 45 nt, from about 12 nt to about 40 nt, from about 12 nt to about 35 nt, from about 12 nt to about 30 nt, from about 12 nt to about 25 nt, from about 12 nt to about 20 nt, from about 12 nt to about 19 nt, from about 19 nt to about 20 nt, from about 19 nt to about 25 nt, from about 19 nt to about 30 nt, from about 19 nt to about 35 nt, from about 19 nt to about 40 nt, from about 19 nt to about 45 nt, from about 19 nt to about 50 nt, from about 19 nt to about 60 nt, from about 20 nt to about 25 nt, from about 20 nt to about 30 nt, from about 20 nt to about 35 nt, from about 20 nt to about 40 nt, from about 20 nt to about 45 nt, from about 20 nt to about 50 nt, or from about 20 nt to about 60 nt. The nucleotide sequence of the DNA-targeting sequence that is complementary to the target polynucleotide sequence of the target DNA can have a length of at least about 12 nt.

In some cases, the DNA-targeting sequence that is complementary to a target polynucleotide sequence of the target DNA is 20 nucleotides in length. In some cases, the DNA-targeting sequence that is complementary to a target polynucleotide sequence of the target DNA is 19 nucleotides in length.

The percent complementarity between the DNA-targeting sequence and the target polynucleotide sequence of the target DNA can be at least 50% (*e.g.*, at least 55%, at least 60%, at least 65%, at least 70%, at least 75%, at least 80%, at least 85%, at least 90%, at least 95%, at least 97%, at least 98%, at least 99%, or 100%). In some cases, the percent complementarity between the DNA-targeting sequence and the target polynucleotide sequence is 100% over the seven or eight contiguous 5'-most nucleotides of the target polynucleotide sequence. In some cases, the percent complementarity between the DNA-targeting sequence and the target polynucleotide sequence is at least 60% over about 20 contiguous nucleotides. In some cases, the percent complementarity between the DNA-targeting sequence and the target polynucleotide sequence is 100% over the 7, 8, 9, 10, 11, 12, 13, or 14 contiguous 5'-most nucleotides of the target polynucleotide sequence (*i.e.*, the 7, 8, 9, 10, 11, 12, 13, or 14 contiguous 3'-most

nucleotides of the DNA-targeting sequence), and as low as 0% over the remainder. In such a case, the DNA-targeting sequence can be considered to be 7, 8, 9, 10, 11, 12, 13, or 14 nucleotides in length, respectively.

b. Cas9-Binding Sequence

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The protein-binding segment or protein-binding sequence of the subject polynucleotide binds to a wt Cas9, or a modified dCas9 protein (*e.g.*, nickase or dCas9) with reduced endonuclease activity, or lacks endonuclease activity. For simplicity, the protein-binding sequence of the subject polynucleotide, which may bind to wt and/or modified Cas9 proteins, may simply be referred to as "Cas9-binding sequence" herein. However, it should be understood that when the Cas9-binding sequence of the invention binds to a dCas9, it is not prevented from binding to a wt Cas9 or a Cas9 nickase. In certain embodiments, the Cas9-binding sequence of the invention binds to dCas9 as well as wt Cas9 and/or Cas9 nickase.

The Cas9-binding sequence interacts with or bind to a Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), and together they bind to the target polynucleotide sequence recognized by the DNA-targeting sequence. The Cas9-binding sequence comprises two complementary stretches of nucleotides that hybridize to one another to form a double stranded RNA duplex (a dsRNA duplex). These two complementary stretches of nucleotides may be covalently linked by intervening nucleotides known as linkers or linker nucleotides (*e.g.*, in the case of a single-molecule polynucleotide), and hybridize to form the double stranded RNA duplex (dsRNA duplex, or "Cas9-binding hairpin") of the Cas9-binding sequence, thus resulting in a stem-loop structure. Alternatively, in some embodiment, the two complementary stretches of nucleotides may not be covalently linked, but instead are held together by hybridization between complementary sequences (*e.g.*, in the case of a two-molecule polynucleotide of the invention).

The Cas9-binding sequence can have a length of from about 10 nucleotides to about 100 nucleotides, *e.g.*, from about 10 nucleotides (nt) to about 20 nt, from about 20 nt to about 30 nt, from about 30 nt to about 40 nt, from about 50 nt, from about 50 nt to about 60 nt, from about 60 nt to about 70 nt, from about 70 nt to about 80 nt, from about 80 nt to about 90 nt, or from about 90 nt to about 100 nt. For example, the Cas9-binding sequence can have a length of from about 15 nucleotides (nt) to about 80 nt, from about 15 nt to about 50 nt, from about 15 nt to about 40 nt, from about 15 nt to about 30 nt, from about 37 nt to about 47 nt (*e.g.*, 42 nt), or from about 15 nt to about 25 nt.

The dsRNA duplex of the Cas9-binding sequence can have a length from about 6 base

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pairs (bp) to about 50 bp. For example, the dsRNA duplex of the Cas9-binding sequence can have a length from about 6 bp to about 40 bp, from about 6 bp to about 30 bp, from about 6 bp to about 25 bp, from about 6 bp to about 20 bp, from about 6 bp to about 15 bp, from about 8 bp to about 40 bp, from about 8 bp to about 30 bp, from about 8 bp to about 25 bp, from about 8 bp to about 20 bp or from about 8 bp to about 15 bp. For example, the dsRNA duplex of the Cas9binding sequence can have a length from about from about 8 bp to about 10 bp, from about 10 bp to about 15 bp, from about 15 bp to about 18 bp, from about 18 bp to about 20 bp, from about 20 bp to about 25 bp, from about 25 bp to about 30 bp, from about 30 bp to about 35 bp, from about 35 bp to about 40 bp, or from about 40 bp to about 50 bp. In some embodiments, the dsRNA duplex of the Cas9-binding sequence has a length of 36 base pairs. The percent complementarity between the nucleotide sequences that hybridize to form the dsRNA duplex of the Cas9-binding sequence can be at least about 60%. For example, the percent complementarity between the nucleotide sequences that hybridize to form the dsRNA duplex of the Cas9-binding sequence can be at least about 65%, at least about 70%, at least about 75%, at least about 80%, at least about 85%, at least about 90%, at least about 95%, at least about 98%, or at least about 99%. In some cases, the percent complementarity between the nucleotide sequences that hybridize to form the dsRNA duplex of the Cas9-binding sequence is 100%.

The linker can have a length of from about 3 nucleotides to about 100 nucleotides. For example, the linker can have a length of from about 3 nucleotides (nt) to about 90 nt, from about 3 nucleotides (nt) to about 70 nt, from about 3 nucleotides (nt) to about 50 nt, from about 3 nucleotides (nt) to about 40 nt, from about 3 nucleotides (nt) to about 30 nt, from about 3 nucleotides (nt) to about 20 nt or from about 3 nucleotides (nt) to about 10 nt. For example, the linker can have a length of from about 3 nt to about 5 nt, from about 5 nt to about 10 nt, from about 10 nt to about 15 nt, from about 15 nt to about 20 nt to about 25 nt, from about 25 nt to about 30 nt, from about 40 nt to about 50 nt, from about 50 nt to about 60 nt, from about 60 nt to about 70 nt, from about 70 nt to about 80 nt, from about 80 nt to about 90 nt, or from about 90 nt to about 100 nt. In some embodiments, the linker is 4 nt.

Non-limiting examples of nucleotide sequences that can be included in a suitable Cas9-binding sequence (*i.e.*, Cas9 handle) are set forth in SEQ ID NOs: 563-682 of WO 2013/176772 (see, for examples, FIGs. 8 and 9 of WO 2013/176772), incorporated herein by reference.

In some cases, a suitable Cas9-binding sequence comprises a nucleotide sequence that

differs by 1, 2, 3, 4, or 5 nucleotides from any one of the above-listed sequences.

c. PUF Domain-Binding Sequence (PBS)

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The subject polynucleotide comprises one or more tandem sequences, each of which can be specifically recognized and bound by a specific PUF domain (*infra*). Since a PUF domain can be engineered to bind virtually any PBS based on the nucleotide-specific interaction between the individual PUF motifs of PUF domain and the single RNA nucleotide they recognize, the PBS sequences can be any designed sequence that bind their corresponding PUF domain.

In certain embodiments, a PBS of the invention has 8-mer. In other embodiments, a PBS of the invention has 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16 or more RNA nucleotides.

In certain embodiments, the PBS of the invention has the sequence 5'-UGUAUAUA-3', and binds the wt human Pumilio 1 PUF domain.

In certain embodiments, the PBS of the invention has the sequence 5'-UGUAUGUA-3', and binds the PUF domain PUF(3-2).

In certain embodiments, the PBS of the invention has the sequence 5'-UUGAUAUA-3', and binds the PUF domain PUF(6-2/7-2).

In certain embodiments, the PBS of the invention has the sequence 5'-UGGAUAUA-3', and binds the PUF domain PUF(6-2).

In certain embodiments, the PBS of the invention has the sequence 5'-UUUAUAUA-3', and binds the PUF domain PUF(7-2).

In certain embodiments, the PBS of the invention has the sequence 5'-UGUGUGUG-3', and binds the PUF domain PUF⁵³¹.

In certain embodiments, the PBS of the invention has the sequence 5'-UGUAUAUG-3', and binds the PUF domain PUF(1-1).

In certain embodiments, the PBS of the invention has the sequence 5'-UUUAUAUA-3' or 5'-UAUAUAUA-3', and binds the PUF domain PUF(7-1).

In certain embodiments, the PBS of the invention has the sequence 5'-UGUAUUUA-3', and binds the PUF domain PUF(3-1).

In certain embodiments, the PBS of the invention has the sequence 5'-UUUAUUUA-3', and binds the PUF domain PUF(7-2/3-1).

Applicant has created 65,536 8-mer PBS and their corresponding PUF domain sequences (see below) that can bind the specific PBS. Applicant has also created a python script to retrieve

any of the 65,536 individual PUF domain sequences that binds a given 8-mer PBS. For example, for the 8-mer UUGAUGUA, one possible PUF domain sequence can be:

GRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGCRFIQLKLERATPAERQLVFNEILQAAYQ
LMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIEKALEFIPSDQQNEMVR
ELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPYGCRVIQRILEHCLP
DQTLPILEELHQHTEQLVQDQYGSYVIEHVLEHGRPEDKSKIVAEIRGNVLVLSQHKFANNVVQ
KCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAEPGQRKIVMHKIRPH
IATLRKYTYGKHILAKLEKYYMKNGVDLG

In certain embodiments, one or more spacer region(s) separates two adjacent PBS sequences. The spacer regions may have a length of from about 3 nucleotides to about 100 nucleotides. For example, the spacer can have a length of from about 3 nucleotides (nt) to about 90 nt, from about 3 nucleotides (nt) to about 80 nt, from about 3 nucleotides (nt) to about 70 nt, from about 3 nucleotides (nt) to about 60 nt, from about 3 nucleotides (nt) to about 50 nt, from about 3 nucleotides (nt) to about 20 nt, from about 3 nucleotides (nt) to about 30 nt, from about 3 nucleotides (nt) to about 20 nt or from about 3 nucleotides (nt) to about 10 nt. For example, the spacer can have a length of from about 3 nt to about 5 nt, from about 5 nt to about 10 nt, from about 10 nt to about 15 nt, from about 15 nt to about 20 nt, from about 20 nt to about 25 nt, from about 25 nt to about 30 nt, from about 30 nt to about 35 nt, from about 35 nt to about 40 nt, from about 40 nt to about 50 nt, from about 50 nt to about 60 nt, from about 60 nt to about 70 nt, from about 70 nt to about 80 nt, from about 80 nt to about 90 nt, or from about 90 nt to about 100 nt. In some embodiments, the spacer is 4 nt.

d. Optional Other Sequences

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A stability control sequence (*e.g.*, transcriptional terminator segment) influences the stability of an RNA (*e.g.*, a subject polynucleotide). One example of a suitable stability control sequence is a transcriptional terminator segment (*i.e.*, a transcription termination sequence). A transcriptional terminator segment of a subject polynucleotide can have a total length of from about 10 nucleotides to about 100 nucleotides, *e.g.*, from about 10 nucleotides (nt) to about 20 nt, from about 20 nt to about 30 nt, from about 30 nt to about 40 nt, from about 40 nt to about 50 nt, from about 50 nt to about 60 nt, from about 60 nt to about 70 nt, from about 70 nt to about 80 nt, from about 80 nt to about 90 nt, or from about 90 nt to about 100 nt. For example, the transcriptional terminator segment can have a length of from about 15 nucleotides (nt) to about 80 nt, from about 15 nt to about 50 nt, from about 15 nt to about 15 nt to about 30 nt or from about 15 nt to about 25 nt.

In some cases, the transcription termination sequence is one that is functional in a eukaryotic cell. In some cases, the transcription termination sequence is one that is functional in a prokaryotic cell.

Non-limiting examples of nucleotide sequences that can be included in a stability control sequence (*e.g.*, transcriptional termination segment, or in any segment of the DNA-targeting RNA to provide for increased stability) include sequences set forth in SEQ ID NO: 683-696 of WO 2013/176772 (incorporated herein by reference), see, for example, SEQ ID NO: 795 of WO 2013/176772, a Rho-independent transcription termination site.

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The stability control sequence may be situated after the Cas9-binding sequence, for example, between the Cas9-binding sequence and the first PBS, between two adjacent PBS, or after the last PBS.

In some embodiments, the polynucleotide of the invention or parts thereof (*e.g.*, the DNA-targeting sequence, the Cas9-binding sequence, and/or the one or more of the PBS), or a polynucleotide encoding the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), or a polynucleotide encoding one of the PUF domain fusions (*infra*), may comprise a modification or sequence that provides for an additional desirable feature, *e.g.*, modified or regulated stability; subcellular targeting; tracking, *e.g.*, a fluorescent label; a binding site for a protein or protein complex; *etc.*).

Non-limiting examples include: a 5' cap (*e.g.*, a 7-methylguanylate cap (m⁷G)); a 3' polyadenylated tail (*i.e.*, a 3' poly(A) tail); a riboswitch sequence or an aptamer sequence (*e.g.*, to allow for regulated stability and/or regulated accessibility by proteins and protein complexes); a terminator sequence; a sequence that forms a dsRNA duplex (*i.e.*, a hairpin)); a modification or sequence that targets the RNA to a subcellular location (*e.g.*, nucleus, mitochondria, chloroplasts, and the like); a modification or sequence that provides for tracking (*e.g.*, direct conjugation to a fluorescent molecule, conjugation to a moiety that facilitates fluorescent detection, a sequence that allows for fluorescent detection, *etc.*); a modification or sequence that provides a binding site for proteins (*e.g.*, proteins that act on DNA, including transcriptional activators, transcriptional repressors, DNA methyltransferases, DNA demethylases, histone acetyltransferases, histone deacetylases, and the like); a modification or sequence that provides for increased, decreased, and/or controllable stability; and combinations thereof.

3. The Cas9 Protein (wt, nickase, or dCas9)

The Cas9 protein (e.g., wt, nickase, or dCas9 protein) of the invention comprises: i) an

RNA-binding portion that interacts with the Cas9-binding sequence of the subject polynucleotide, and ii) an activity portion that exhibits wt, reduced endonuclease (*e.g.*, endodeoxyribonuclease) activity, or lacks endonuclease (*e.g.*, endodeoxyribonuclease) activity, depending on the identity of the Cas9 protein.

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The Cas9-binding sequence of the polynucleotide and the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) can form a complex that binds to a specific target polynucleotide sequence, based on the sequence complementarity between the DNA-targeting sequence and the target polynucleotide sequence. The DNA-targeting sequence of the subject polynucleotide provides target specificity to the complex via its sequence complementarity to the target polynucleotide sequence of a target DNA. If the target polynucleotide sequence is at or adjacent to a transcription regulatory element or an epigenetic modification site of a target gene, the complex, together with transcription regulators or effectors that modulate epigenetic modification fused to PBS-binding PUF domain, can selectively modulate transcription or epigenetic modulation of the target gene.

In certain embodiments, the modified Cas9 protein has reduced or lacks endonuclease (*e.g.*, endodeoxyribonuclease) activity. For example, a modified Cas9 suitable for use in a method of the present invention may be a Cas9 nickase, or exhibits less than about 20%, less than about 15%, less than about 15%, less than about 15%, less than about 1%, or less than about 0.1%, of the endonuclease (*e.g.*, endodeoxyribonuclease) activity of a wild-type Cas9 polypeptide, *e.g.*, a wild-type Cas9 polypeptide comprising an amino acid sequence as depicted in FIG. 3 and SEQ ID NO: 8 of WO 2013/176772 (incorporated herein by reference). In some embodiments, the dCas9 has substantially no detectable endonuclease (*e.g.*, endodeoxyribonuclease) activity. In some embodiments when a dCas9 has reduced catalytic activity (*e.g.*, when a Cas9 protein has a D10, G12, G17, E762, H840, N854, N863, H982, H983, A984, D986, and/or a A987 mutation, *e.g.*, D10A, G12A, G17A, E762A, H840A, N854A, N863A, H982A, H983A, A984A, and/or D986A), the polypeptide can still bind to target DNA in a site-specific manner, because it is still guided to a target polynucleotide sequence by a DNA-targeting sequence of the subject polynucleotide, as long as it retains the ability to interact with the Cas9-binding sequence of the subject polynucleotide.

In some cases, a suitable Cas9 protein (e.g., wt, nickase, or dCas9 protein) comprises an amino acid sequence having at least about 75%, at least about 80%, at least about 85%, at least about 90%, at least about 95%, at least about 99% or 100% amino acid sequence identity to amino acids 7-166 or 731-1003 of the Cas9/Csnl amino acid sequence (of *Streptococcus*

pyogenes), as depicted in FIG. 3 and SEQ ID NO: 8 of WO 2013/176772 (incorporated by reference), or to the corresponding portions in any one of the amino acid sequences SEQ ID NOs: 1-256 and 795-1346 of WO 2013/176772 (incorporated by reference), preferably to the corresponding portions in any one of the amino acid sequences of the orthogonal Cas9 sequences from *S. pyogenes*, *N. meningitidis*, *S. thermophilus* and *T. denticola* (see, Esvelt *et al.*, *Nature Methods*, 10(11): 1116-1121, 2013, incorporated by reference).

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In some cases, the Cas9 nickase can cleave the complementary strand of the target DNA but has reduced ability to cleave the non-complementary strand of the target DNA. For example, the Cas9 nickase can have a mutation (amino acid substitution) that reduces the function of the RuvC domain. As a non-limiting example, in some cases, the Cas9 nickase is a D10A (aspartate to alanine) mutation of the amino acid sequence depicted in FIG. 3 of WO 2013/176772, or the corresponding mutation of any of the amino acid sequences set forth in SEQ ID NOs: 1-256 and 795-1346 of WO 2013/176772 (all such sequences incorporated by reference).

In some cases, the Cas9 nickase can cleave the non-complementary strand of the target DNA but has reduced ability to cleave the complementary strand of the target DNA. For example, the Cas9 nickase can have a mutation (amino acid substitution) that reduces the function of the HNH domain (RuvC/HNH/RuvC domain motifs). As a non-limiting example, in some cases, the Cas9 nickase is a H840A (histidine to alanine at amino acid position 840 of SEQ ID NO: 8 of WO 2013/176772, incorporated by reference) or the corresponding mutation of any of the amino acid sequences set forth in SEQ ID NOs: 1-256 and 795-1346 of WO 2013/176772 (all such sequences incorporated by reference).

In some cases, the dCas9 has a reduced ability to cleave both the complementary and the non-complementary strands of the target DNA. As a non-limiting example, in some cases, the dCas9 harbors both D10A and H840A mutations of the amino acid sequence depicted in FIG. 3 of WO 2013/176772 or the corresponding mutations of any of the amino acid sequences set forth in SEQ ID NOs: 1-256 and 795-1346 of WO 2013/176772 (all such sequences incorporated by reference).

Other residues can be mutated to achieve the same effect (*i.e.* inactivate one or the other nuclease portions). As non-limiting examples, residues D10, G12, G17, E762, H840, N854, N863, H982, H983, A984, D986, and/or A987 (or the corresponding mutations of any of the proteins set forth as SEQ ID NOs: 1-256 and 795-1346) can be altered (*i.e.*, substituted) (see FIGs. 3, 5, 11A, and Table 1 of WO 2013/176772 (all incorporated by reference) for more

information regarding the conservation of Cas9 amino acid residues). Also, mutations other than alanine substitutions are suitable.

In some cases, the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) is optionally a fusion polypeptide comprising: i) a Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein); and b) a covalently linked heterologous polypeptide (also referred to as a "fusion partner"), which can be the same or different from the fusion partner fused to the PUF domains (*infra*).

4. PUF Domain (and the optional Cas9) Fusion Proteins

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PUF proteins (named after *Drosophila* Pumilio and *C. elegans* fern-3 binding factor) are known to be involve in mediating mRNA stability and translation. These protein contain a unique RNA-binding domain known as the PUF domain. The RNA-binding PUF domain, such as that of the human Pumilio 1 protein (referred here also as PUM), contains 8 repeats (each repeat called a PUF motif or a PUF repeat) that bind consecutive bases in an anti-parallel fashion, with each repeat recognizing a single base - *i.e.*, PUF repeats R1 to R8 recognize nucleotides N8 to N1, respectively. For example, PUM is composed of eight tandem repeats, each repeat consisting of 34 amino acids that folds into tightly packed domains composed of alpha helices.

Each PUF repeat uses two conserved amino acids from the center of each repeat to specifically recognize the edge of one individual base within the RNA recognition sequence, and a third amino acid (Tyr, His or Arg) to stack between adjacent bases, causing a very specific binding between a PUF domain and an 8-mer RNA. For example, the code to recognize base U is the amino acid sequence "NYxxQ", whereas "(C/S)RxxQ" recognizes A and "SNxxE" recognizes G. These amino acids correspond to positions 12, 13, and 16 in the human Pumilio 1 PUF motif. The two recognition amino acid side chains at positions 12 and 16 in each PUF α - α - α repeat recognize the Watson-Crick edge of the corresponding base and largely determine the specificity of that repeat.

Therefore, the sequence specificity of the PUF domains can be altered precisely by changing the conserved amino acid (*e.g.*, by site-directed mutagenesis) involved in base recognition within the RNA recognition sequence. By changing two amino acids in each repeat, a PUF domain can be modified to bind almost any 8-nt RNA sequence. This unique binding mode makes PUF and its derivatives a programmable RNA-binding domain that can be used in the instant invention, as part of a PUF domain-fusion that brings any effector domain to a specific PBS on the subject polynucleotide.

As used herein, "PUF domain" refers to a wildtype or naturally existing PUF domain, as well as a PUF homologue domain that is based on / derived from a natural or existing PUF domain, such as the prototype human Pumilio 1 PUF domain. The PUF domain of the invention specifically binds to an RNA sequence (*e.g.*, an 8-mer RNA sequence), wherein the overall binding specificity between the PUF domain and the RNA sequence is defined by sequence specific binding between each PUF motif / PUF repeat within the PUF domain and the corresponding single RNA nucleotide.

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In certain embodiments, the PUF domain comprises or consists essentially of 8 PUF motifs, each specifically recognizes and binds to one RNA nucleotide (*e.g.*, A, U, G, or C).

Applicant has created 65,536 8-mer PBS and their corresponding PUF domain sequences (each about 350 amino acids long) that can bind the specific PBS. Applicant has also created a python script to retrieve any of the 65,536 individual PUF domain sequences that binds a given 8-mer PBS.

In certain embodiments, the PUF domain has more or less than 8 PUF motifs / repeats, e.g., the PUF domain comprises or consists essentially of 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, or more PUF repeats / motifs, each specifically recognizes and binds to one RNA nucleotide (e.g., A, U, G, or C), so long as the PUF domain binds the RNA of 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16 or more nucleotides. By increasing or decreasing the number of PUF motifs, the length of the recognized RNA will be correspondingly increased or decreased. Since each PUF motif recognizes one RNA base, decreasing the domain by one motif decreases the length of the RNA recognized by one base; while increasing the domain by one motif increases the length of the RNA recognized by one base. Any number of motifs may be present. Therefore, in such embodiments, the specificity of the PUF domain-fusions of the invention may be altered due to changes in PUF domain length. In certain embodiments, the additional PUF motifs are inserted between two of the original PUF motifs, e.g., before the 1st, between the 1st and the 2nd, the 2nd and the 3rd, the 3rd, and the 4th, the 4th and the 5th, the 5th and the 6th, the 6th and the 7th, the 7th and the 8th, or after the 8th. In certain embodiments, there are 1, 2, 3, 4, 5, 6, 7, 8, or more inserted PUF motifs between any of the insertion points above. For example, in certain embodiments, there are 1, 2, 3, 4, 5, 6, 7, 8, or more inserted PUF motifs between the 5th and the 6th original PUF motif. Filipovska et al. (Nature Chemical Biology doi: 10.1038/NChemBio.577, published online: 15 may 2011) have reported an engineered PUF domain with 16 PUF motifs, including 8 additional PUF motifs inserted between the 5^{th} and 6^{th} original PUF motifs.

In certain embodiments, the PUF domain comprises PUF motifs from different PUF domains from different proteins. For example, a PUF domain of the invention may be constructed with PUF motifs from the human Pumilio 1 protein and one or more other PUF motifs from one or more other PUF proteins, such as PuDp or FBF. The RNA binding pockets of PUF domains have natural concave curvatures. Since different PUF proteins may have different curvatures, different PUF motifs in a PUF domain may be used to alter the curvature of the PUF domain. Altering the curvature is another method for altering the specificity and/or binding affinity of the PUF domain since flatter curvatures may allow for the recognition of more RNA bases.

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Also included in the scope of the invention are functional variants of the subject PUF domains or fusions thereof. The term "functional variant" as used herein refers to a PUF domain having substantial or significant sequence identity or similarity to a parent PUF domain, which functional variant retains the biological activity of the PUF domain of which it is a variant - e.g., one that retains the ability to recognize target RNA to a similar extent, the same extent, or to a higher extent in terms of binding affinity, and/or with substantially the same or identical binding specificity, as the parent PUF domain. The functional variant PUF domain can, for instance, be at least about 30%, 50%, 75%, 80%, 90%, 98% or more identical in amino acid sequence to the parent PUF domain. The functional variant can, for example, comprise the amino acid sequence of the parent PUF domain with at least one conservative amino acid substitution, for example, conservative amino acid substitutions in the scaffold of the PUF domain (i.e., amino acids that do not interact with the RNA). Alternatively or additionally, the functional variants can comprise the amino acid sequence of the parent PUF domain with at least one non-conservative amino acid substitution. In this case, it is preferable for the non-conservative amino acid substitution to not interfere with or inhibit the biological activity of the functional variant. The non-conservative amino acid substitution may enhance the biological activity of the functional variant, such that the biological activity of the functional variant is increased as compared to the parent PUF domain, or may alter the stability of the PUF domain to a desired level (e.g., due to substitution of amino acids in the scaffold). The PUF domain can consist essentially of the specified amino acid sequence or sequences described herein, such that other components, e.g., other amino acids, do not materially change the biological activity of the functional variant.

In certain embodiments, the PUF domain is a Pumilio homology domain (PU-HUD). In a particular embodiment, the PU-HUD is a human Pumilio 1 domain. The sequence of the human PUM is known in the art and is reproduced below:

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Gly Arg Ser Arg Leu Leu Glu Asp Phe Arg Asn Asn Arg Tyr Pro Asn
    Leu Gln Leu Arg Glu Ile Ala Gly His Ile Met Glu Phe Ser Gln Asp
    Gln His Gly Ser Arg Phe Ile Gln Leu Lys Leu Glu Arg Ala Thr Pro
    Ala Glu Arg Gln Leu Val Phe Asn Glu Ile Leu Gln Ala Ala Tyr Gln
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    Leu Met Val Asp Val Phe Gly Asn Tyr Val Ile Gln Lys Phe Phe Glu
    Phe Gly Ser Leu Glu Gln Lys Leu Ala Leu Ala Glu Arg Ile Arg Gly
    His Val Leu Ser Leu Ala Leu Gln Met Tyr Gly Cys Arg Val Ile Gln
    Lys Ala Leu Glu Phe Ile Pro Ser Asp Gln Gln Asn Glu Met Val Arg
    Glu Leu Asp Gly His Val Leu Lys Cys Val Lys Asp Gln Asn Gly Asn
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    His Val Val Gln Lys Cys Ile Glu Cys Val Gln Pro Gln Ser Leu Gln
    Phe Ile Ile Asp Ala Phe Lys Gly Gln Val Phe Ala Leu Ser Thr His
    Pro Tyr Gly Cys Arg Val Ile Gln Arg Ile Leu Glu His Cys Leu Pro
    Asp Gln Thr Leu Pro Ile Leu Glu Glu Leu His Gln His Thr Glu Gln
    Leu Val Gln Asp Gln Tyr Gly Asn Tyr Val Ile Gln His Val Leu Glu
    His Gly Arg Pro Glu Asp Lys Ser Lys Ile Val Ala Glu Ile Arg Gly
15
    Asn Val Leu Val Leu Ser Gln His Lys Phe Ala Ser Asn Val Val Glu
    Lys Cys Val Thr His Ala Ser Arg Thr Glu Arg Ala Val Leu Ile Asp
    Glu Val Cys Thr Met Asn Asp Gly Pro His Ser Ala Leu Tyr Thr Met
    Met Lys Asp Gln Tyr Ala Asn Tyr Val Val Gln Lys Met Ile Asp Val Ala Glu Pro Gly Gln Arg Lys Ile Val Met His Lys Ile Arg Pro His
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    Ile Ala Thr Leu Arg Lys Tyr Thr Tyr Gly Lys His Ile Leu Ala Lys
    Leu Glu Lys Tyr Tyr Met Lys Asn Gly Val Asp Leu Gly
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The wt human PUM specifically binds the Nanos Response Element (NRE) RNA, bearing a core 8-nt sequence 5'-UGUAUAUA-3'.

In certain embodiments, the PUF domain of the invention is any PUF protein family member with a Pum-HD domain. Non-limiting examples of a PUF family member include FBF in *C. elegans*, Ds pum in *Drosophila*, and PUF proteins in plants such as *Arabidopsis* and rice. A phylogenetic tree of the PUM-HDs of *Arabidopsis*, rice and other plant and non-plant species is provided in Tam *et al.* ("The Puf family of RNA-binding proteins in plants: phylogeny, structural modeling, activity and subcellular localization." *BMC Plant Biol.* 10:44, 2010, the entire contents of which are incorporated by reference herein).

PUF family members are highly conserved from yeast to human, and all members of the family bind to RNA in a sequence specific manner with a predictable code. The accession number for the domain is PS50302 in the Prosite database (Swiss Institute of Bioinformatics) and a sequence alignment of some of the members of this family is shown in FIG. 5 & 6 of WO 2011-160052 A2 (ClustalW multiple sequence alignment of human, mouse, rat Pumilio 1 (hpum1, Mpum1, Ratpum1) and human and mouse Pumilio 2 (hpum2, Mpum2), respectively.

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The *Drosophila* Pumilio (PumDr) is very different in length from other mammalian Pumilio 1 homologues, thus only the C-terminal PUF HUD domain is shown in the sequence alignment with human PUM1 and PUM2 in FIG. 6 of WO 2011/160052A2. The N-terminal part of human and fly Pum proteins shows weak homology (40% similarity) and differs

significantly in size and protein sequence. The C-terminal part shows a very high degree of homology and evolutionary conservation (78% identity, 86% similarity for PUM1 and 79% identity, 88% similarity for PUM2), with highly conserved protein sequence and structure of the Pum RNA-binding domain. In all three proteins PUM-HD is composed of the N-terminal conserved part of 20 amino acids, eight Pum repeats of 36 amino acids each, and the C-terminal conserved region. In human Pumilio proteins, the C-conserved part is 44 amino acids long, whereas *Drosophila* protein has an insert of additional 85 amino acids in the C-conserved region. The nucleotide and amino acid sequences can be found in the DDBJ/EMBL/GENBANK® databases under accession nos. AF315592 (PUM1) and AF315591 (PUM2) (Spassov & Jurecic, "Cloning and comparative sequence analysis of PUM1 and PUM2 genes, human members of the Pumilio family of RNA-binding proteins," *Gene*, 299:195-204, October 2002, the entire contents of each of which (publication and sequences) are incorporated by reference herein).

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In addition, all aligned sequences, *i.e.*, SEQ ID NOs:55-60 of WO 2011/160052A2, are incorporated herein by reference.

In some embodiments, the PUF domain of the invention can be made up of eight 36 mers, in which 33 of the amino acids are conserved and the 34th, 35th and 36th amino acids can vary, imparting specificity for a particular base in an RNA sequence. In particular embodiments, the RNA binding domain is about 300 (*e.g.*, 310, 309, 308, 307, 306, 305, 304, 303, 302, 301, 300, 299, 298, 297, 296, 295, 294, 293, 292, 291, 290, *etc.*) amino acids in length. In some embodiments, the PUF domain of this invention is designed to bind to a specific RNA sequence of about 8 nucleotides (*e.g.*, 8-16 contiguous RNA bases). In particular embodiments, the fifth nucleotide of the 8-nt sequence is a U or C, while the other 7 nucleotides can vary.

In some embodiments, the PUF domain is modified from a wt PUF domain to bind an RNA sequence that is different from the RNA sequence bound by the unmodified (*i.e.*, wild type) RNA binding PUF domain. The RNA sequence can be about an 8mer (*e.g.*, an 8mer, 9 mer, 10mer, 11mer, 12mer, 13mer, 14mer, 15mer, 16mer, *etc.*). The ability to introduce modifications into the amino acid sequence of the RNA binding domain to alter its specificity for a target RNA sequence is based on the known interactions of bases with the different amino acid side chains of the RNA binding domain (*e.g.*, PUF proteins). The RNA recognition code of the PUF domain is shown below, which can be generally written as:

SerXXXGlu for G (guanine), such as SNxxE;

CysXXXGln, such as CysArgXXGln or SerArgXXGln (*i.e.*, (C/S)RxxQ) for A (adenine);

AsnXXXGln for U (uracil), such as NYxxQ, and,

5 SnXXXArg for C (cytosine), such as SerTyrXXArg.

where X is any amino acid, and Sn represents a small or nucleophilic residue such as Gly, Ala, Ser, Thr, or Cys.

Based on the guidelines above, at least one PUF domain can be constructed based on any given 8-mer sequences. Specifically, a PUF domain binding to an 8-mer RNA sequence of 5'- $N_1N_2N_3N_4N_5N_6N_7N_8$ -3' can have the following sequence formula, in which R1-R8 each represents a PUF motif peptide sequence listed in the tables below, depending on the specific identity of the ribonucleotide (*i.e.*, A, U, C, or G) at any of the N_1 - N_8 locations. Note that R1 binds N_8 , R2 binds N_7 , etc.

GlyArgSerArgLeuLeuGluAspPheArgAsnAsnArgTyrProAsnLeuGlnLeuArgGluIleAlaG
lyHisIleMetGluPheSerGlnAsp[R1]ThrProAlaGluArgGlnLeuValPheAsnGluIleLeuG
lnAlaAlaTyrGlnLeuMetValAsp[R2]SerLeuGluGlnLysLeuAlaLeuAlaGluArgIleArgG
lyHisValLeuSerLeuAlaLeuGln[R3]ProSerAspGlnGlnAsnGluMetValArgGluLeuAspG
lyHisValLeuLysCysValLysAsp[R4]GlnProGlnSerLeuGlnPheIleIleAspAlaPheLysG
lyGlnValPheAlaLeuSerThrHis[R5]LeuProAspGlnThrLeuProIleLeuGluGluLeuHisG
lnHisThrGluGlnLeuValGlnAsp[R6]ArgProGluAspLysSerLysIleValAlaGluIleArgG
lyAsnValLeuValLeuSerGlnHis[R7]SerArgThrGluArgAlaValLeuIleAspGluValCysT
hrMetAsnAspGlyProHisSerAlaLeuTyrThrMetMetLysAsp[R8]GluProGlyGlnArgLysI
leValMetHisLysIleArgProHisIleAlaThrLeuArgLysTyrThrTyrGlyLysHisIleLeuAl
aLysLeuGluLysTyrTyrMetLysAsnGlyValAspLeuGly

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N ₈ nucleotide	R1 peptide sequence(s)
A	GlnHisGly CysArg PheIle Gln LeuLysLeuGluArgAla
	GlnHisGly SerArg PheIle Gln LeuLysLeuGluArgAla
С	GlnHisGly Ser ArgPheIle Arg LeuLysLeuGluArgAla
	GlnHisGly Gly ArgPheIle Arg LeuLysLeuGluArgAla
	GlnHisGly Ala ArgPheIle Arg LeuLysLeuGluArgAla
	GlnHisGly Thr ArgPheIle Arg LeuLysLeuGluArgAla
	GlnHisGly Cys ArgPheIle Arg LeuLysLeuGluArgAla
G	GlnHisGly Ser ArgPheIle Glu LeuLysLeuGluArgAla
U	GlnHisGly Asn ArgPheIle Gln LeuLysLeuGluArgAla

N ₇ nucleotide	R2 peptide sequence(s)
A	ValPheGly CysArg ValIle Gln LysPhePheGluPheGly
	ValPheGly SerArg ValIle Gln LysPhePheGluPheGly
	ValPheGly Cys TyrValIle Gln LysPhePheGluPheGly
	ValPheGly Ser TyrValIle Gln LysPhePheGluPheGly
С	ValPheGly Ser TyrValIle Arg LysPhePheGluPheGly
	ValPheGly Gly TyrValIle Arg LysPhePheGluPheGly
	ValPheGly Ala TyrValIle Arg LysPhePheGluPheGly
	ValPheGly Thr TyrValIle Arg LysPhePheGluPheGly

N ₇ nucleotide	R2 peptide sequence(s)
	ValPheGly Cys TyrValIle Arg LysPhePheGluPheGly
G	ValPheGly Ser TyrValIle Glu LysPhePheGluPheGly
U	ValPheGly Asn TyrValIle Gln LysPhePheGluPheGly

N ₆ nucleotide	R3 peptide sequence(s)
A	MetTyrGly CysArg ValIle Gln LysAlaLeuGluPheIle
	MetTyrGly SerArg ValIle Gln LysAlaLeuGluPheIle
С	MetTyrGly Ser ArgValIle Arg LysAlaLeuGluPheIle
	MetTyrGly Gly ArgValIle Arg LysAlaLeuGluPheIle
	MetTyrGly Ala ArgValIle Arg LysAlaLeuGluPheIle
	MetTyrGly Thr ArgValIle Arg LysAlaLeuGluPheIle
	MetTyrGly Cys ArgValIle Arg LysAlaLeuGluPheIle
G	MetTyrGly Ser ArgValIle Glu LysAlaLeuGluPheIle
U	MetTyrGly Asn ArgValIle Gln LysAlaLeuGluPheIle

N ₅ nucleotide	R4 peptide sequence(s)
A	GlnAsnGly CysArg ValVal Gln LysCysIleGluCysVal
	GlnAsnGly SerArg ValVal Gln LysCysIleGluCysVal
	GlnAsnGly Cys HisValVal Gln LysCysIleGluCysVal
	GlnAsnGly Ser HisValVal Gln LysCysIleGluCysVal
С	GlnAsnGly Ser HisValVal Arg LysCysIleGluCysVal
	GlnAsnGly Gly HisValVal Arg LysCysIleGluCysVal
	GlnAsnGly Ala HisValVal Arg LysCysIleGluCysVal
	GlnAsnGly Thr HisValVal Arg LysCysIleGluCysVal
	GlnAsnGly Cys HisValVal Arg LysCysIleGluCysVal
G	GlnAsnGly Ser HisValVal Glu LysCysIleGluCysVal
U	GlnAsnGly Asn HisValVal Gln LysCysIleGluCysVal

N ₄ nucleotide	R5 peptide sequence(s)
A	ProTyrGly CysArg ValIle Gln ArgIleLeuGluHisCys
	ProTyrGly SerArg ValIle Gln ArgIleLeuGluHisCys
С	ProTyrGly Ser ArgValIle Arg ArgIleLeuGluHisCys
	ProTyrGly Gly ArgValIle Arg ArgIleLeuGluHisCys
	ProTyrGly Ala ArgValIle Arg ArgIleLeuGluHisCys
	ProTyrGly Thr ArgValIle Arg ArgIleLeuGluHisCys
	ProTyrGly Cys ArgValIle Arg ArgIleLeuGluHisCys
G	ProTyrGly Ser ArgValIle Glu ArgIleLeuGluHisCys
U	ProTyrGly Asn ArgValIle Gln ArgIleLeuGluHisCys

N ₃ nucleotide	R6 peptide sequence(s)
A	GlnTyrGly CysArg ValIle Gln HisValLeuGluHisGly
	GlnTyrGly SerArg ValIle Gln HisValLeuGluHisGly
	GlnTyrGly Cys TyrValIle Gln HisValLeuGluHisGly
	GlnTyrGly Ser TyrValIle Gln HisValLeuGluHisGly
С	GlnTyrGly Ser TyrValIle Arg HisValLeuGluHisGly
	GlnTyrGly Gly TyrValIle Arg HisValLeuGluHisGly
	GlnTyrGly Ala TyrValIle Arg HisValLeuGluHisGly
	GlnTyrGly Thr TyrValIle Arg HisValLeuGluHisGly
	GlnTyrGly Cys TyrValIle Arg HisValLeuGluHisGly
G	GlnTyrGly Ser TyrValIle Glu HisValLeuGluHisGly
U	GlnTyrGly Asn TyrValIle Gln HisValLeuGluHisGly

N ₂ nucleotide	R7 peptide sequence(s)
A	LysPheAla CysArg ValVal Gln LysCysValThrHisAla
	LysPheAla SerArg ValVal Gln LysCysValThrHisAla
	LysPheAla Cys AsnValVal Gln LysCysValThrHisAla
	LysPheAla Ser AsnValVal Gln LysCysValThrHisAla

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N ₂ nucleotide	R7 peptide sequence(s)	
С	LysPheAla Ser AsnValVal Arg LysCysValThrHisAla	
	LysPheAla Gly AsnValVal Arg LysCysValThrHisAla	
	LysPheAla Ala AsnValVal Arg LysCysValThrHisAla	
	LysPheAla Thr AsnValVal Arg LysCysValThrHisAla	
	LysPheAla Cys AsnValVal Arg LysCysValThrHisAla	
G	LysPheAla Ser AsnValVal Glu LysCysValThrHisAla	
U	LysPheAla Asn AsnValVal Gln LysCysValThrHisAla	

N ₁ nucleotide	R8 peptide sequence(s)
A	GlnTyrAla CysArg ValVal Gln LysMetIleAspValAla
	GlnTyrAla SerArg ValVal Gln LysMetIleAspValAla
	GlnTyrAla Cys TyrValVal Gln LysMetIleAspValAla
	GlnTyrAla Ser TyrValVal Gln LysMetIleAspValAla
С	GlnTyrAla Ser TyrValVal Arg LysMetIleAspValAla
	GlnTyrAla Gly TyrValVal Arg LysMetIleAspValAla
	GlnTyrAla Ala TyrValVal Arg LysMetIleAspValAla
	GlnTyrAla Thr TyrValVal Arg LysMetIleAspValAla
	GlnTyrAla Cys TyrValVal Arg LysMetIleAspValAla
G	GlnTyrAla Ser TyrValVal Glu LysMetIleAspValAla
U	GlnTyrAla Asn TyrValVal Gln LysMetIleAspValAla

Several exemplary PUF domains with modified RNA binding specificity, constructed based on the above RNA recognition code, are provided below, each can be used to construct PUF domain-fusions of the invention.

PUF (3-2)

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```
Gly Arg Ser Arg Leu Leu Glu Asp Phe Arg Asn Asn Arg Tyr Pro Asn
    Leu Gln Leu Arg Glu Ile Ala Gly His Ile Met Glu Phe Ser Gln Asp
    Gln His Gly Ser Arg Phe Ile Gln Leu Lys Leu Glu Arg Ala Thr Pro
10
    Ala Glu Arq Gln Leu Val Phe Asn Glu Ile Leu Gln Ala Ala Tyr Gln
    Leu Met Val Asp Val Phe Gly Asn Tyr Val Ile Gln Lys Phe Phe Glu
    Phe Gly Ser Leu Glu Gln Lys Leu Ala Leu Ala Glu Arg Ile Arg Gly
    His Val Leu Ser Leu Ala Leu Gln Met Tyr Gly Ser Arq Val Ile Glu
    Lys Ala Leu Glu Phe Ile Pro Ser Asp Gln Gln Asn Glu Met Val Arg
15
    Glu Leu Asp Gly His Val Leu Lys Cys Val Lys Asp Gln Asn Gly Asn
    His Val Val Gln Lys Cys Ile Glu Cys Val Gln Pro Gln Ser Leu Gln
    Phe Ile Ile Asp Ala Phe Lys Gly Gln Val Phe Ala Leu Ser Thr His
    Pro Tyr Gly Cys Arq Val Ile Gln Arq Ile Leu Glu His Cys Leu Pro
    Asp Gln Thr Leu Pro Ile Leu Glu Glu Leu His Gln His Thr Glu Gln
20
    Leu Val Gln Asp Gln Tyr Gly Asn Tyr Val Ile Gln His Val Leu Glu
    His Gly Arg Pro Glu Asp Lys Ser Lys Ile Val Ala Glu Ile Arg Gly
    Asn Val Leu Val Leu Ser Gln His Lys Phe Ala Ser Asn Val Val Glu
    Lys Cys Val Thr His Ala Ser Arg Thr Glu Arg Ala Val Leu Ile Asp
    Glu Val Cys Thr Met Asn Asp Gly Pro His Ser Ala Leu Tyr Thr Met
25
    Met Lys Asp Gln Tyr Ala Asn Tyr Val Val Gln Lys Met Ile Asp Val
    Ala Glu Pro Gly Gln Arg Lys Ile Val Met His Lys Ile Arg Pro His
    Ile Ala Thr Leu Arg Lys Tyr Thr Tyr Gly Lys His Ile Leu Ala Lys
    Leu Glu Lys Tyr Tyr Met Lys Asn Gly Val Asp Leu Gly
```

PUF(3-2) has two point mutations (C935S/Q939E) in the PUF repeat 3, and recognizes a cognate RNA with a mutation at position 6 of the NRE (A6G; 5'- UGUAUGUA-3').

PUF (6-2/7-2)

```
Gly Arg Ser Arg Leu Leu Glu Asp Phe Arg Asn Asn Arg Tyr Pro Asn
    Leu Gln Leu Arg Glu Ile Ala Gly His Ile Met Glu Phe Ser Gln Asp
    Gln His Gly Ser Arg Phe Ile Gln Leu Lys Leu Glu Arg Ala Thr Pro
    Ala Glu Arq Gln Leu Val Phe Asn Glu Ile Leu Gln Ala Ala Tyr Gln
    Leu Met Val Asp Val Phe Gly Asn Tyr Val Ile Gln Lys Phe Phe Glu
    Phe Gly Ser Leu Glu Gln Lys Leu Ala Leu Ala Glu Arg Ile Arg Gly
    His Val Leu Ser Leu Ala Leu Gln Met Tyr Gly Cys Arg Val Ile Gln
    Lys Ala Leu Glu Phe Ile Pro Ser Asp Gln Gln Asn Glu Met Val Arg
10
    Glu Leu Asp Gly His Val Leu Lys Cys Val Lys Asp Gln Asn Gly Asn
    His Val Val Gln Lys Cys Ile Glu Cys Val Gln Pro Gln Ser Leu Gln
    Phe Ile Ile Asp Ala Phe Lys Gly Gln Val Phe Ala Leu Ser Thr His
    Pro Tyr Gly Cys Arg Val Ile Gln Arg Ile Leu Glu His Cys Leu Pro
    Asp Gln Thr Leu Pro Ile Leu Glu Glu Leu His Gln His Thr Glu Gln
    Leu Val Gln Asp Gln Tyr Gly Ser Tyr Val Ile Glu His Val Leu Glu
15
    His Gly Arg Pro Glu Asp Lys Ser Lys Ile Val Ala Glu Ile Arg Gly
    Asn Val Leu Val Leu Ser Gln His Lys Phe Ala Asn Asn Val Val Gln
    Lys Cys Val Thr His Ala Ser Arg Thr Glu Arg Ala Val Leu Ile Asp
    Glu Val Cys Thr Met Asn Asp Gly Pro His Ser Ala Leu Tyr Thr Met
20
    Met Lys Asp Gln Tyr Ala Asn Tyr Val Val Gln Lys Met Ile Asp Val
    Ala Glu Pro Gly Gln Arg Lys Ile Val Met His Lys Ile Arg Pro His
    Ile Ala Thr Leu Arg Lys Tyr Thr Tyr Gly Lys His Ile Leu Ala Lys
    Leu Glu Lys Tyr Tyr Met Lys Asn Gly Val Asp Leu Gly
```

PUF (6-2/7-2) has double point mutations (N1043S/Q1047E and S1079N/E1083Q) in repeats 6 and 7, respectively, and recognizes a cognate RNA sequence with two mutations at positions 2 and 3 of the NRE (GU/UG; 5'- UUGAUAUA-3').

A related PUF (6-2) has point mutations (N1043S/Q1047E) in repeats 6, and recognizes a cognate RNA sequence with a mutation at position 3 of the NRE (5'- UGGAUAUA-3').

Another related PUF (7-2) has point mutations (S1079N/E1083Q) in repeats 7, and recognizes a cognate RNA sequence with a mutation at position 2 of the NRE (5'-UUUAUAUA-3').

$PIJF^{531}$

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```
Gly Arg Ser Arg Leu Leu Glu Asp Phe Arg Asn Asn Arg Tyr Pro Asn
    Leu Gln Leu Arg Glu Ile Ala Gly His Ile Met Glu Phe Ser Gln Asp
35
    Gln His Gly Ser Arg Phe Ile Glu Leu Lys Leu Glu Arg Ala Thr Pro
    Ala Glu Arg Gln Leu Val Phe Asn Glu Ile Leu Gln Ala Ala Tyr Gln
    Leu Met Val Asp Val Phe Gly Asn Tyr Val Ile Gln Lys Phe Phe Glu
    Phe Gly Ser Leu Glu Gln Lys Leu Ala Leu Ala Glu Arg Ile Arg Gly
40
    His Val Leu Ser Leu Ala Leu Gln Met Tyr Gly Ser Arg Val Ile Glu
    Lys Ala Leu Glu Phe Ile Pro Ser Asp Gln Gln Asn Glu Met Val Arq
    Glu Leu Asp Gly His Val Leu Lys Cys Val Lys Asp Gln Asn Gly Asn
    His Val Val Gln Lys Cys Ile Glu Cys Val Gln Pro Gln Ser Leu Gln
    Phe Ile Ile Asp Ala Phe Lys Gly Gln Val Phe Ala Leu Ser Thr His
45
    Pro Tyr Gly Ser Arg Val Ile Glu Arg Ile Leu Glu His Cys Leu Pro
    Asp Gln Thr Leu Pro Ile Leu Glu Glu Leu His Gln His Thr Glu Gln
    Leu Val Gln Asp Gln Tyr Gly Asn Tyr Val Ile Gln His Val Leu Glu
```

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His Gly Arg Pro Glu Asp Lys Ser Lys Ile Val Ala Glu Ile Arg Gly Asn Val Leu Val Leu Ser Gln His Lys Phe Ala Ser Asn Val Val Glu Lys Cys Val Thr His Ala Ser Arg Thr Glu Arg Ala Val Leu Ile Asp Glu Val Cys Thr Met Asn Asp Gly Pro His Ser Ala Leu Tyr Thr Met Met Lys Asp Gln Tyr Ala Asn Tyr Val Val Gln Lys Met Ile Asp Val Ala Glu Pro Gly Gln Arg Lys Ile Val Met His Lys Ile Arg Pro His Ile Ala Thr Leu Arg Lys Tyr Thr Tyr Gly Lys His Ile Leu Ala Lys Leu Glu Lys Tyr Tyr Met Lys Asn Gly Val Asp Leu Gly
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The PUF domain PUF⁵³¹ has mutations (Q867E/Q939E/C935S/Q1011E/C1007S) in wild type PUF repeats 1, 3 and 5, and recognizes the sequence 5'-UGUGUGUG-3'. The PUF⁵³¹ can recognize its new target sequence with very high affinity, compared to the wild type PUF RNA.

Another modified PUF domain PUF(1-1) has one point mutation (Q867E) in the PUF repeat 1, and recognizes a cognate RNA with a mutation at position 8 of the NRE (A8G; 5'-UGUAUAUG-3').

Yet another modified PUF domain PUF(7-1) has one point mutation (E1083Q) in the PUF repeat 7, and recognizes a cognate RNA with a mutation at position 2 of the NRE (G2U; 5'- UUUAUAUA-3'; or G2A; 5'- UAUAUAUA-3').

Still another modified PUF domain PUF(3-1) has one point mutation (C935N) in the PUF repeat 3, and recognizes a cognate RNA with a mutation at position 6 of the NRE (A6U; 5'- UGUAUUUA-3').

A further modified PUF (7-2/3-1) has point mutations (C935N/S1079N/E1083Q) in repeats 7 and 3, and recognizes a cognate RNA sequence with mutations at positions 2 and 6 of the NRE (5'- UUUAUUUA-3').

The sequences of certain modified PUF domains are represented below.

```
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    Gly Arg Ser Arg Leu Leu Glu Asp Phe Arg Asn Asn Arg Tyr Pro Asn
    Leu Gln Leu Arg Glu Ile Ala Gly His Ile Met Glu Phe Ser Gln Asp
    Gln His Gly Ser Arg Phe Ile Glu Leu Lys Leu Glu Arg Ala Thr Pro
    Ala Glu Arg Gln Leu Val Phe Asn Glu Ile Leu Gln Ala Ala Tyr Gln
    Leu Met Val Asp Val Phe Gly Cys Arg Val Ile Gln Lys Phe Phe Glu
30
    Phe Gly Ser Leu Glu Gln Lys Leu Ala Leu Ala Glu Arg Ile Arg Gly
    His Val Leu Ser Leu Ala Leu Gln Met Tyr Gly Cys Arq Val Ile Gln
    Lys Ala Leu Glu Phe Ile Pro Ser Asp Gln Gln Asn Glu Met Val Arg
    Glu Leu Asp Gly His Val Leu Lys Cys Val Lys Asp Gln Asn Gly Asn
    His Val Val Gln Lys Cys Ile Glu Cys Val Gln Pro Gln Ser Leu Gln
35
    Phe Ile Ile Asp Ala Phe Lys Gly Gln Val Phe Ala Leu Ser Thr His
    Pro Tyr Gly Cys Arg Val Ile Gln Arg Ile Leu Glu His Cys Leu Pro
    Asp Gln Thr Leu Pro Ile Leu Glu Glu Leu His Gln His Thr Glu Gln
    Leu Val Gln Asp Gln Tyr Gly Ser Tyr Val Ile Glu His Val Leu Glu
    His Gly Arg Pro Glu Asp Lys Ser Lys Ile Val Ala Glu Ile Arg Gly
40
    Asn Val Leu Val Leu Ser Gln His Lys Phe Ala Asn Asn Val Val Gln
    Lys Cys Val Thr His Ala Ser Arg Thr Glu Arg Ala Val Leu Ile Asp
    Glu Val Cys Thr Met Asn Asp Gly Pro His Ser Ala Leu Tyr Thr Met
    Met Lys Asp Gln Tyr Ala Ser Tyr Val Val Glu Lys Met Ile Asp Val
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Ala Glu Pro Gly Gln Arg Lys Ile Val Met His Lys Ile Arg Pro His

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Ile Ala Thr Leu Arg Lys Tyr Thr Tyr Gly Lys His Ile Leu Ala Lys
    Leu Glu Lys Tyr Tyr
    Gly Arg Ser Arg Leu Leu Glu Asp Phe Arg Asn Asn Arg Tyr Pro Asn
    Leu Gln Leu Arg Glu Ile Ala Gly His Ile Met Glu Phe Ser Gln Asp
    Gln His Gly Asn Arg Phe Ile Gln Leu Lys Leu Glu Arg Ala Thr Pro
    Ala Glu Arg Gln Leu Val Phe Asn Glu Ile Leu Gln Ala Ala Tyr Gln
    Leu Met Val Asp Val Phe Gly Ser Tyr Val Ile Glu Lys Phe Phe Glu
    Phe Gly Ser Leu Glu Gln Lys Leu Ala Leu Ala Glu Arg Ile Arg Gly
10
    His Val Leu Ser Leu Ala Leu Gln Met Tyr Gly Ser Arg Val Ile Glu
    Lys Ala Leu Glu Phe Ile Pro Ser Asp Gln Gln Asn Glu Met Val Arg
    Glu Leu Asp Gly His Val Leu Lys Cys Val Lys Asp Gln Asn Gly Asn
    His Val Val Gln Lys Cys Ile Glu Cys Val Gln Pro Gln Ser Leu Gln
    Phe Ile Ile Asp Ala Phe Lys Gly Gln Val Phe Ala Leu Ser Thr His
15
    Pro Tyr Gly Ser Arg Val Ile Glu Arg Ile Leu Glu His Cys Leu Pro
    Asp Gln Thr Leu Pro Ile Leu Glu Glu Leu His Gln His Thr Glu Gln
    Leu Val Gln Asp Gln Tyr Gly Ser Tyr Val Ile Glu His Val Leu Glu
    His Gly Arg Pro Glu Asp Lys Ser Lys Ile Val Ala Glu Ile Arg Gly
20
    Asn Val Leu Val Leu Ser Gln His Lys Phe Ala Cys Asn Val Val Gln
    Lys Cys Val Thr His Ala Ser Arg Thr Glu Arg Ala Val Leu Ile Asp
    Glu Cys Val Thr Met Asn Asp Gly Pro His Ser Ala Leu Tyr Thr Met
    Met Lys Asp Gln Tyr Ala Ser Tyr Val Val Glu Lys Met Ile Asp Val
    Ala Glu Pro Gly Gln Arg Lys Ile Val Met His Lys Ile Arg Pro His
25
    Ile Ala Thr Leu Arg Lys Tyr Thr Tyr Gly Lys His Ile Leu Ala Lys
    Leu Glu Lys Tyr Tyr
    Gly Arg Ser Arg Leu Leu Glu Asp Phe Arg Asn Asn Arg Tyr Pro Asn
    Leu Gln Leu Arg Glu Ile Ala Gly His Ile Met Glu Phe Ser Gln Asp
30
    Gln His Gly Cys Arg Phe Ile Gln Leu Lys Leu Glu Arg Ala Thr Pro
    Ala Glu Arg Gln Leu Val Phe Asn Glu Ile Leu Gln Ala Ala Tyr Gln
    Leu Met Val Asp Val Phe Gly Ser Tyr Val Ile Glu Lys Phe Phe Glu
    Phe Gly Ser Leu Glu Gln Lys Leu Ala Leu Ala Glu Arg Ile Arg Gly
    His Val Leu Ser Leu Ala Leu Gln Met Tyr Gly Asn Arg Val Ile Gln
35
    Lys Ala Leu Glu Phe Ile Pro Ser Asp Gln Gln Asn Glu Met Val Arg
    Glu Leu Asp Gly His Val Leu Lys Cys Val Lys Asp Gln Asn Gly Asn
    His Val Val Gln Lys Cys Ile Glu Cys Val Gln Pro Gln Ser Leu Gln
    Phe Ile Ile Asp Ala Phe Lys Gly Gln Val Phe Ala Leu Ser Thr His
    Pro Tyr Gly Cys Arg Val Ile Gln Arg Ile Leu Glu His Cys Leu Pro
40
    Asp Gln Thr Leu Pro Ile Leu Glu Glu Leu His Gln His Thr Glu Gln
    Leu Val Gln Asp Gln Tyr Gly Ser Tyr Val Ile Glu His Val Leu Glu
    His Gly Arg Pro Glu Asp Lys Ser Lys Ile Val Ala Glu Ile Arg Gly
    Asn Val Leu Val Leu Ser Gln His Lys Phe Ala Cys Asn Val Val Gln
    Lys Cys Val Thr His Ala Ser Arg Thr Glu Arg Ala Val Leu Ile Asp
45
    Glu Cys Val Thr Met Asn Asp Gly Pro His Ser Ala Leu Tyr Thr Met
    Met Lys Asp Gln Tyr Ala Cys Tyr Val Val Gln Lys Met Ile Asp Val Ala Glu Pro Gly Gln Arg Lys Ile Val Met His Lys Ile Arg Pro His
    Ile Ala Thr Leu Arg Lys Tyr Thr Tyr Gly Lys His Ile Leu Ala Lys
    Leu Glu Lys Tyr Tyr
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According to the invention, heterologous polypeptide (also referred to as a "fusion partner") can be fused to the PUF domain of the invention that binds to at least one of the PBS on the subject polynucleotide. In addition, if desired, the same or different fusion partner can also optionally be fused to the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein). Thus as described herein, unless specifically disclaimed, any of the fusion partners are intended to be fused to PUF domain, and optionally also fused to the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein). The fusion partner fused to the PUF domain can be the same or different from the optional fusion partner fused to the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) (*infra*).

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The fusion partner may exhibit an activity (*e.g.*, enzymatic activity). Suitable fusion partners include, but are not limited to, a polypeptide that provides for methyltransferase activity, demethylase activity, acetyltransferase activity, deacetylase activity, kinase activity, phosphatase activity, ubiquitin ligase activity, deubiquitinating activity, adenylation activity, deadenylation activity, SUMOylating activity, deSUMOylating activity, ribosylation activity, deribosylation activity, myristoylation activity, or demyristoylation activity, any of which can be directed at modifying the DNA directly (*e.g.*, methylation of DNA) or at modifying a DNA-associated polypeptide (*e.g.*, a histone or DNA binding protein).

Protein name	Function			
Transcriptional Activators				
GAL4	Transcription activation			
VP16	Transcription activation			
VP64	Transcription activation			
p65 subdomain (NFkB)	Transcription activation			
Transcriptional repressors				
KRAB	Transcription repression			
Mad mSIN3 interaction domain (SID)	Transcription repression			
the ERF repressor domain (ERD)	Transcription repression			
Histone lysine methyltransferases (KMT)				
KMT1 family: SUV39H1, SUV39H2, G9A,	Heterochromatin formation/			
ESET/SETDB1, and homologs (Clr4, Su(var)3-9)	transcription repression			
KMT2 family: hSET1A, hSET1B, MLL1 to 5, ASH1,	Transcription activation			
and homologs (Trx, Trr, Ash1)	Transcription activation			
KMT3 family: SYMD2, NSD1	Transcription activation			
KMT4: DOT1L and homologs	Transcription activation			
KMT5 family: Pr-SET7/8, SUV4-20H1, and	DNA damage response,			
homologs (PR-set7, Suv4-20, Set9)	transcription repression			
KMT6: EZH2	Polycomb silencing			
KMT8: RIZ1	Transcription repression			
Histone lysine demethylates (KDM)				
KDM1: LSD1/BHC110 and homologs	Transcription activation and repression,			
(SpLsd1/Swm1/Saf110, Su(var)3-3)	heterochromatin formation			
KDM3 family: JHDM2a/b	Androgen receptor gene activation,			
	spermatogenesis			
KDM4 family: JMJD2A/JHDM3A, JMJD2B,	Transcription elongation, transcription			
JMJD2C/GASC1, JMJD2D, and homologs (Rph1)	repression, heterochromatin formation,			
	genome integrity			
KDM5 family: JARID1A/RBP2, JARID1B/PLU-1,	Transcription repression			
JARID1C/SMCX, JARID1D/SMCY,				
and homologs (Lid, Jhn2, Jmj2)				
KDM6 family: UTX, JMJD3	Transcription activation			

Protein name	Function		
Histone lysine acetyltransferases (KAT)			
KAT2 family: hGCN5, PCAF, and homologs	Town influence for the DNA consistence		
(dGCN5/PCAF, Gcn5)	Transcription activation, DNA repair		
KAT3 family: CBP, p300, and homologs (dCBP/NEJ)	Transcription activation, DNA repair		
KAT4: TAF1 and homologs (dTAF1)	Transcription activation		
KAT5: TIP60/PLIP, and homologs	Transcription activation, DNA repair		
KAT6: MOZ/MYST3, MORF/MYST4, and homologs	Transcription activation and elongation,		
(Mst2, Sas3, CG1894)	DNA replication		
KAT7: HBO1/MYST2, and homologs (CHM, Mst2)	Transcription, DNA replication		
KAT8: HMOF/MYST1, and homologs (dMOF,	Chromatin boundaries, dosage		
CG1894, Sas2, Mst2)	compensation, DNA repair		
KAT13 family: SRC1, ACTR, P160, CLOCK, and	Transcription activation		
homologs			
Histone lysine deacetylases			
Class I: HDAC1, HDAC2, HDAC3, HDAC8, and its	Transcription repression,		
homologs (Rpd3, Hos1 , Cir6)	heterochromatin formation		
Class lla: HDAC4, HDAC5, HDAC7, HDAC9, and	Transcription repression,		
its homologs (Hda1, Cir3 etc.)	heterochromatin formation		
Class III: SIRT1, SIRT2, and its homologs (Sir2,	Transcription repression,		
Hst1, Hst2, Hst3, Hst4)	heterochromatin formation		
Class IV: HDAC11	Transcription repression		
DNA methylases(adenosine or cytosine modification)			
Dam (E. coli)	Restriction system		
Dcm (E. coli)	Restriction system		
M. Sssl (Spiroplasma sp)	Restriction system		
DNMT1	Transcription repression. imprinting,		
DIVIVITI	heterochromatin formation		
DNMT3a/DNMT3b, METI, DRM3 (plants), and	Transcription repression. imprinting,		
homologs	heterochromatin formation		
Chromomethylases e.g. ZMET2, CMT1, CMT2 (plants)	Transcription repression. imprinting,		
Official official ses e.g. Zivie 12, Owi 11, Owi 12 (plants)	heterochromatin formation		
DNA demethylases			
AID/Apobec deaminase family: AID	Transcription activation, genome integrity		
TET dioxygenase family: TET1	Transcription activation, genome integrity		
DEMETER glycosylase family: DME, DML1, DML2,	Transcription activation, genome		
ROS1	integrity		

Protein name	Function
Boundary elements	
CTCF	Chromatin insulation, heterochromatin spreading suppression
Periphery recruitment elements	
Lamin A	Transcription repression
Lamin B	Transcription repression
Protein docking elements	
FKBP/FRB (S. pombe)	rapamycin dependent recruitment
Pil1/Aby1 (E. coli)	ABA dependent recruitment

Additional fusion partners may include the various fluorescent protein, polypeptides, variants, or functional domains thereof, such as GFP, Superfolder GFP, EGFP, BFP, EBFP, EBFP2, Azurite, mKalama1, CFP, ECFP, Cerulean, CyPet, mTurquoise2, YFP, Citrine, Venus, Ypet, BFPms1, roGFP, and bilirubin-inducible fluorescent proteins such as UnaG, dsRed, eqFP611, Dronpa, TagRFPs, KFP, EosFP, Dendra, IrisFP, *etc*.

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Further suitable fusion partners include, but are not limited to boundary elements (*e.g.*, CTCF), proteins and fragments thereof that provide periphery recruitment (*e.g.*, Lamin A, Lamin B, *etc.*), and protein docking elements (*e.g.*, FKBP/FRB, Pill/Abyl, *etc.*).

Additional non-limiting examples of fusion partners to accomplish increased or decreased transcription are listed below, and include transcription activator and transcription repressor domains (*e.g.*, the Kruppel associated box (KRAB or SKD); the Mad mSIN3 interaction domain (SID); the ERF repressor domain (ERD), *etc.*).

In some embodiments, the heterologous sequence can be fused to the C-terminus of the PUF domain or Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein). In some embodiments, the heterologous sequence can be fused to the N-terminus of the PUF domain or Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein). In some embodiments, the heterologous sequence can be fused to an internal portion (*i.e.*, a portion other than the N- or C- terminus) of the PUF domain or Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein).

In some embodiments, a PUF domain fusion is generated by fusing a PUF domain with a heterologous sequence that provides for subcellular localization (*i.e.*, the heterologous sequence is a subcellular localization sequence, *e.g.*, a nuclear localization signal (NLS, such as PPKKKRKV) for targeting to the nucleus; a mitochondrial localization signal for targeting to the mitochondria; a chloroplast localization signal for targeting to a chloroplast; an ER retention signal; and the like). In some embodiments, the heterologous sequence can provide a tag (*i.e.*,

the heterologous sequence is a detectable label) for ease of tracking and/or purification (e.g., a fluorescent protein, e.g., green fluorescent protein (GFP), YFP, RFP, CFP, mCherry, tdTomato, and the like; a histidine tag, e.g., a 6×His tag; a hemagglutinin (HA) tag; a FLAG tag; a Myc tag; and the like). In some embodiments, the heterologous sequence can provide for increased or decreased stability (i.e., the heterologous sequence is a stability control peptide, e.g., a degron, which in some cases is controllable (e.g., a temperature sensitive or drug controllable degron sequence, see below). In some embodiments, the heterologous sequence can provide for increased or decreased transcription from the target DNA (i.e., the heterologous sequence is a transcription modulation sequence, e.g., a transcription factor / activator or a fragment thereof, a protein or fragment thereof that recruits a transcription factor / activator, a transcription repressor or a fragment thereof, a protein or fragment thereof that recruits a transcription repressor, a small molecule / drug-responsive transcription regulator, etc.). In some embodiments, the heterologous sequence can provide a binding domain (i.e., the heterologous sequence is a protein binding sequence, e.g., to provide the ability of a chimeric PUF domain or Cas9 protein (e.g., wt, nickase, or dCas9 protein) to bind to another protein of interest, e.g., a DNA or histone modifying protein, a transcription factor or transcription repressor, a recruiting protein, etc.).

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Suitable fusion partners that provide for increased or decreased stability include, but are not limited to degron sequences. Degrons are readily understood by one of ordinary skill in the art to be amino acid sequences that control the stability of the protein of which they are part. For example, the stability of a protein comprising a degron sequence is controlled at least in part by the degron sequence. In some cases, a suitable degron is constitutive such that the degron exerts its influence on protein stability independent of experimental control (i.e., the degron is not drug inducible, temperature inducible, etc.). In some cases, the degron provides the PUF domain or Cas9 protein (e.g., wt, nickase, or dCas9 protein) with controllable stability such that the PUF domain or Cas9 protein (e.g., wt, nickase, or dCas9 protein) can be turned "on" (i.e., stable) or "off (i.e., unstable, degraded) depending on the desired conditions. For example, if the degron is a temperature sensitive degron, the PUF domain or Cas9 protein (e.g., wt, nickase, or dCas9 protein) may be functional (i.e., "on", stable) below a threshold temperature (e.g., 42°C, 41°C, 40°C, 39°C, 38°C, 37°C, 36°C, 35°C, 34°C, 33°C, 32°C, 31°C, 30°C, etc.) but non-functional (i.e., "off, degraded) above the threshold temperature. As another example, if the degron is a drug inducible degron, the presence or absence of drug can switch the protein from an "off (i.e., unstable) state to an "on" (i.e., stable) state or vice versa. An exemplary drug inducible degron is derived from the FKBP12 protein. The stability of the degron is controlled by the presence or

absence of a small molecule that binds to the degron.

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Examples of suitable degrons include, but are not limited to those degrons controlled by Shield-1, DHFR, auxins, and/or temperature. Non-limiting examples of suitable degrons are known in the art (*e.g.*, Dohmen *et al.*, *Science*, 263(5151): 1273-1276, 1994: "Heat-inducible degron: a method for constructing temperature-sensitive mutants"; Schoeber *et al.*, *Am. J. Physiol. Renal. Physiol.*, 296(1):F204-211, 2009: "Conditional fast expression and function of multimeric TRPV5 channels using Shield-1"; Chu *et al.*, *Bioorg. Med. Chem. Lett.*, 18(22): 5941-4, 2008: "Recent progress with FKBP-derived destabilizing domains"; Kanemaki, *Pflugers Arch.*, 2012: "Frontiers of protein expression control with conditional degrons"; Yang *et al.*, *Mol. Cell.*, 48(4):487-8, 2012: "Titivated for destruction: the methyl degron"; Barbour *et al.*, *Biosci. Rep.*, 33(1), 2013: "Characterization of the bipartite degron that regulates ubiquitin-independent degradation of thymidylate synthase"; and Greussing *et al.*, *J. Vis. Exp.*, (69), 2012: "Monitoring of ubiquitin-proteasome activity in living cells using a Degron (dgn)-destabilized green fluorescent protein (GFP)-based reporter protein"; all of which are incorporated in their entirety by reference).

Exemplary degron sequences have been well-characterized and tested in both cells and animals. Thus, fusing Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) to a degron sequence produces a "tunable" and "inducible" PUF domain or Cas9 (*e.g.*, wt, nickase, or dCas9 protein).

Any of the fusion partners described herein can be used in any desirable combination. As one non-limiting example to illustrate, each PUF domain can be independently fused to the same or different fusion partners, and they may bind in any order on the series of PBS of the subject polynucleotide. For example, one PUF domain can be fused to a YFP sequence for detection, a second PUF domain fused to a degron sequence for stability, and a third PUF domain fused to a transcription activator sequence to increase transcription of the target DNA. Any of these types of PUF domain fusions can have more than 1 binding sites or PBS on the subject polynucleotide, in any desired order. The number of fusion partners that can be used in the PUF domain fusions is largely unlimited (*e.g.*, at least 2, 5, 10, 20, 30, 40, 50 or more).

In some embodiments, any PUF domain or Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) fusion protein may comprises one or more (*e.g.* two or more, three or more, four or more, or five or more) heterologous sequences or fusion partners.

In some embodiments, any of the subject Cas9 protein (e.g., wt, nickase, or dCas9 protein) or PUF domain fusions can be codon-optimized. This type of optimization is known in the art and entails the mutation of foreign-derived DNA to mimic the codon preferences of the

intended host organism or cell while encoding the same protein. Thus, the codons are changed, but the encoded protein remains unchanged. For example, if the intended target cell was a human cell, a human codon-optimized PUF domain or Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) fusion would be a better suited PUF domain or Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) fusion. As another non-limiting example, if the intended host cell were a mouse cell, than a mouse codon-optimized PUF domain fusion or Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) would be a suitable PUF domain fusion or Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein). While codon optimization is not required, it is acceptable and may be preferable in certain cases.

Any of the subject PUF domain can be made using, for example, a Golden Gate Assembly kit (see Abil *et al.*, *Journal of Biological Engineering* 8:7, 2014), which is available at Addgene (Kit # 1000000051).

5. Modulation of Transcription

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The PUF domain and/or Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) fusion protein of the invention is targeted by the DNA-targeting sequence of the subject polynucleotide to a specific location (*i.e.*, target polynucleotide sequence) in the target DNA, and exerts locusspecific regulation, such as blocking RNA polymerase binding to a promoter (which selectively inhibits transcription activator function), and/or modifying the local chromatin status (*e.g.*, when a fusion sequence is used that modifies the target DNA or modifies a polypeptide associated with the target DNA). In some cases, the changes are transient (*e.g.*, transcription repression or activation). In some cases, the changes are inheritable (*e.g.*, when epigenetic modifications are made to the target DNA or to proteins associated with the target DNA, *e.g.*, nucleosomal histones).

The biological effects of a method using a subject PUF domain or Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) fusion protein can be detected by any convenient method (*e.g.*, gene expression assays; chromatin-based assays, *e.g.*, Chromatin immunoPrecipitation (ChiP), Chromatin *in vivo* Assay (CiA), *etc.*; and the like).

In some cases, a subject method involves using two or more different DNA-targeting sequences. For example, two different DNA-targeting sequences can be used in a single host cell, where the two different DNA-targeting sequences target two different target polynucleotide sequences in the same target nucleic acid. Thus, for example, a subject transcriptional modulation method can further comprise introducing into the host cell a second DNA-targeting

sequence, or a nucleic acid comprising a nucleotide sequence encoding the second DNA-targeting sequence. In some cases, use of two different DNA-targeting sequences targeting two different targeting sequences in the same target nucleic acid provides for increased modulation (e.g., reduction or increase) in transcription of the target nucleic acid.

As another example, two different DNA-targeting sequences can be used in a single host cell, where the two different DNA-targeting sequences target two different target nucleic acids.

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Thus, in certain embodiments, a transcription modulation method of the present invention provides for selective modulation (*e.g.*, reduction or increase) of a target nucleic acid in a host cell. For example, "selective" reduction of transcription of a target nucleic acid reduces transcription of the target nucleic acid by at least about 10%, at least about 20%, at least about 30%, at least about 40%, at least about 50%, at least about 60%, at least about 70%, at least about 80%, at least about 90%, or greater than 90%, compared to the level of transcription of the target nucleic acid in the absence of a DNA-targeting sequence / modified Cas9 polypeptide / PUF domain-fusion complex. Selective reduction of transcription of a target nucleic acid reduces transcription of the target nucleic acid, but does not substantially reduce transcription of a non-target nucleic acid, *e.g.*, transcription of a non-target nucleic acid is reduced, if at all, by less than 10% compared to the level of transcription of the non-target nucleic acid in the absence of the DNA-targeting sequence / modified Cas9 polypeptide / PUF domain-fusion complex.

On the other hand, "selective" increased transcription of a target DNA can increase transcription of the target DNA by at least about 1.1 fold (*e.g.*, at least about 1.2 fold, at least about 1.3 fold, at least about 1.4 fold, at least about 1.5 fold, at least about 1.6 fold, at least about 1.7 fold, at least about 1.8 fold, at least about 1.9 fold, at least about 2 fold, at least about 2.5 fold, at least about 3 fold, at least about 3.5 fold, at least about 4 fold, at least about 4.5 fold, at least about 5 fold, at least about 6 fold, at least about 7 fold, at least about 8 fold, at least about 9 fold, at least about 10 fold, at least about 12 fold, at least about 15 fold, or at least about 20-fold) compared to the level of transcription of the target DNA in the absence of the DNA-targeting sequence / modified Cas9 polypeptide / PUF domain-fusion complex. Selective increase of transcription of a target DNA increases transcription of the target DNA, but does not substantially increase transcription of a non-target DNA, *e.g.*, transcription of a non-target DNA is increased, if at all, by less than about 5-fold (*e.g.*, less than about 4-fold, less than about 3-fold, less than about 2-fold, less than about 1.8-fold, less than about 1.6-fold, less than about 1.4-fold, less than about 1.2-fold, or less than about 1.1 -fold) compared to the level of transcription of the non-targeted DNA in the absence of the DNA-targeting sequence / modified Cas9

polypeptide / PUF domain-fusion complex.

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As a non-limiting example, increased transcription can be achieved by fusing dCas9 to a heterologous sequence, and/or by fusing the heterologous sequence to one of the PUF domains that binds to a PBS of the subject polynucleotide. Suitable fusion partners include, but are not limited to, a polypeptide that provides an activity that indirectly increases transcription by acting directly on the target DNA or on a polypeptide (*e.g.*, a histone or other DNA-binding protein) associated with the target DNA. Suitable fusion partners include, but are not limited to, a polypeptide that provides for methyltransferase activity, demethylase activity, acetyltransferase activity, deacetylase activity, kinase activity, phosphatase activity, ubiquitin ligase activity, deubiquitinating activity, adenylation activity, deadenylation activity, SUMOylating activity, or demyristoylation activity, ribosylation activity, deribosylation activity, myristoylation activity, or demyristoylation activity.

Additional suitable fusion partners include, but are not limited to, a polypeptide that directly provides for increased transcription of the target nucleic acid (e.g., a transcription activator or a fragment thereof, a protein or fragment thereof that recruits a transcription activator, a small molecule/drug-responsive transcription regulator, etc.). See section entitled "PUF domain (and the optional dCas9) Fusion Proteins."

A non-limiting example of a subject method using a dCas9 fusion protein and/or a PUF domain-fusion protein to increase transcription in a prokaryote includes a modification of the bacterial one-hybrid (B1H) or two-hybrid (B2H) system. In the B1H system, a DNA binding domain (BD) is fused to a bacterial transcription activation domain (AD, e.g., the alpha subunit of the E. coli RNA polymerase (RNAPα)). Thus, a subject dCas9 or PUF domain can be fused to a heterologous sequence comprising an AD. When the subject dCas9 or PUF domain fusion protein arrives at the upstream region of a promoter (targeted there by the DNA-targeting sequence) the AD (e.g., RNAPα) of the dCas9 or PUF domain fusion protein recruits the RNAP holoenzyme, leading to transcription activation. In the B2H system, the BD is not directly fused to the AD; instead, their interaction is mediated by a protein-protein interaction (e.g., GAL11P-GAL4 interaction). To modify such a system for use in the subject methods, dCas9 or PUF domain can be fused to a first protein sequence that provides for protein-protein interaction (e.g., the yeast GAL11P and/or GAL4 protein) and RNAPa can be fused to a second protein sequence that completes the protein-protein interaction (e.g., GAL4 if GAL11P is fused to dCas9 or PUF domain, GAL11P if GAL4 is fused to dCas9 or PUF domain, etc.). The binding affinity between GAL11P and GAL4 increases the efficiency of binding and transcription rate.

A non-limiting example of a subject method using a dCas9 and/or PUF domain fusion protein to increase transcription in a eukaryotes includes fusion of dCas9 and/or PUF domain to an activation domain (AD) (*e.g.*, GAL4, herpesvirus activation protein VP16 or VP64, human nuclear factor NF-kB p65 subunit, *etc.*). To render the system inducible, expression of the dCas9 / PUF domain fusion protein can be controlled by an inducible promoter (*e.g.*, Tet-ON, Tet-OFF, *etc.*). The DNA-targeting sequence can be designed to target known transcription response elements (*e.g.*, promoters, enhancers, *etc.*), known upstream activating sequences (UAS), sequences of unknown or known function that are suspected of being able to control expression of the target DNA, *etc.*

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In some embodiments, multiple subject polynucleotides are used simultaneously in the same cell to simultaneously modulate transcription at different locations on the same target DNA or on different target DNAs. In some embodiments, two or more subject polynucleotides target the same gene or transcript or locus. In some embodiments, two or more subject polynucleotides target different unrelated loci. In some embodiments, two or more subject polynucleotides target different, but related loci.

Because the subject polynucleotides are small and robust, they can be simultaneously present on the same expression vector and can even be under the same transcriptional control if so desired. In some embodiments, two or more (*e.g.*, 3 or more, 4 or more, 5 or more, 10 or more, 15 or more, 20 or more, 25 or more, 30 or more, 35 or more, 40 or more, 45 or more, or 50 or more) subject polynucleotides are simultaneously expressed in a target cell, from the same or different vectors. The expressed subject polynucleotides can be differently recognized by orthogonal dCas9 proteins from different bacteria, such as *S. pyogenes*, *S. thermophilus*, *L. innocua*, and *N. meningitidis*.

To express multiple subject polynucleotides, an artificial RNA processing system mediated by the Csy4 endoribonuclease can be used. Multiple subject polynucleotides can be concatenated into a tandem array on a precursor transcript (*e.g.*, expressed from a U6 promoter), and separated by Csy4-specific RNA sequence. Co-expressed Csy4 protein cleaves the precursor transcript into multiple subject polynucleotides. Advantages for using an RNA processing system include: first, there is no need to use multiple promoters or vectors; second, since all subject polynucleotides are processed from a precursor transcript, their concentrations are normalized for similar wt Cas9 / Cas9 nickase / dCas9-binding.

Csy4 is a small endoribonuclease (RNase) protein derived from bacteria *Pseudomonas* aeruginosa. Csy4 specifically recognizes a minimal 17-bp RNA hairpin, and exhibits rapid (<1

min) and highly efficient (>99.9) RNA cleavage. Unlike most RNases, the cleaved RNA fragment remains stable and functionally active. The Csy4-based RNA cleavage can be repurposed into an artificial RNA processing system. In this system, the 17-bp RNA hairpins are inserted between multiple RNA fragments that are transcribed as a precursor transcript from a single promoter. Co-expression of Csy4 is effective in generating individual RNA fragments.

6. Host Cells

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A method of the present invention to modulate transcription may be employed to induce transcriptional modulation in mitotic or post-mitotic cells *in vivo* and/or ex vivo and/or *in vitro*. Because the subject polynucleotide provides specificity by hybridizing to target polynucleotide sequence of a target DNA, a mitotic and/or post-mitotic cell can be any of a variety of host cell, where suitable host cells include, but are not limited to, a bacterial cell; an archaeal cell; a single-celled eukaryotic organism; a plant cell; an algal cell, *e.g.*, *Botryococcus braunii*, *Chlamydomonas reinhardtii*, *Nannochloropsis gaditana*, *Chlorella pyrenoidosa*, *Sargassum patens*, *C. agardh*, and the like; a fungal cell; an animal cell; a cell from an invertebrate animal (*e.g.*, an insect, a cnidarian, an echinoderm, a nematode, *etc.*); a eukaryotic parasite (*e.g.*, a malarial parasite, *e.g.*, *Plasmodium falciparum*; a helminth; *etc.*); a cell from a vertebrate animal (*e.g.*, fish, amphibian, reptile, bird, mammal); a mammalian cell, *e.g.*, a rodent cell, a human cell, a non-human primate cell, *etc.* Suitable host cells include naturally-occurring cells; genetically modified cells (*e.g.*, cells genetically modified in a laboratory, *e.g.*, by the "hand of man"); and cells manipulated *in vitro* in any way. In some cases, a host cell is isolated or cultured.

Any type of cell may be of interest (*e.g.*, a stem cell, *e.g.* an embryonic stem (ES) cell, an induced pluripotent stem (iPS) cell, a germ cell; a somatic cell, *e.g.* a fibroblast, a hematopoietic cell, a neuron, a muscle cell, a bone cell, a hepatocyte, a pancreatic cell; an *in vitro* or *in vivo* embryonic cell of an embryo at any stage, *e.g.*, a 1-cell, 2-cell, 4-cell, 8-cell, *etc.* stage zebrafish embryo; *etc.*). Cells may be from established cell lines or they may be primary cells, where "primary cells," "primary cell lines," and "primary cultures" are used interchangeably herein to refer to cells and cells cultures that have been derived from a subject and allowed to grow *in vitro* for a limited number of passages, *i.e.* splittings, of the culture. For example, primary cultures include cultures that may have been passaged 0 times, 1 time, 2 times, 4 times, 5 times, 10 times, or 15 times, but not enough times go through the crisis stage. Primary cell lines can be are maintained for fewer than 10 passages *in vitro*. Target cells are in many embodiments unicellular organisms, or are grown in culture.

If the cells are primary cells, such cells may be harvest from an individual by any

convenient method. For example, leukocytes may be conveniently harvested by apheresis, leukocytapheresis, density gradient separation, *etc.*, while cells from tissues such as skin, muscle, bone marrow, spleen, liver, pancreas, lung, intestine, stomach, *etc.* are most conveniently harvested by biopsy. An appropriate solution may be used for dispersion or suspension of the harvested cells. Such solution will generally be a balanced salt solution, *e.g.* normal saline, phosphate-buffered saline (PBS), Hank's balanced salt solution, *etc.*, conveniently supplemented with fetal calf serum or other naturally occurring factors, in conjunction with an acceptable buffer at low concentration, *e.g.*, from 5-25 mM. Convenient buffers include HEPES, phosphate buffers, lactate buffers, *etc.* The cells may be used immediately, or they may be stored, frozen, for long periods of time, being thawed and capable of being reused. In such cases, the cells will usually be frozen in 10% dimethyl sulfoxide (DMSO), 50% serum, 40% buffered medium, or other solutions commonly used in the art to preserve cells at such freezing temperatures, and thawed in a manner as commonly known in the art for thawing frozen cultured cells.

15 7. Introducing Nucleic Acid into a Host Cell

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A subject polynucleotide, a nucleic acid comprising a nucleotide sequence encoding same, or a nucleic acid comprising a nucleotide sequence encoding the subject Cas9 protein (e.g., wt, nickase, or dCas9 protein) or PUF domain fusion, can be introduced into a host cell by any of a variety of well-known methods.

Methods of introducing a nucleic acid into a host cell are known in the art, and any known method can be used to introduce a nucleic acid (*e.g.*, vector or expression construct) into a stem cell or progenitor cell. Suitable methods include, include *e.g.*, viral or bacteriophage infection, transfection, conjugation, protoplast fusion, lipofection, electroporation, calcium phosphate precipitation, polyethyleneimine (PEI)-mediated transfection, DEAE-dextran mediated transfection, liposome-mediated transfection, particle gun technology, calcium phosphate precipitation, direct micro injection, nanoparticle-mediated nucleic acid delivery (see, *e.g.*, Panyam *et al.*, *Adv. Drug Deliv. Rev.*, pii: S0169-409X(12)00283-9.doi:10.1016 / j.addr.2012.09.023), and the like.

Thus the present invention also provides an isolated nucleic acid comprising a nucleotide sequence encoding a subject polynucleotide. In some cases, a subject nucleic acid also comprises a nucleotide sequence encoding a subject Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) and/or a subject PUF domain fusion.

In some embodiments, a subject method involves introducing into a host cell (or a population of host cells) one or more nucleic acids (*e.g.*, vectors) comprising nucleotide sequences encoding a subject polynucleotide and/or a subject Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) and/or a subject PUF domain fusion. In some embodiments a host cell comprising a target DNA is *in vitro*. In some embodiments a host cell comprising a target DNA is *in vivo*. Suitable nucleic acids comprising nucleotide sequences encoding a subject polynucleotide and/or a subject Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) and/or a subject PUF domain fusion include expression vectors, where the expression vectors may be recombinant expression vector.

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In some embodiments, the recombinant expression vector is a viral construct, *e.g.*, a recombinant adeno-associated virus construct (see, *e.g.*, U.S. Patent No. 7,078,387), a recombinant adenoviral construct, a recombinant lentiviral construct, a recombinant retroviral construct, *etc*.

Suitable expression vectors include, but are not limited to, viral vectors (e.g. viral vectors based on vaccinia virus; poliovirus; adenovirus (see, e.g., Li et al., Invest Opthalmol. Vis. Sci., 35:2543-2549, 1994; Borras et al., Gene Ther., 6:515-524, 1999; Li and Davidson, Proc. Natl. Acad. Sci. USA, 92:7700-7704, 1995; Sakamoto et al., Hum. Gene Ther., 5:1088-1097, 1999; WO 94/12649, WO 93/03769; WO 93/19191; WO 94/28938; WO 95/11984 and WO 95/00655); adeno-associated virus (see, e.g., Ali et al., Hum. Gene Ther., 9:81-86, 1998, Flannery et al., Proc. Natl. Acad. Sci. USA, 94:6916-6921, 1997; Bennett et al., Invest Opthalmol Vis Sci 38:2857-2863, 1997; Jomary et al., Gene Ther., 4:683-690, 1997, Rolling et al., Hum. Gene Ther., 10:641-648, 1999; Ali et al., Hum. Mol. Genet., 5:591-594, 1996; Srivastava in WO 93/09239, Samulski et al., J. Vir., 63:3822-3828, 1989; Mendelson et al., Virol., 166: 154-165, 1988; and Flotte et al., Proc. Natl. Acad. Sci. USA, 90: 10613-10617, 1993); SV40; herpes simplex virus; human immunodeficiency virus (see, e.g., Miyoshi et al., Proc. Natl. Acad. Sci. USA, 94: 10319-23, 1997; Takahashi et al., J. Virol., 73:7812-7816, 1999); a retroviral vector (e.g., Murine Leukemia Virus, spleen necrosis virus, and vectors derived from retroviruses such as Rous Sarcoma Virus, Harvey Sarcoma Virus, avian leukosis virus, a lentivirus, HIV virus, myeloproliferative sarcoma virus, and mammary tumor virus); and the like.

Numerous suitable expression vectors are known to those skilled in the art, and many are commercially available. The following vectors are provided by way of example; for eukaryotic host cells: pXTl, pSG5 (Stratagene), pSVK3, pBPV, pMSG, and pSVLSV40 (Pharmacia). However, any other vector may be used so long as it is compatible with the host cell.

Depending on the host/vector system utilized, any of a number of suitable transcription and translation control elements, including constitutive and inducible promoters, transcription enhancer elements, transcription terminators, *etc.* may be used in the expression vector (see *e.g.*, Bitter *et al.*, *Methods in Enzymology*, 153:516-544, 1987).

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In some embodiments, a nucleotide sequence encoding a subject polynucleotide and/or a subject Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) and/or a subject PUF domain fusion is operably linked to a control element, *e.g.*, a transcriptional control element, such as a promoter. The transcriptional control element may be functional in either a eukaryotic cell, *e.g.*, a mammalian cell; or a prokaryotic cell (*e.g.*, bacterial or archaeal cell). In some embodiments, a nucleotide sequence encoding a subject polynucleotide and/or a subject Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) and/or a subject PUF domain fusion is operably linked to multiple control elements that allow expression of the nucleotide sequence encoding the subject polynucleotide and/or a subject Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) and/or a subject PUF domain fusion in both prokaryotic and eukaryotic cells.

A promoter can be a constitutively active promoter (*i.e.*, a promoter that is constitutively in an active/"ON" state), it may be an inducible promoter (*i.e.*, a promoter whose state, active/"ON" or inactive/"OFF", is controlled by an external stimulus, *e.g.*, the presence of a particular temperature, compound, or protein.), it may be a spatially restricted promoter (*i.e.*, transcriptional control element, enhancer, *etc.*) (*e.g.*, tissue specific promoter, cell type specific promoter, *etc.*), and it may be a temporally restricted promoter (*i.e.*, the promoter is in the "ON" state or "OFF" state during specific stages of embryonic development or during specific stages of a biological process, *e.g.*, hair follicle cycle in mice).

Suitable promoters can be derived from viruses and can therefore be referred to as viral promoters, or they can be derived from any organism, including prokaryotic or eukaryotic organisms. Suitable promoters can be used to drive expression by any RNA polymerase (*e.g.*, pol I, pol II, pol III). Exemplary promoters include, but are not limited to the SV40 early promoter, mouse mammary tumor virus long terminal repeat (LTR) promoter; adenovirus major late promoter (Ad MLP); a herpes simplex virus (HSV) promoter, a cytomegalovirus (CMV) promoter such as the CMV immediate early promoter region (CMVIE), a rous sarcoma virus (RSV) promoter, a human U6 small nuclear promoter (U6) (Miyagishi *et al.*, *Nature Biotech.*, 20:497-500, 2002), an enhanced U6 promoter (*e.g.*, Xia *et al.*, *Nucleic Acids Res.*, 31(17):e100, 2003), a human HI promoter (HI), and the like.

Examples of inducible promoters include, but are not limited to T7 RNA polymerase

promoter, T3 RNA polymerase promoter, Isopropyl-beta-D-thiogalactopyranoside (IPTG)-regulated promoter, lactose induced promoter, heat shock promoter, Tetracycline-regulated promoter (*e.g.*, Tet-ON, Tet-OFF, *etc.*), Steroid-regulated promoter, Metal-regulated promoter, estrogen receptor-regulated promoter, *etc.* Inducible promoters can therefore be regulated by molecules including, but not limited to, doxycycline; RNA polymerase, *e.g.*, T7 RNA polymerase; an estrogen receptor; an estrogen receptor fusion; *etc.*

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In some embodiments, the promoter is a spatially restricted promoter (*i.e.*, cell type specific promoter, tissue specific promoter, *etc.*) such that in a multi-cellular organism, the promoter is active (*i.e.*, "ON") in a subset of specific cells. Spatially restricted promoters may also be referred to as enhancers, transcriptional control elements, control sequences, *etc.* Any convenient spatially restricted promoter may be used and the choice of suitable promoter (*e.g.*, a brain specific promoter, a promoter that drives expression in a subset of neurons, a promoter that drives expression in the lungs, a promoter that drives expression in the lungs, a promoter that drives expression in muscles, a promoter that drives expression in islet cells of the pancreas, *etc.*) will depend on the organism. For example, various spatially restricted promoters are known for plants, flies, worms, mammals, mice, *etc.* Thus, a spatially restricted promoter can be used to regulate the expression of a nucleic acid encoding a subject Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) or PUF domain fusion in a wide variety of different tissues and cell types, depending on the organism. Some spatially restricted promoters are also temporally restricted such that the promoter is in the "ON" state or "OFF" state during specific stages of embryonic development or during specific stages of a biological process (*e.g.*, hair follicle cycle in mice).

For illustration purposes, examples of spatially restricted promoters include, but are not limited to, neuron-specific promoters, adipocyte-specific promoters, cardiomyocyte-specific promoters, smooth muscle-specific promoters, photoreceptor-specific promoters, *etc*. Neuron-specific spatially restricted promoters include, but are not limited to, a neuron-specific enolase (NSE) promoter (see, *e.g.*, EMBL HSEN02, X51956); an aromatic amino acid decarboxylase (AADC) promoter; a neurofilament promoter (see, *e.g.*, GenBank HUMNFL, L04147); a synapsin promoter (see, *e.g.*, GenBank HUMSYNIB, M55301); a thy-1 promoter (see, *e.g.*, Chen *et al.*, *Cell*, 51:7-19, 1987; and Llewellyn *et al.*, *Nat. Med.*, 16(10): 1161-1166, 2010); a serotonin receptor promoter (see, *e.g.*, GenBank S62283); a tyrosine hydroxylase promoter (TH) (see, *e.g.*, Oh *et al.*, *Gene Ther.*, 16:437, 2009; Sasaoka *et al.*, *Mol. Brain Res.*, 16:274, 1992; Boundy *et al.*, *Neurosci.*, 18:9989, 1998; and Kaneda *et al.*, *Neuron*, 6:583-594, 1991); a GnRH promoter (see, *e.g.*, Radovick *et al.*, *Proc. Natl. Acad. Sci. USA*, 88:3402-3406, 1991); an L7

promoter (see, *e.g.*, Oberdick *et al.*, *Science*, 248:223-226, 1990); a DNMT promoter (see, *e.g.*, Bartge *et al.*, *Proc. Natl. Acad. Sci. USA*, 85:3648-3652, 1988); an enkephalin promoter (see, *e.g.*, Comb *et al.*, *EMBO J.*, 17:3793-3805, 1988); a myelin basic protein (MBP) promoter; a Ca²⁺-calmodulin- dependent protein kinase II-alpha (CamKIIa) promoter (see, *e.g.*, Mayford *et al.*, *Proc. Natl. Acad. Sci. USA*, 93: 13250, 1996; and Casanova *et al.*, *Genesis*, 31:37, 2001); a CMV enhancer / platelet-derived growth factor-β promoter (see, *e.g.*, Liu *et al.*, *Gene Therapy*, 11:52-60, 2004); and the like.

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Adipocyte-specific spatially restricted promoters include, but are not limited to aP2 gene promoter/enhancer, *e.g.*, a region from -5.4 kb to +21 bp of a human aP2 gene (see, *e.g.*, Tozzo *et al.*, *Endocrinol.* 138: 1604, 1997; Ross *et al.*, *Proc. Natl. Acad. Sci. USA*, 87:9590, 1990; and Pavjani *et al.*, *Nat. Med.*, 11:797, 2005); a glucose transporter-4 (GLUT4) promoter (see, *e.g.*, Knight *et al.*, *Proc. Natl. Acad. Sci. USA*, 100: 14725, 2003); a fatty acid translocase (FAT/CD36) promoter (see, *e.g.*, Kuriki *et al.*, *Biol. Pharm. Bull.*, 25: 1476, 2002; and Sato *et al.*, *Biol. Chem.* 277: 15703, 2002); a stearoyl-CoA desaturase-1 (SCD1) promoter (Tabor *et al.*, *Biol. Chem.* 274:20603, 1999); a leptin promoter (see, *e.g.*, Mason *et al.*, *Endocrinol.* 139: 1013, 1998; and Chen *et al.*, *Biochem. Biophys. Res. Comm.*, 262: 187, 1999); an adiponectin promoter (see, *e.g.*, Kita *et al.*, *Biochem. Biophys. Res. Comm.*, 331:484, 2005; and Chakrabarti, *Endocrinol.* 151:2408, 2010); an adipsin promoter (see, *e.g.*, Piatt *et al.*, *Proc. Natl. Acad. Sci. USA*, 86:7490, 1989); a resistin promoter (see, *e.g.*, Seo *et al.*, *Molec. Endocrinol.*, 17: 1522, 2003); and the like.

Cardiomyocyte-specific spatially restricted promoters include, but are not limited to control sequences derived from the following genes: myosin light chain-2, a-myosin heavy chain, AE3, cardiac troponin C, cardiac actin, and the like. Franz *et al.*, *Cardiovasc. Res.*, 35:560-566, 1997; Robbins *et al.*, *Ann. N.Y. Acad. Sci.*, 752:492-505, 1995; Linn *et al.*, *Circ. Res.*, 76:584-591, 1995; Parmacek *et al.*, *Mol. Cell. Biol.*, 14:1870-1885, 1994; Hunter *et al.*, *Hypertension*, 22:608-617, 1993; and Sartorelli *et al.*, *Proc. Natl. Acad. Sci.*, 89:4047-4051, 1992.

Smooth muscle-specific spatially restricted promoters include, but are not limited to an SM22a promoter (see, *e.g.*, Akyurek *et al.*, *Mol. Med.*, 6:983, 2000; and U.S. Patent No. 7,169,874); a smoothelin promoter (see, *e.g.*, WO 2001/018048); an a-smooth muscle actin promoter; and the like. For example, a 0.4 kb region of the SM22a promoter, within which lie two CArG elements, has been shown to mediate vascular smooth muscle cell-specific expression

(see, e.g., Kim et al., Mol. Cell. Biol., 17:2266-2278, 1997; Li et al., J. Cell Biol., 132:849-859, 1996; and Moessler et al., Development, 122:2415-2425, 1996).

Photoreceptor-specific spatially restricted promoters include, but are not limited to, a rhodopsin promoter; a rhodopsin kinase promoter (Young *et al.*, *Ophthalmol. Vis. Sci.*, 44:4076, 2003); a beta phosphodiesterase gene promoter (Nicoud *et al.*, *Gene Med.*, 9: 1015, 2007); a retinitis pigmentosa gene promoter (Nicoud *et al.*, 2007, supra); an interphotoreceptor retinoid-binding protein (IRBP) gene enhancer (Nicoud *et al.* (2007) supra); an IRBP gene promoter (Yokoyama *et al.*, *Exp. Eye Res.*, 55:225, 1992); and the like.

8. Libraries

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The present invention also provides a plurality or library of the subject polynucleotide sequences, or a plurality or library of the vectors encoding the same. The latter may comprise a library of recombinant expression vectors comprising nucleotides encoding the subject polynucleotides.

A subject library can comprise from about 10 individual members to about 10^{12} individual members; *e.g.*, a subject library can comprise from about 10 individual members to about 10^2 individual members, from about 10^3 individual members to about 10^3 individual members, from about 10^3 individual members to about 10^5 individual members to about 10^7 individual members to about 10^7 individual members to about 10^7 individual members to about 10^9 individual members to about 10^9 individual members to about 10^{12} individual members.

In certain embodiments, two of the vectors differ in the encoded polynucleotides in their respective DNA-targeting sequences, Cas9-binding sequences, and/or the copy number, identity (*e.g.*, sequence, or binding specificity), or relative order of the PBS.

For example, in certain embodiments, an "individual member" of a subject library differs from other members of the library in the nucleotide sequence of the DNA-targeting sequence of the subject polynucleotide. Thus, *e.g.*, each individual member of a subject library can comprise the same or substantially the same nucleotide sequence of the Cas9-binding sequence as all other members of the library; and can comprise the same or substantially the same nucleotide sequence of the PBS as all other members of the library; but differs from other members of the library in the nucleotide sequence of the DNA-targeting sequence of the subject polynucleotide. In this way, the library can comprise members that bind to different target polynucleotide sequences that are either on the same target gene or on different target genes.

In a related embodiment, members of the library may differ such that different DNA-targeting sequences are associated with different PBS, such that different target DNA can be independently regulated - *e.g.*, some target genes are transcriptionally activated (and optionally labeled by a first fluorescent color), while others are transcriptionally repressed (and optionally labeled by a second fluorescent color).

In certain other embodiments, an individual member of a subject library differs from other members of the library in the nucleotide sequence of the Cas9-binding sequence of the subject polynucleotide. Thus, *e.g.*, each individual member of a subject library can comprise the same or substantially the same nucleotide sequence of the DNA-targeting sequence as all other members of the library; and can comprise the same or substantially the same nucleotide sequence of the PBS as all other members of the library; but differs from other members of the library in the nucleotide sequence of the Cas9-binding sequence of the subject polynucleotide. In this way, the library can comprise members that bind to different orthogonal Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) from different species, allowing separately and parallelly regulatable systems in the same host cell.

In certain other embodiments, an individual member of a subject library differs from other members of the library in the nucleotide sequence of the PBS of the subject polynucleotide. Thus, *e.g.*, each individual member of a subject library can comprise the same or substantially the same nucleotide sequence of the DNA-targeting sequence as all other members of the library; and can comprise the same or substantially the same nucleotide sequence of the Cas9-binding sequence as all other members of the library; but differs from other members of the library in the nucleotide sequence of the PBS of the subject polynucleotide.

9. Exemplary Utilities

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A method for modulating transcription according to the present invention finds use in a variety of applications, including research applications; diagnostic applications; industrial applications; and treatment applications.

Research applications may include, e.g., determining the effect of reducing or increasing transcription of a target nucleic acid on, e.g., development, metabolism, expression of a downstream gene, and the like.

High through-put genomic analysis can be carried out using a subject transcription modulation method, in which only the DNA-targeting sequence of the subject polynucleotide

needs to be varied, while the Cas9-binding sequence and the PBS can (in some cases) be held constant. A library (e.g., a subject library) comprising a plurality of nucleic acids used in the genomic analysis would include: a promoter operably linked to a subject polynucleotide-encoding nucleotide sequence, where each nucleic acid would include a different DNA-targeting sequence, a common Cas9-binding sequence, and a common PBS. A chip could contain over 5×10^4 unique polynucleotide of the invention.

Applications would include large-scale phenotyping, gene-to-function mapping, and meta-genomic analysis.

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The subject methods disclosed herein can also find use in the field of metabolic engineering. Because transcription levels can be efficiently and predictably controlled by designing an appropriate DNA-targeting RNA, as disclosed herein, the activity of metabolic pathways (*e.g.*, biosynthetic pathways) can be precisely controlled and tuned by controlling the level of specific enzymes (*e.g.*, via increased or decreased transcription) within a metabolic pathway of interest. Metabolic pathways of interest include those used for chemical (fine chemicals, fuel, antibiotics, toxins, agonists, antagonists, *etc.*) and/or drug production.

Biosynthetic pathways of interest include but are not limited to (1) the mevalonate pathway (*e.g.*, HMG-CoA reductase pathway) (converts acetyl-CoA to dimethylallyl pyrophosphate (DMAPP) and isopentenyl pyrophosphate (IPP), which are used for the biosynthesis of a wide variety of biomolecules including terpenoids/isoprenoids), (2) the non-mevalonate pathway (*i.e.*, the "2-C-methyl-D-erythritol 4-phosphate/l-deoxy-D-xylulose 5-phosphate pathway" or "MEP/DOXP pathway" or "DXP pathway") (also produces DMAPP and IPP, instead by converting pyruvate and glyceraldehyde 3 -phosphate into DMAPP and IPP via an alternative pathway to the mevalonate pathway), (3) the polyketide synthesis pathway (produces a variety of polyketides via a variety of polyketide synthase enzymes. Polyketides include naturally occurring small molecules used for chemotherapy (e. g., tetracyclin, and macrolides) and industrially important polyketides include rapamycin (immunosuppressant), erythromycin (antibiotic), lovastatin (anticholesterol drug), and epothilone B (anticancer drug)), (4) fatty acid synthesis pathways, (5) the DAHP (3-deoxy-D-arabino-heptulosonate 7-phosphate) synthesis pathway, (6) pathways that produce potential biofuels (such as short-chain alcohols and alkane, fatty acid methyl esters and fatty alcohols, isoprenoids, *etc.*), *etc.*

The methods disclosed herein can also be used to design integrated networks (*i.e.*, a cascade or cascades) of control. For example, a subject polynucleotide / Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) / PUF domain fusion may be used to control (*i.e.*, modulate, *e.g.*,

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increase, decrease) the expression of another polynucleotide / Cas9 protein (e.g., wt, nickase, or dCas9 protein) / PUF domain fusion. For example, a first subject polynucleotide may be designed to target the modulation of transcription of a second Cas9 protein (e.g., wt, nickase, or dCas9 protein) or PUF domain fusion with a function that is different than the first PUF domain fusion (e.g., methyltransferase activity, demethylase activity, acetyltansferase activity, deacetylase activity, etc.). In addition, because different Cas9 proteins (e.g., wt, nickase, or dCas9 protein) (e.g., derived from different species) may require a different Cas9 handle (i.e., Cas9-binding sequence), the second Cas9 protein (e.g., wt, nickase, or dCas9 protein) can be derived from a different species than the first Cas9 protein (e.g., wt, nickase, or dCas9 protein) above. Thus, in some cases, the second Cas9 protein (e.g., wt, nickase, or dCas9 protein) can be selected such that it may not interact with the first subject polynucleotide. In other cases, the second Cas9 protein (e.g., wt, nickase, or dCas9 protein) can be selected such that it does interact with the first subject polynucleotide. In some such cases, the activities of the two (or more) Cas9 proteins (e.g., wt, nickase, or dCas9 protein) / PUF domain fusions may compete (e.g., if the polypeptides have opposing activities) or may synergize (e.g., if the polypeptides have similar or synergistic activities). Likewise, as noted above, any of the complexes (i.e., polynucleotide / Cas9 protein (e.g., wt, nickase, or dCas9 protein) / PUF domain fusion) in the network can be designed to control other polynucleotide / Cas9 protein (e.g., wt, nickase, or dCas9 protein) / PUF domain fusion. Because a subject polynucleotide / Cas9 protein (e.g., wt, nickase, or dCas9 protein) / PUF domain fusion can be targeted to any desired DNA sequence, the methods described herein can be used to control and regulate the expression of any desired target. The integrated networks (i.e., cascades of interactions) that can be designed range from very simple to very complex, and are without limit.

In a network wherein two or more components (*e.g.*, polynucleotide / Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) / PUF domain fusion) are each under regulatory control of another polynucleotide / Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) / PUF domain fusion complex, the level of expression of one component of the network may affect the level of expression (*e.g.*, may increase or decrease the expression) of another component of the network. Through this mechanism, the expression of one component may affect the expression of a different component in the same network, and the network may include a mix of components that increase the expression of other components, as well as components that decrease the expression of other components. As would be readily understood by one of skill in the art, the above examples whereby the level of expression of one component may affect the level of

expression of one or more different component(s) are for illustrative purposes, and are not limiting. An additional layer of complexity may be optionally introduced into a network when one or more components are modified (as described above) to be manipulatable (*i.e.*, under experimental control, *e.g.*, temperature control; drug control, *i.e.*, drug inducible control; light control; *etc.*).

As one non-limiting example, a first subject polynucleotide can bind to the promoter of a second subject polynucleotide, which controls the expression of a target therapeutic / metabolic gene. In such a case, conditional expression of the first subject polynucleotide indirectly activates the therapeutic / metabolic gene. RNA cascades of this type are useful, for example, for easily converting a repressor into an activator, and can be used to control the logics or dynamics of expression of a target gene.

A subject transcription modulation method can also be used for drug discovery and target validation.

10. Kits

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The present invention also provides a kit for carrying out a subject method. A subject kit may comprise: a) a polynucleotide of the present invention, or a nucleic acid (*e.g.*, vector) comprising a nucleotide sequence encoding the same; optionally, b) a subject Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), or a vector encoding the same (including an expressible mRNA encoding the same); and optionally, c) one or more subject PUF domain fusion each comprising a PUF domain fused to an effector domain that may be the same or different among the different PUF domain fusions, or a vector encoding the same (including an expressible mRNA encoding the same).

In certain embodiments, one or more of a) - c) may be encoded by the same vector.

In certain embodiments, the kit also comprises one or more buffers or reagents that facilitate the introduction of any one of a) - c) into a host cell, such as reagents for transformation, transfection, or infection.

For example, a subject kit can further include one or more additional reagents, where such additional reagents can be selected from: a buffer; a wash buffer; a control reagent; a control expression vector or RNA polynucleotide; a reagent for *in vitro* production of the wt or dCas9 or PUF domain fusion from DNA; and the like.

Components of a subject kit can be in separate containers; or can be combined in a single container.

In addition to above-mentioned components, a subject kit can further include instructions for using the components of the kit to practice the subject methods. The instructions for practicing the subject methods are generally recorded on a suitable recording medium. For example, the instructions may be printed on a substrate, such as paper or plastic, *etc*. As such, the instructions may be present in the kits as a package insert, in the labeling of the container of the kit or components thereof (*i.e.*, associated with the packaging or subpackaging) *etc*. In other embodiments, the instructions are present as an electronic storage data file present on a suitable computer readable storage medium, *e.g.* CD-ROM, diskette, flash drive, *etc*. In yet other embodiments, the actual instructions are not present in the kit, but means for obtaining the instructions from a remote source, *e.g.* via the internet, are provided. An example of this embodiment is a kit that includes a web address where the instructions can be viewed and/or from which the instructions can be downloaded. As with the instructions, this means for obtaining the instructions is recorded on a suitable substrate.

15 EXAMPLES

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Example 1 sgRNA Scaffold Remains Functional with Insertion of 47 Copies of Engineered Pumilio Binding Sites

This example demonstrates that the subject 3-component CRISPR/Cas complex / system can have at least 47 copies of the engineered 8-mer Pumilio homologue domain-binding sequences (PBSs) at the 3' end of sgRNA, without substantially affecting the function of the dCas9/sgRNA complex.

In particular, to test whether appending PBS to the 3' end of sgRNA affects sgRNA function, a series of modified Tet-targeting (sgTetO) or non-targeting control (sgControl) sgRNA were generated, with 0 copy, 5 copies, 15 copies, 25 copies, and 47 copies of the 8-mer Pumilio homologue domain-binding sequence (PBS) for PUF (3-2) (also simply referred to as PUFa) [PBS32 or PBSa: 5'-UGUAUgUA-3'], PUF(6-2/7-2) (also simply referred to as PUFb) [PBS6272 or PBSb: 5'-UugAUAUA-3']. See FIG. 1A. The ability of these constructs to direct the dCas9-VP64 transcriptional activator to activate tdTomato expression in a HEK293T/TetO::tdTomato cell line was tested.

Cells were transfected with dCas9-VP64 with the different sgRNA scaffolds, and were analyzed by fluorescent-activated cell sorting (FACS) two days after transfection (FIG. 1B). All the control non-targeting sgRNAs did not activate tdTomato expression. Meanwhile, all the Tet-

targeting sgRNAs with different number of PBS could direct dCas9-VP64 to activate tdTomato expression, showing that insertion of at least 47 copies of 8-mer sites do not substantially impact the activity of sgRNA in directing dCas9-VP64 to its targets (FIG. 1C).

Under the test condition, and for both PUFa-VP64/PBSa and PUFb-VP64/PBSb, 5-10 copies of PBS appended to the sgRNA were best able to activate the target transgene.

Meanwhile, 15, 20, and 47 copies of PBS led to slightly lower, albeit still substantial transgene activation (FIG. 1C).

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Example 2 The Subject 3-Component CRISPR/Cas Complexes / Systems are Orthogonal to Each Other Due to the Specificity of the Engineered Pumilio with the Cognate 8-mer Binding Sites

This example demonstrates that specificity between the differently programmed PUF domains and their corresponding sgRNA with their cognate 8-mer motifs provide independence or orthogonality between each of the subject 3-component CRISPR/Cas complex / system.

Fusions of PUF(3-2)::VP64 and PUF(6-2/7-2)::VP64, which interacts with sgRNA (sgRNA-PBS32) with 5'-UGUAUgUA-3' binding sites and sgRNA-PBS6272 with 5'-UugAUAUA-3' binding sites, respectively, were created, and their activity to turn on tdTomato expression in conjunction with dCas9 was tested. In addition, two additional pairs, PUFw-VP64 recognizing PBSw (5'-UGUAUAUA-3') and PUFc-VP64 recognizing PBSc (5'-UugAUgUA-3'), were also constructed to test their ability to activate the same TetO::tdTomato expression in conjunction with dCas9 (FIG. 1D).

As shown in FIG. 1D, PUF::VP64 can activate tdTomato expression only when the sgRNA with the cognate binding sites were provided. This demonstrates that the subject 3-component CRISPR/Cas complex / system provides independence or orthogonality of effector function based on the pairing of PUF domains and their 8-mer binding sites on the sgRNA-PBS. Impressively, although PBSa and PBSw binding sites only differ by one nucleotide, their gene activation remains target-specific, demonstrating the high specificity of the subject 3-component CRISPR/Cas complex / system.

Example 3 The Subject 3-Component CRISPR/Cas Complex / System allows Assembly of Protein Complex at Target Loci

This example demonstrates that protein complexes with two or more different protein components can be assembled on sgRNA and operate at defined loci using the subject system.

Specifically, p65-HSF1 has recently been shown to be a potent activator domain. An sgRNA with both PBS32 and PBS6272 positioned next to each other, and PUF(3-2)::VP64 and PUF(6-2/7-2)::p65-HSF1 fusions that would occupy the two different sites, were generated (FIG. 2A). Co-transfection of both PUF(3-2)::VP64 and PUF(6-2/7-2)::p65-HSF1 induced a tdTomato fluorescence, with an intensity about the sum of the fluorescent intensity resulting from transfecting the single activators alone. This indicates that sgRNA with binding sites for both PUF(3-2) and PUF(6-2/7-2) allows both fusion proteins of both types to assemble on the targeted genomic locus.

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A recent paper has tested both VP64 and p65HSF1 as transcriptional activation domains, and found p65HSF1 to be a more potent activator. To directly compare these two transcriptional activation domains, p65HSF1 PUF fusion (PUFa-p65HSF1) and VP64 PUF fusion (PUFa-VP64) were used to activate the TetO::tdTomato transgene using sgRNA with different number of PBSa (FIG. 2C). PUFa-p65HSF1 provided up to 3 times more activation as did PUFa-VP64. Activation was observed even with only one PBSa (previously not observed with PUFa-VP64 module). Thus p65HSF1 is confirmed to be a more potent transcriptional activation domain than VP64.

Example 4 The Subject 3-Component CRISPR/Cas Complex / System can Activate Endogenous Genes More Efficiently than dCas9 Direct Fusion with Activator

We previously used a cocktail of 3-4 sgRNAs per gene to achieve robust endogenous gene activation of *OCT4* and *SOX2* using a dCas9-VP160 direct fusion while single sgRNAs failed to induce much activation (data not shown).

This example demonstrates that recruitment of multiple molecules of activator domains via multiple PBS on the sgRNA-PBS in the subject system increases transactivation activity, thus allowing the use of fewer sgRNAs to achieve endogenous gene activation.

Specifically, activation of endogenous genes *OCT4* and *SOX2* in HEK293T were compared using the subject system with a direct dCas9-p65HSF1 activator using either a cocktail of four sgRNA-PBS per gene, or individual sgRNA-PBS (FIGs. 3A and 3B). Higher activation was observed using the subject 3-Component CRISPR/Cas Complex / System compared to direct fusion in the mixed sgRNA-PBS cocktail, as well as in single guide experiments in both *OCT4* and *SOX2* activation experiments (FIGs. 3A and 3B). Little to no activation by single guide targeting of direct fusion dCas9-p65HSF1 to *OCT4* and *SOX2* was observed, while robust activation was observed in the corresponding 3-component system

experiments, showing the superior activity of the subject 3-Component CRISPR/Cas Complex / System activator over the direct fusion.

To determine the optimal number of PBSa sites on the sgRNA for *OCT4* and *SOX2* activation, sgRNA-PBS targeting either *OCT4* or *SOX2* proximal promoter with 1, 5, 15 or 25 copies of PBSa were constructed. In both *OCT4* and *SOX2* experiments, we observed highest activation using 5×PBSa, in either sgRNA-5×PBSa cocktail experiments and single sgRNA-5×PBSa experiments, recapitulating the finding in the TetO::tdTomato reporter experiments (FIGs. 3D and 3E).

Example 5 The Subject 3-Component CRISPR/Cas Complex / System Allows Simultaneous Activation and Repression of Target Genes

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This example demonstrates that different effector functions can be assigned to each of the subject 3-component CRISPR/Cas complex / system.

The KRAB::PUF(6-2/7-2) repression fusion and the sgRNA targeting SV40 promoter were first generated. A HEK293T reporter cell line having a tdTomato reporter under the control of the TetO promoter, and an EGFP reporter under the control of the SV40 promoter (HEK293T / TetO::tdTomato / SV40::EGFP) was then used to test simultaneous (1) activation of tdTomato via dCas9/sgTetO-PBS32/PUF(3-2)::VP64 binding to TetO promoter, and (2) repression of EGFP expression via binding of dCas9/sgSV40-PBS6272/KRAB::PUF(6-2/7-2) at the SV40 promoter (FIG. 4A). Expression of the 3-component CRISPR/Cas activator complex consisting of dCas9, sgTetO-5×PBS32 and PUF(3-2)::VP64 activated tdTomato fluorescence (FIG 4B; sample 2) while expression of the 3-component CRISPR/Cas repressor complex consisting of dCas9, sgSV40-5×PBS6272 reduced EGFP fluorescence (FIG4B; sample 4). Co-expression of both activator and repressor complexes induced simultaneous activation of the tdTomato and repression of the EGFP transgene, respectively (FIG4B, sample 6), demonstrating that the subject 3-component CRISPR/Cas complexes with different effector functions can operate within the same cell and produce different output at their targets.

To further confirm the versatility of the subject system in recruiting various effectors, a KRAB-PUFa repressor fusion and as well as a PUFc-p65HSF1 activator fusion were constructed. In a reporter cell line HEK293T/TetO::tdTomato/SV40::EGFP, the TetO::tdTomato reporter gene can be efficiently activated by dCas9/PUFc-p65HSF1/sgTetO-PBSc, while SV40::EGFP expression is significantly repressed by dCas9/KRAB-PUFa /sgSV40-PBSa (FIG. 4C). When both systems were applied, simultaneous activation of

TetO::tdTomato and repression of SV40::EGFP expression were achieved (FIG. 4C). When non-targeting (sgCtl) sgRNA were used, or when the PUF fusions were omitted, the fluorescent levels of the respective reporters were not affected, showing that the effects on the reporters are specific and are due to the action of the effectors recruited by the cognate dCas9/sgRNA-PBS at the targets.

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Next, it was tested whether the expression of multiple endogenous genes can be independently regulated using this strategy. The subject 3-component modules were directed to endogenous target genes by changing the targeting sequence of sgRNA-PBSb and sgRNA-PBSa so that PUFb-p65HSF1 was recruited to the *OCT4* promoter and BFPKRAB-PUFa to the *SOX2* promoter. Similar to the results from reporter gene experiments, effector-mediated simultaneous as well as independent activation of *OCT4* and repression of *SOX2* were achieved (FIG. 4D).

Example 6 Recruitment of Histone Acetyltransferase (HAT) Domain by the Subject 3-Component CRISPR/Cas Complex / System Achieves Enhancer Activation

Artificial transcription factor systems can be used to recruit epigenetic modifiers to activate or repress genes. Recent experiments have used histone acetyltransferase (HAT) to activate enhancers. To demonstrate that the subject 3-component system can recruit multiple molecules of HAT domain to increase the efficiency of epigenetic editing, *OCT4* was used as a model gene since its enhancers and the promoter are well defined, and the choice of enhancer usage is of biological significance corresponding to the embryonic stem cell states.

In this experiment, the Proximal Promoter (PP), Proximal Enhancer (PE) and Distal Enhancer (DE) were targeted, each with four different sgRNA-PBS (FIG. 5A). Direct fusion between HAT from CREB-binding protein (CBP) and the C-terminus of dCas9 (dCas9::CBPHAT) was constructed, so were an N-terminal fusion module CBPHAT::PUFa, and a C-terminal fusion module PUFa::CBPHAT. Their activity in activating *OCT4* expression via binding to PP, PE and DE were then tested.

As shown in FIG. 5B, dCas9::CBPHAT and CBPHAT::PUFa have similar activity at proximal promoter (PP). Interestingly, when coupled with sgRNA with 5×PBSa, the subject 3-component modules have higher efficiency activating *OCT4* gene via both enhancers PE and DE, with N-terminal fusion CBPHAT::PUFa giving the highest activation. Next, it was analyzed the activity of CBPHAT::PUFa directed by single sgRNA-5×PBSa to PP, PE and DE by sgRNA-5×PBSa (FIG. 5C). Although with smaller fold changes than using cocktails of 4 sgRNA-5×PBSa, single sgRNA-5×PBSa were able to activate the expression of *OCT4* gene

through targeting of these elements (FIG. 5C).

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Example 7 The Subject 3-Component CRISPR/Cas Complex / System Allows Fluorescent Tagging of Telomeres

In addition to transcriptional regulation, another important application of dCas9-effector is to label genomic loci for live cell imaging. This example demonstrates that the subject 3-component CRISPR/Cas complex / system can be used for fluorescent tagging of chromosomal loci, such as labeling of telomeres.

We appended sgRNA designed to target telomeres (sgTelomere) with 0, 5, 15, or 25 copies of PBSa to recruit fluorescent proteins fused to a PUFa domain (FIG. 6A). While expression of sgTelomere-5×PBSa, 15×PBSa and 25×PBSa with dCas9 and Clover::PUFa produced green fluorescent foci consistent with telomere labeling, expression of sgRNA harboring no PBSa site did not produce any foci (FIG. 6B). To confirm that subject 3-component system-directed fluorescent signal is indeed localized at telomeres, co-labeling experiment with antibody against telomeric repeat binding factor TRF2 was performed. The 3-component system telomere signals largely overlapped with the TRF labeling (FIG. 6C), indicating highly specific labeling of telomeres by sgRNA appended with PBSa sites that recruit Clover-PUFa.

Interestingly, the strength of telomere labeling increased as more copies of PBS were appended to the Telomere-sgRNAs (FIG. 6B). Quantification of foci number and signal-to-noise (%GFP in foci/total GFP in nucleus) showed progressive increase from experiment using sgRNA with 5, 15 to 25× PBSa (FIGs. 6D and 6E), indicating the multimerization feature of the subject 3-component system allows for titration of labeling intensity at target loci.

Example 8 The Subject 3-Component CRISPR/Cas Complex / System Allows Simultaneous Fluorescent Tagging of Telomeres and Centromeres

This example demonstrates that the subject 3-component CRISPR/Cas complex / system can label more than one (e.g., two) genomic loci simultaneously in the same cells by using the multiplexing feature.

To further demonstrate the ability of the subject 3-component system to label two genomic loci simultaneously, an sgRNA was designed to target centromeres with appended binding sites for PUFc (sgCentromere-20×PBSc). Labeling of centromeres by the subject 3-component system and immunostaining using anti-CREST antibody were observed and confirmed (FIG. 6F). When Clover-PUFb/sgCentromere-20×PBSc, Ruby-PUFa/sgTelomere-

25×PBSa and dCas9 were co-introduced into HEK293T cells, independent labeling of both centromeres and telomeres in the same cells were observed (FIG. 6G), demonstrating that the subject 3-component system can be used to independently label multiple genomic loci.

Example 9 The Subject 3-Component CRISPR/Cas Complex / System Allows Fluorescent Tagging of Non-repeat Chromosomal Loci

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A previous study using dCas9::GFP to label non-repetitive DNA reported the requirement of >32 targeting events to concentrate enough signal to label such non-repeat regions. This example demonstrates that, by incorporating multiple binding sites for PUF-fluorescent protein fusions, fluorescent signals can be concentrated at a target site, thus reducing the amount of targeting sites needed for detection of non-repeat DNA.

The non-repeat region at the MUC4 locus was tested in this example. Seven (7) sgRNAs each harboring 15×PBS32, Clover::PUF(3-2) and dCas9, labeling pattern reminiscent of that of MUC4 labeling was successfully detected (FIG. 7). This demonstrates that the subject 3-component CRISPR/Cas complex / system can be used to "polymerize" proteins at defined genomic loci, which enables and greatly expands the application of the subject 3-component CRISPR/Cas complex / system in the field of imaging.

The above examples demonstrate the ability of the subject 3-component CRISPR/Cas complex / system to achieve multiplexing (FIG. 8A), complex formation (FIG. 8C), and polymerization of proteins (FIG. 8B), including transcriptional regulators, epigenetic modifiers, and fluorescent proteins, and the system can independently direct them to defined genomic loci. This enables construction of complex molecular behavior at multiple loci, and allows studying and reconstitution of protein complexes with defined stoichiometry. The polymerization feature of the subject 3-component CRISPR/Cas complex / system allows concentration of enzymatic activity or other proteins to defined genomic loci, to increase the effect of the enzymatic activity or to concentrate signal enrichment for applications like chromosomal imaging.

More specifically, some main advantages of the subject 3-component system include: (A) *Multiplexing*. Different modules of the subject 3-component system can be simultaneously delivered into a cell and each can operate at their defined target sites with independence (i.e., without interference with other modules and their target sites). Since PUF domains can be easily programmed to recognize any 8-mer RNA motifs, this expands the potential number of independent modules to a theoretical maximum of 4^8 (65536). By inserting a PUF array within

another, the recognition site can be programmed to a 16-mer RNA motif, with a sequence space of 4¹⁶ (4.29 billion). (B) Multimerization: Simplicity of the linear 8-mer PBS motif allows extensive multimerization of PUF fusions on sgRNA-PBS without hindering sgRNA transcription or Cas9/sgRNA DNA binding activity. This feature allows multiple molecules of PUF fusions to be assembled on the sgRNA, allowing for localized concentration of effectors or protein tags. This is particularly beneficial for fluorescent imaging or transcriptional regulation. As shown with the above experiments labeling repeat sequences such as telomeres, sgRNA-PBS with more PBS increases signal at the telomeric foci. This feature may facilitate labeling of nonrepeat sequences where usually tiling of more than 30 sgRNAs were required. Higher efficiency of HAT-mediated enhancer activation using the subject system versus direct dCas9-HAT fusion was observed. It is contemplated that multimerization can facilitate spreading of the epigenetic modification directed by the artificial epigenetic factors useful for reprogramming of large epigenetic domains such as super-enhancers or imprinted loci. (C) Stoichiometrically defined Complex formation: although not directly tested here, it is contemplated that the sgRNA-PBS can act as RNA scaffold for PUF-directed assembly of Stoichiometrically defined protein complexes. Specifically, Varying numbers of PBS copies with varying specificities can be appended to the sgRNA to allow for multiprotein complex formation with defined stoichiometry, as well as with defined ordering along the sgRNA-PBS.

The materials and methods used in the examples above are compiled below.

20 Cloning

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A list of vectors, links to their Addgene entries are provided in Table S1 below. Detailed description of cloning strategies and sequences are given below.

PUFa [PUF(3-2)] and PUFb [PUF(6-2/7-2)] with N-terminal NLS were amplified from constructs containing these coding sequences with primers containing *SgrAI* and *PacI* sites and were used to replace *SgrAI*-dCas9-*FseI* from pAC164:pmax-dCas9Master_VP64 to create pAC1355:pmax-NLSPUFa_VP64 and pAC1356:pmax-NLSPUFb_VP64. A fusion PCR with 5' fragment up to repeat 4 of NLSPUFb and 3' fragment from repeat 5 to the end of NLSPUFa was used to create pAC1357:pmax-NLSPUFw_VP64. A fusion PCR of 5' fragment of NLSPUFa with 3' fragment of NLSPUb was used to create pAC1358:pmax-NLSPUFc_VP64.

p65HSF1 activator ORF was amplified from MS2-P65-HSF1_GFP (Addgene: 61423) with *FseI PacI* sites to replace VP64 fragment in pAC164 to create pAC1410:pmax-dCas9_p65HSF1, and replace VP64 in pAC1355 and pAC1358 to create pAC1393: pmax-

NLSPUFa_p65HSF1 and pAC1411:pmax-NLSPUFc_p65HSF1, respectively.

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Clover and mRuby2 were amplified from pcDNA3-Clover (Addgene #40259) and pcDNA3-mRuby2 (Addgene #40260) respectively with primers containing *SgrAI* and *FseI* cloning site, ligated with various *FseI*-PUF-*PacI* amplified from the above pAC1356~1358 and vector digested from pAC149:pCR8-dCas9VP160 (Addgene #48221) to create gateway donor vectors pAC1402, pAC1403 and pAC1404 containing ORFs of Clover_PUFa and Clover_PUFc, mRuby2_PUFa, respectively. These ORFs are then transferred to PB3-neo vector by recombining with pAC1119:PB3-neo(-)-pmaxDEST(+) by LR Clonase (Invitrogen) to create expression vectors pAC1360 (Clover_PUFa), pAC1381 (Clover_PUFc) and pAC1362 (mRuby2_PUFa).

NLSKRAB repressor domain was amplified from SOX2 TALE Repressor (KRAB 1-75) (Addgene #42945) with primers containing *AgeI-ClaI* sites and ligated with NLSPUFa amplified with primers containing *AcII PacI* and with pAC1360 digested with *SgrAI-PacI* as vector to create pAC1412: PB3-neo(-)-pmax-NLSKRAB NLSPUFa.

The *FseI*-p65HSF1-*PacI* fragment was released from pAC1393 and ligated with *SgrAI*-NLSPUMb fragment released from pAC1356 and pAC1360 digested with *SgrAI*-PacI as vector to create pAC1413: PB3-neo(-)-pmax-NLSPUFb_p65HSF1. The BFPKRAB fragment was amplified from pHR-SFFV-dCas9-BFP-KRAB (Addgene #46911) and was used to replace Clover fragment from pAC1360 to create pAC1414: PB3-neo(-)-pmax-BFPKRAB_NLSPUFa. Then, an *NheI*-CAGGS-NLSPUFb_p65HSF1-*NheI* fragment was amplified from pAC1413 and inserted into pAC1414 digested with *NheI* to create a dual expression vector for BFPKRAB-NLSPUFa and NLSPUFb-p65HSF1 (pAC1414: PB3-NLSPUFb_p65HSF1(-)neo(-)-BFPKRAB2_NLSPUFa).

Four gateway donor vectors with improved linker sequences and three extra NLS on the N-terminal and one additional NLS on the C-terminal of PUF as well as cloning sites for N-terminal (*SgrAI,ClaI*) and C-terminal (*FseI-PacI*) insertions were created (pAC1404~1408). HAT sequence was amplified from mouse Crebbp gene using mouse cDNA with primers containing *FseI-PacI* site and inserted into pAC164 to create pAC1364: pmax-dCas9Master_CBPHAT and into pAC1405 to create pAC1415: pCR8-4×NLSPUFa_2×NLS_CBPHAT. HAT sequence was amplified with another pair of primers containing *SgrAI-AcII* site and cloned into *SgrAI-ClaI* site of pAC1405 to create pAC1416: pCR8-CBPHAT_4×NLSPUFa_2×NLS. pAC1415 and pAC1416 were recombined into pAC90:pmax-DEST (Addgene #48222) to create expression vectors pAC1417: pmax-

4×NLSPUFa_2×NLS_CBPHAT and pAC1418: pmax-CBPHAT_4×NLSPUFa_2×NLS, respectively. *FseI*-mCherry-*PacI* fragment was amplified from a plasmid containing mCherry sequence and ligated with *SgrAI*-dCas9-*FseI* to PB3-neo(-)-pmax to generate pAC1419: PB3-neo(-)-pmax-dCas9Master_mCherry.

Expression vectors for sgRNA-PBS were constructed as follows: First, a sgRNA scaffold based on sgF+E with *BbsI* for oligo cloning of guide sequence and with 3' *BsaI* (right upstream of the terminator) for insertion of PBS were ordered as a gBlock (IDT), and were cloned into pX330 (Addgene #42230) replacing the *AfIIII-NotI* region to create vector pAC1394: pX-sgFE-BsaI(AGAT). Then, oligos encoding 5×PBSa sites each separated by ggc-spacer flanked by 5'-AGAT-3' overhangs on one side and 5'-ATCT-3' on the other side were treated with T4PNK and annealed and ligated into pAC1394 digested with *BsaI* (to create compatible overhangs). Clones were then screened for 1 copy (5×PBS), 2 copies (10×PBS), etc of the oligo insertions for the different number of PBS. For 1×PBS and 2×PBS vectors, they were constructed using oligo containing one PBS site. Guide sequence for each target were then cloned onto the sgRNA-PBS expression vectors via *BbsI* site as previously described. For sgRNA expression vectors with GFP expression markers, they were constructed by transferring the sgRNA-PBS expression cassette from the pX vectors onto a PB-GFP vector via AscI site. The different sgRNA expression constructs are listed in Table S1.

Cell Culture for Experiments

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HEK293T cells were cultivated in Dulbecco's modified Eagle's medium (DMEM)(Sigma) with 10% fetal bovine serum (FBS)(Lonza), 4% Glutamax (Gibco), 1% Sodium Pyruvate (Gibco) and penicillin-streptomycin (Gibco). Incubator conditions were 37°C and 5% CO₂. For activation experiments, cells were seeded into 12-well plates at 100,000 cells per well the day before being transfected with 200 ng of dCas9 construct, 100 ng of modified sgRNA and 100 ng of PUF-fusion with Attractene transfection reagent (Qiagen). After transfection, cells were grown for 48 hrs and harvested for either RNA extraction or fluorescent-activated cell sorting (FACS). For dual activation-repression experiments, transfection remained the same, however cells were seeded into 12-well plates at 150,000 cells per well and were grown for 72 hrs before being harvested for FACS. For experiments with *OCT4* and *SOX2* dual activation-repression, cells were triple-sorted by BFP (for the activator-repressor module PUFb-p65HSF1/BFPKRAB-PUFa), mCherry (for dCas9mCherry) and GFP (for the sgRNA-PBS on vectors co-expressing EGFP) before RNA extraction. For imaging experiments, cells were seeded into 6-well plates with 22×22×1 microscope cover glass at 300,000 cells per well the day

before being transfected with 50 ng of dCas9 construct, 500 ng of modified sgRNA, and 50 ng of a PUF-fluorescent fusion with Attractene transfection reagent. After transfection, cells were grown for 48 hrs then immunostained.

Quantitative RT-PCR Analysis

Cells were harvested with trypsin, washed with Dulbecco's phosphate-buffered saline (dPBS), centrifuged at 125 g for 5 mins and then RNA was extracted using RNeasy Plus Mini Kit (Qiagen). A cDNA library was made using Applied Biosystems High Capacity RNA-to-cDNA kit with 1 μg of RNA. TaqMan Gene expression assays (Applied Biosystems) were designed using GAPDH (Hs03929097, VIC) as endogenous control and OCT4 (Hs00999632, FAM) and SOX2 (Hs01053049, FAM) as targets. TaqMan Universal Master Mix II, with UNG (Applied Biosystems) was used for Quantitative PCR (qPCR), with 2 μl of 1:10 diluted cDNA used for each reaction. Activation was analyzed with the Applied Biosystems ViiA7 instrument. Gene expression levels were calculated by "delta delta Ct" algorithm and normalized to control samples.

15 Fluorescent-Activated Cell Sorting

Cells were trypisinized and fixed for 10 min with 2% paraformaldehyde. Afterwards, the cells were centrifuged at 125 g for 5 min and resuspended in dPBS. Samples were analyzed on a FACScalibur flow cytometer using CellQuest Pro software (BD Bioscience). thousands events were collected in each run.

20 Immunostaining and Microscopy

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While adherent to a cover glass, cells were fixed in 2% paraformaldehyde, washed with 0.1% Triton X-100 in dPBS, permeabilized with 0.4% Triton X-100 in dPBS for 5 min at 4°C, blocked in 5% Blotting-grade blocking buffer (BIO-RAD) for 30 min, incubated with the primary antibody in blocking buffer at 4°C overnight, washed three times with dPBS, then incubated in the dark with a respective Alexa Fluor-conjugated secondary antibody at room temperature for 3 hours, washed again, and stained with DAPI. The cover glass was mounted on a slide with glycerol before imaging. Immunostaining of telomeres was performed with a 1:100 dilution of an anti-TRF2 primary antibody (Novus Biologicals: NB110-57130) and a 1:500 dilution of an Alexa fluor 594-conjugated anti-Rabbit IgG secondary antibody (Invitrogen, A11037). A 1:100 dilution of CREST antibody (Antibodies Incorporated: 15-235-0001) was used in conjunction with a 1:500 dilution of an Alexa fluor 594-conjugated anti-Human IgG

secondary antibody (Invitrogen, A11014) to detect centromeres.

DALDDFDLDMLGSDALDDFDLDMLGSDALDDFDLDMLYID

Sequences of some of the constructs used in the examples above and the related sequences are listed herein below.

>NLSPUFa_VP64 Key: NLS PUFa VP64

5 MGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATPAE RQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIEKA LEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPY GCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGNYVIQHVLEHGRPEDKSKIVAEIRGNV LVLSQHKFASNVVEKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAE PGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLGGPAGSGR**ADALDDFDLDMLGS**

In the above sequence, the NLS sequence is residues 6-12, PUFa is residues 15-363, and VP64 is residues 371-421.

>NLSPUFb VP64 Key: NLS PUFb VP64

15 MGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATPAE RQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGCRVIQKA LEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPY GCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGSYVIEHVLEHGRPEDKSKIVAEIRGNV LVLSQHKFANNVVQKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAE PGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLGGPAGSGR**ADALDDFDLDMLGS DALDDFDLDMLGSDALDDFDLDML**YID

In the above sequence, the NLS sequence is residues 6-12, PUFb is residues 15-363, and VP64 is residues 371-421.

>NLSPUFw VP64 Key: NLS PUFw VP64

25 MGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATPAE RQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGCRVIQKA LEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPY GCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGNYVIQHVLEHGRPEDKSKIVAEIRGNV LVLSQHKFASNVVEKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAE PGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLGGPAGSGRADALDDFDLDMLGS DALDDFDLDMLGSDALDDFDLDMLGSDALDDFDLDMLYID

In the above sequence, the NLS sequence is residues 6-12, PUFw is residues 15-363, and VP64 is residues 371-421.

>NLSPUFc VP64 Key: NLS PUFc VP64

- 35 MGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATPAE RQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIEKA LEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPY GCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGSYVIEHVLEHGRPEDKSKIVAEIRGNV LVLSQHKFANNVVQKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAE
- 40 PGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLGGPAGSGR**ADALDDFDLDMLGS**DALDDFDLDMLGSDALDDFDLDMLGSDALDDFDLDMLYID

In the above sequence, the NLS sequence is residues 6-12, PUFc is residues 15-363, and VP64 is residues 371-421.

>Clover_NLSPUFa Key: Clover NLS PUFa

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MVSKGEELFTGVVPILVELDGDVNGHKFSVRGEGEGDATNGKLTLKFICTTGKLPVPWPTLVTT
FGYGVACFSRYPDHMKQHDFFKSAMPEGYVQERTISFKDDGTYKTRAEVKFEGDTLVNRIELKG
IDFKEDGNILGHKLEYNFNSHNVYITADKQKNGIKANFKIRHNVEDGSVQLADHYQQNTPIGDG
PVLLPDNHYLSHQSALSKDPNEKRDHMVLLEFVTAAGITHGMDELYKSRGPYSIVSPKCGGGGS
GPAGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATP
AERQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIE
KALEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTH
PYGCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGNYVIQHVLEHGRPEDKSKIVAEIRG
NVLVLSQHKFASNVVEKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDV
AEPGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLG

In the above sequence, the NLS sequence is residues 264-270, PUFa is residues 273-621, and Clover is residues 1-251.

>Clover_NLSPUFc Key: Clover NLS PUFc

MVSKGEELFTGVVPILVELDGDVNGHKFSVRGEGEGDATNGKLTLKFICTTGKLPVPWPTLVTT
FGYGVACFSRYPDHMKQHDFFKSAMPEGYVQERTISFKDDGTYKTRAEVKFEGDTLVNRIELKG
IDFKEDGNILGHKLEYNFNSHNVYITADKQKNGIKANFKIRHNVEDGSVQLADHYQQNTPIGDG
PVLLPDNHYLSHQSALSKDPNEKRDHMVLLEFVTAAGITHGMDELYKSRGPYSIVSPKCGGGGS
GPAGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATP
AERQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIE
KALEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTH
PYGCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGSYVIEHVLEHGRPEDKSKIVAEIRG
NVLVLSQHKFANNVVQKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDV
AEPGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLG

In the above sequence, the NLS sequence is residues 264-270, PUFc is residues 273-621, and Clover is residues 1-251.

>mRuby2_NLSPUFa Key: 6×His-mRuby2 NLS PUFa

30 MVRGSHHHHHGMASMTGGQQMGRDLYDDDDKDPMVSKGEELIKENMRMKVVMEGSVNGHQFKC
TGEGEGNPYMGTQTMRIKVIEGGPLPFAFDILATSFMYGSRTFIKYPKGIPDFFKQSFPEGFTW
ERVTRYEDGGVVTVMQDTSLEDGCLVYHVQVRGVNFPSNGPVMQKKTKGWEPNTEMMYPADGGL
RGYTHMALKVDGGGHLSCSFVTTYRSKKTVGNIKMPGIHAVDHRLERLEESDNEMFVVQREHAV
AKFAGLGGGMDELYKGGGGSGPAGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIME

35 FSQDQHGSRFIQLKLERATPAERQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAE
RIRGHVLSLALQMYGSRVIEKALEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQP
QSLQFIIDAFKGQVFALSTHPYGCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGNYVIQ
HVLEHGRPEDKSKIVAEIRGNVLVLSQHKFASNVVEKCVTHASRTERAVLIDEVCTMNDGPHSA
LYTMMKDQYANYVVQKMIDVAEPGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDL

In the above sequence, the NLS sequence is residues 284-290, PUFa is residues 293-641, and 6×His-mRuby2 is residues 1-271, including the 6×His tag at residues 6-11.

>NLSPUFa_p65HSF1 Key: PUFa NLS p65HSF1

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MGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATPAE
RQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIEKA
LEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPY
GCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGNYVIQHVLEHGRPEDKSKIVAEIRGNV
LVLSQHKFASNVVEKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAE
PGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLGGPAGGGGSGGGGSGGGSGPK
KKRKVAAAGSPSGQISNQALALAPSSAPVLAQTMVPSSAMVPLAQPPAPAPVLTPGPPQSLSAP
VPKSTQAGEGTLSEALLHLQFDADEDLGALLGNSTDPGVFTDLASVDNSEFQQLLNQGVSMSHS
TAEPMLMEYPEAITRLVTGSQRPPDPAPTPLGTSGLPNGLSGDEDFSSIADMDFSALLSQISSS
GQGGGGSGFSVDTSALLDLFSPSVTVPDMSLPDLDSSLASIQELLSPQEPPRPPEAENSSPDSG
KQLVHYTAQPLFLLDPGSVDTGSNDLPVLFELGEGSYFSEGDGFAEDPTISLLTGSEPPKAKDP
TVSID

In the above sequence, the NLS sequence is residues 6-12, PUFa is residues 15-363, p65 is residues 427-575, and HSF1 is residues 584-707.

>NLSKRAB_NLSPUFa Key: NLSKRAB PUFa

MGSPKKRKVEASMDAKSLTAWSRTLVTFKDVFVDFTREEWKLLDTAQQIVYRNVMLENYKNLV SLGYQLTKPDVILRLEKGEEPWLVSRGSIVGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLRE IAGHIMEFSQDQHGSRFIQLKLERATPAERQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLE QKLALAERIRGHVLSLALQMYGSRVIEKALEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQK CIECVQPQSLQFIIDAFKGQVFALSTHPYGCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQ YGNYVIQHVLEHGRPEDKSKIVAEIRGNVLVLSQHKFASNVVEKCVTHASRTERAVLIDEVCTM NDGPHSALYTMMKDQYANYVVQKMIDVAEPGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYY MKNGVDLG

In the above sequence, the two NLS sequences are residues 4-10 and residues 99-105, PUFa is residues 108-456, and KRAB is residues 11-92.

>BFPKRAB_NLSPUFa Key: HA-2×NLS-BFPKRAB NLS PUFa MAYPYDVPDYASLGSGSPKKKRKVEDPKKKRKVDGIGSGSNGSSGSSELIKENMHMKLYMEGTV

DNHHFKCTSEGEGKPYEGTQTMRIKVVEGGPLPFAFDILATSFLYGSKTFINHTQGIPDFFKQS
FPEGFTWERVTTYEDGGVLTATQDTSLQDGCLIYNVKIRGVNFTSNGPVMQKKTLGWEAFTETL
YPADGGLEGRNDMALKLVGGSHLIANIKTTYRSKKPAKNLKMPGVYYVDYRLERIKEANNETYV
EQHEVAVARYCDLPSKLGHKLNGGGGGMDAKSLTAWSRTLVTFKDVFVDFTREEWKLLDTAQQI
VYRNVMLENYKNLVSLGYQLTKPDVILRLEKGEEPGGSGGSGPAGILPPKKKRKVSRGRSRLL
EDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATPAERQLVFNEILQAAYQLMVDVF
GNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIEKALEFIPSDQQNEMVRELDGHV
LKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPYGCRVIQRILEHCLPDQTLPI
LEELHQHTEQLVQDQYGNYVIQHVLEHGRPEDKSKIVAEIRGNVLVLSQHKFASNVVEKCVTHA
SRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAEPGQRKIVMHKIRPHIATLRK
YTYGKHILAKLEKYYMKNGVDLG

In the above sequence, the NLS sequence is residues 370-376, PUFa is residues 379-727, and HA-2×NLS-BFPKRAB is residues 1-355, including the HA tag at residues 3-11.

>dCas9Master_mCherry HATag NLS dCas9 mCherry

MIDGGGGSGGGGGGSMYPYDVPDYASPKKKRKVEASDKKYSIGLAIGTNSVGWAVITDEYK VPSKKFKVLGNTDRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEM

AKVDDSFFHRLEESFLVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLI YLALAHMIKFRGHFLIEGDLNPDNSDVDKLFIQLVQTYNQLFEENPINASGVDAKAILSARLSK SRRLENLIAQLPGEKKNGLFGNLIALSLGLTPNFKSNFDLAEDAKLQLSKDTYDDDLDNLLAQI GDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEHHQDLTLLKALVRQQLPEK 5 YKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGTEELLVKLNREDLLRKQRTFDNGSI PHOIHLGELHAILRROEDFYPFLKDNREKIEKILTFRIPYYVGPLARGNSRFAWMTRKSEETIT PWNFEEVVDKGASAOSFIERMTNFDKNLPNEKVLPKHSLLYEYFTVYNELTKVKYVTEGMRKPA FLSGEQKKAIVDLLFKTNRKVTVKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIK DKDFLDNEENEDILEDIVLTLTLFEDREMIEERLKTYAHLFDDKVMKQLKRRRYTGWGRLSRKL 10 INGIRDKQSGKTILDFLKSDGFANRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGS PAIKKGILQTVKVVDELVKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEGIKELGSQ ILKEHPVENTQLQNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDAIVPQSFLKDDSIDNKVLT RSDKNRGKSDNVPSEEVVKKMKNYWROLLNAKLITORKFDNLTKAERGGLSELDKAGFIKROLV ETROITKHVAOILDSRMNTKYDENDKLIREVKVITLKSKLVSDFRKDFOFYKVREINNYHHAHD 15 AYLNAVVGTALIKKYPKLESEFVYGDYKVYDVRKMIAKSEOEIGKATAKYFFYSNIMNFFKTEI TLANGEIRKRPLIETNGETGEIVWDKGRDFATVRKVLSMPQVNIVKKTEVQTGGFSKESILPKR NSDKLIARKKDWDPKKYGGFDSPTVAYSVLVVAKVEKGKSKKLKSVKELLGITIMERSSFEKNP IDFLEAKGYKEVKKDLIIKLPKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHY EKLKGSPEDNEQKQLFVEQHKHYLDEIIEQISEFSKRVILADANLDKVLSAYNKHRDKPIREQA 20 ENIIHLFTLTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQSITGLYETRIDLSQLGGDSP KKKRKVEASGGGGGGGGGGGGGGAMVSKGEEDNMAIIKEFMRFKVHMEGSVNGHEFEIEGE GEGRP YEGTOTAKLKVTKGGPLPFAWDILSPOFMYGSKAYVKHPADIPDYLKLSFPEGFKWERV MNFEDGGVVTVTQDSSLQDGEFIYKVKLRGTNFPSDGPVMQKKTMGWEASSERMYPEDGALKGE IKQRLKLKDGGHYDAEVKTTYKAKKPVQLPGAYNVNIKLDITSHNEDYTIVEQYERAEGRHSTG 25 **GMDELYKID**

In the above sequence, the two NLS sequences are residues 30-36 and 1408-1414, dCas9 is residues 40-1406, mCherry is residues 1436-1671, and the HA tag is at residues 20-28.

>CBPHAT_4×NLS_PUFa_2×NLS Key: CBPHAT NLS PUFa

30

35

40

45

MIFKPEELRQALMPTLEALYRQDPESLPFRQPVDPQLLGIPDYFDIVKNPMDLSTIKRKLDTGQ YQEPWQYVDDVWLMFNNAWLYNRKTSRVYKFCSKLAEVFEQEIDPVMQSLGYCCGRKYEFSPQT LCCYGKQLCTIPRDAAYYSYQNRYHFCEKCFTEIQGENVTLGDDPSQPQTTISKDQFEKKKNDT LDPEPFVDCKECGRKMHQICVLHYDIIWPSGFVCDNCLKKTGRPRKENKFSAKRLQTTRLGNHL EDRVNKFLRRQNHPEAGEVFVRVVASSDKTVEVKPGMKSRFVDSGEMSESFPYRTKALFAFEEI DGVDVCFFGMHVQEYGSDCPPPNTRRVYISYLDSIHFFRPRCLRTAVYHEILIGYLEYVKKLGY VTGHIWACPPSEGDDYIFHCHPPDQKIPKPKRLQEWYKKMLDKAFAERIINDYKDIFKQANEDR LTSAKELPYFEGDFWPNVLEESIKELEQEEEERKKEESTAASETPEGSQGDSKNAKKKNNKKTN KNKSSISRANKKKPSMPNVSNDLSOKLYATMEKHKEVFFVIHLHAGPVISTOPPIVDPDPLLSC DLMDGRDAFLTLARDKHWEFSSLRRSKWSTLCMLVELHTQGQDRFVYTCNECKHHVETRWHCTV CEDYDLCINCYNTKSHTHKMVKWGLGLDDEGSSQGEPQSKSPQESRRLSIQRCIQSLVHACQCR NANCSLPSCQKMKRVVQHTKGCKRKTNGGCPVCKQLIALCCYHAKHCQENKCPVPFCLNINDGG GGSDPKKKRKVDPKKKRKVDPKKKRKVGSTGSRNDGGGGSGGGGGGGGGGGGGAGILPPKKKRKV SRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATPAERQLVFNEILQAA YQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIEKALEFIPSDQQNEM VRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPYGCRVIQRILEHC LPDOTLPILEELHOHTEOLVODOYGNYVIOHVLEHGRPEDKSKIVAEIRGNVLVLSOHKFASNV VEKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAEPGQRKIVMHKIR PHIATLRKYTYGKHILAKLEKYYMKNGVDLGDPKKKRKVDPKKKRKVGGRGGGGGGGGGGGGG SGPA

In the above sequence, the six 7-residue NLS sequences begin at residues 773, 781, 789, 826, 1185, and 1193, PUFa is residues 835-1183, and CBPHAT is residues 2-764.

> 4×NLS_PUFa_2×NLS_CBPHAT Key: NLS PUFa CBPHAT

25

MIDGGGGSDPKKKRKVDPKKKRKVDPKKKRKVGSTGSRNDGGGGSGGGGGGGGGGGAGILPPK 5 KKRKVSRGRSRLLEDFRNNRYPNLOLREIAGHIMEFSODOHGSRFIOLKLERATPAEROLVFNE ILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIEKALEFIPSD QQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPYGCRVIQR ILEHCLPDQTLPILEELHQHTEQLVQDQYGNYVIQHVLEHGRPEDKSKIVAEIRGNVLVLSQHK FASNVVEKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAEPGQRKIV 10 MHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLGDPKKKRKVDPKKKRKVGGRGGGGGGG SGGGGSGPAIFKPEELRQALMPTLEALYRQDPESLPFRQPVDPQLLGIPDYFDIVKNPMDLSTI KRKLDTGQYQEPWQYVDDVWLMFNNAWLYNRKTSRVYKFCSKLAEVFEQEIDPVMQSLGYCCGR KYEFSPQTLCCYGKQLCTIPRDAAYYSYQNRYHFCEKCFTEIQGENVTLGDDPSQPQTTISKDQ FEKKKNDTLDPEPFVDCKECGRKMHQICVLHYDIIWPSGFVCDNCLKKTGRPRKENKFSAKRLQ 15 TTRLGNHLEDRVNKFLRRQNHPEAGEVFVRVVASSDKTVEVKPGMKSRFVDSGEMSESFPYRTK ALFAFEEIDGVDVCFFGMHVOEYGSDCPPPNTRRVYISYLDSIHFFRPRCLRTAVYHEILIGYL EYVKKLGYVTGHIWACPPSEGDDYIFHCHPPDQKIPKPKRLQEWYKKMLDKAFAERIINDYKDI FKQANEDRLTSAKELPYFEGDFWPNVLEESIKELEQEEEERKKEESTAASETPEGSQGDSKNAK KKNNKKTNKNKSSISRANKKKPSMPNVSNDLSQKLYATMEKHKEVFFVIHLHAGPVISTQPPIV 20 DPDPLLSCDLMDGRDAFLTLARDKHWEFSSLRRSKWSTLCMLVELHTQGQDRFVYTCNECKHHV ETRWHCTVCEDYDLCINCYNTKSHTHKMVKWGLGLDDEGSSQGEPQSKSPQESRRLSIQRCIQS LVHACQCRNANCSLPSCQKMKRVVQHTKGCKRKTNGGCPVCKQLIALCCYHAKHCQENKCPVPF CLNI

In the above sequence, the six 7-residue NLS sequences begin at residues 10, 18, 26, 63, 422, and 430, PUFa is residues 72-420, and CBPHAT is residues 458-1220.

Name and Description	DNA sequence		
sgRNA-PBS expression casses	sgRNA-PBS expression cassettes:		
U6::sgRNA-0×PBS expression cassette containing the target sequences as Ns without PBS sequences	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtattcg atttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNNNNNNNNNNNNN		
U6::sgRNA-1×PBS32 expression cassette containing the target sequences as Ns and 1 copy of PBS32 (UGUAUGUA)	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtattcg atttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNNNNNNNNNNNNN		

Name and Description	DNA sequence
U6::sgRNA-5×PBS32	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
expression cassette	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca
containing the target	aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa
sequences as Ns and 5	ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcgatttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNN
copies of PBS32	NNNNNNNgtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag
(UGUAUGUA) separated	gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt
by GCC spacer sequence	ctccagatTGTATGTAGCCTGTATGTAGCCTGTATGTAGCC
attached at 3' region of the	TGTATGTAagatCTTTTTTTgttttagagctagaaatagcaagttaaaataa qqctaqtccqtaqcqcqtqcqccaattctqcaqacaaatqqc
sgRNA	gyctayteegtayegegegecaatteegeagacaaatgge
U6::sgRNA-15×PBS32	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
expression and cloning	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca
cassette containing the	aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa
target sequences as Ns and	ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcgatttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNN
15 copies of PBS32	NNNNNNNngtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag
(UGUAUGUA) separated	gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt
by GCC spacer sequence	ctccagatTGTATGTAGCCTGTATGTAGCCTGTATGTAGCC
attached at 3' region of the	TGTATGTAAGATTGTATGTAGCTGTATGTAGCCTGTATGTA
sgRNA	TGTAGCCTGTATGTAGACTTTTTTTTTttttagagctagaaatagcaagtt
	aaaataaggctagtccgtagcgcgtgcgccaattctgcagacaaatggc
U6::sgRNA-25×PBS32	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
expression and cloning	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca
cassette containing the	aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa
target sequences as Ns and	ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcgatttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNN
25 copies of PBS32	NNNNNNNngtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag
(UGUAUGUA) separated	gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt
by GCC spacer sequence	ctccagatTGTATGTAGCCTGTATGTAGCCTGTATGTAGCC
attached at 3' region of the	TGTATGTAAGATTGTATGTAGCCTGTATGTAGCCTGTATGTA
sgRNA	ATGTAGCCTGTATGTAAGATTGTATGTAGCCTGTATGTAGCCTGTATGTA
	CTGTATGTAGCCTGTATGTAAGATTGTATGTAGCCTGTATG
	TAGCCTGTATGTAGCCTGTATGTAagatCTTTTTTTgttttagagctagaaa
	tagcaagttaaaataaggctagtccgtagcgcgtgcgccaattctgcagaca
LIGUADNIA 15 DDCC070	aatggc gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
U6::sgRNA-1×PBS6272	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca
expression cassette	aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa
containing the target	ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcg
sequences as Ns and 1 copy of PBS6272	atttettggetttatatatettGTGGAAAGGACGAAACACCNNNNNNNNNN
	NNNNNNNNgtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt
(UUGAUAUA)	ctccagatgccTtgATATAgccagatCTTTTTTTgttttagagctagaaata
	gcaagttaaaataaggctagtccgtagcgcgtgcgccaattctgcagacaaa
	tggc
U6::sgRNA-2×PBS6272	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
expression cassette	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa
containing the target	ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcg
sequences as Ns and 2	atttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNN
copies of PBS6272	NNNNNNN gtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag
(UUGAUAUA) separated	gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt
by GCC spacer sequence	ctccagatTTGATATAGCCTTGATATAagatCTTTTTTTgttttagagctag aaatagcaagttaaaataaggctagtccgtagcgcgtgcgccaattctgcag
attached at 3' region of the	acaaatggc
sgRNA	

Name and Description	DNA sequence
U6::sgRNA-5×PBS6272	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
expression cassette	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca
containing the target	aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa
sequences as Ns and 5	ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcg
1	atttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNN
copies of PBS6272	NNNNNNNNgtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag
(UUGAUAUA) separated	gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt ctccagatTTGATATAGCCTTGATATAGCCTTGATATAGCC
by GCC spacer sequence	TTGATATAagatCTTTTTTTgttttagagctagaaatagcaagttaaaataa
attached at 3' region of the	ggctagtccgtagcgcgtgcgccaattctgcagacaaatggc
sgRNA	
U6::sgRNA-10×PBS6272	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
expression cassette	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca
containing the target	aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa
sequences as Ns and 10	ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcgatttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNN
copies of PBS6272	NNNNNNNngtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag
(UUGAUAUA) separated	gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt
by GCC spacer sequence	ctccagatTTGATATAGCCTTGATATAGCCTTGATATAGCC
1	TTGATATAAGATTTGATATAGCCTTGATATAGCCTTGATAT
attached at 3' region of the	AGCCTTGATATAagatCTTTTTTTgttttagagctagaaatagcaagttaaa
sgRNA	ataaggctagtccgtagcgcgtgcgccaattctgcagacaaatggc
U6::sgRNA-15×PBS6272	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
expression cassette	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa
containing the target	ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcg
sequences as Ns and 15	atttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNN
copies of PBS6272	NNNNNNN gtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag
(UUGAUAUA) separated	gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt
by GCC spacer sequence	ctccagatTTGATATAGCCTTGATATAGCCTTGATATAGCC
attached at 3' region of the	TTGATATAAGATTTGATATAGCCTTGATATAGCCTTGATAT
_	AGCCTTGATATAAGATTTGATATAGCCTTGATATAGCCTTG
sgRNA	ATATAGCCTTGATATAagatCTTTTTTTTttttagagctagaaatagcaagt
LIGUARDNA 20VDDS6272	taaaataaggctagtccgtagcgcgtgcgccaattctgcagacaaatggc gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
U6::sgRNA-20×PBS6272	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca
expression cassette	aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa
containing the target	ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcg
sequences as Ns and 20	atttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNNNNN
copies of PBS6272	NNNNNNNN gtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag
(UUGAUAUA) separated	gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt
by GCC spacer sequence	ctccagatTTGATATAGCCTTGATATAGCCTTGATATAGCCTTGATATAGCC
attached at 3' region of the	TTGATATAAGATTTGATATAGCCTTGATATAGCCTTGATATAGCCTTGATAT AGCCTTGATATAAGATTTGATATAGCCTTGATATAGCCTTGATATAGCCTTG
sgRNA	ATATAGCCTTGATATAAGATTTGATATAGCCTTGATATAGCCTTGATATAGC
-6	CTTGATATAGCCTTGATATAagatCTTTTTTTgttttagagctagaaatagc
	aagttaaaataaggctagtccgtagcgcgtgcgccaattctgcagacaaatg
	gc

Name and Description U6::sgRNA-25×PBS6272

U6::sgRNA-25×PBS6272 expression cassette containing the target sequences as Ns and 25 copies of PBS6272 (UUGAUAUA) separated by GCC spacer sequence attached at 3' region of the sgRNA

DNA sequence

gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcg atttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNN NNNNNNNNNNNTSTTAAGGCTTGGAAACAGCATAGCATTGGGT ctccagatTTGATATAGCCTTGATAGCCTTGATATAGCCTTGATATAGCCTTGAT

U6::sgRNA-47×PBS6272 expression cassette containing the target sequences as Ns and 47 copies of PBS6272 (UUGAUAUA) separated by GCC spacer sequence attached at 3' region of the sgRNA

gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcg **NNNNNNN**gtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt ctccagatTTGATATAGCCTTGATATAGCCTTGATATAGCC TTGATATAAGATTTGATATAGCCTTGATATAGCCTTGATAT AGCCTTGATATAAGATTTGATATACCTTGATATAGCCTTGATATAGCCTTGA TATAGCCTTGATATAAGATTTGATATAGCCTTGATATAGCCTTGATATAGCC TTGATATAGCCTTGATATAGCCTTGATATAGCCTTGATATAAGATTTGATAT AGCCTTGATATAGCCTTGATATAGCCTTGATATAGCCTTGATATAAGATTTG ATATAGCCTTGATATAGCCTTGATATAGCCTTGATATAAGA TTTGATATAGCCTTGATATAGCCTTGATATAGCCTTGATAT AAGATTTGATATAGCCTTGATATAGCCTTGATATAGCCTTG ATATAAGATTTGATATAGCCTTGATATAGCCTTGATATAGC CTTGATATAagatCTTTTTTgttttagagctagaaatagcaagttaaaata aggctagtccgtagcgcgtgcgccaattctgcagacaaatggc

U6::sgRNA-2×PBS6272-10Spacer expression cassette containing the target sequences as Ns and 2 copies of PBS6272 (UUGAUAUA) separated by GCCAGATGCC spacer sequence attached at 3' region of the sgRNA

U6::sgRNA-6×PBS6272-10Spacer expression cassette containing the target sequences as Ns and 6 copies of PBS6272 (UUGAUAUA) separated by GCCAGATGCC spacer sequence attached at 3' region of the sgRNA

Name and Description

U6::sgRNA-15×PBS6272-10Spacer expression cassette containing the target sequences as Ns and 15 copies of PBS6272 (UUGAUAUA) separated by GCCAGATGCC spacer sequence attached at 3' region of the sgRNA

DNA sequence

U6::sgRNA-20×PBS6272-10Spacer expression cassette containing the target sequences as Ns and 20 copies of PBS6272 (UUGAUAUA) separated by GCCAGATGCC spacer sequence attached at 3' region of the sgRNA

gagggeetattteeeatgatteetteatatttgeatataegataeaaggetg ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcg ${\bf NNNNNNN} \\ {\bf gtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag}$ gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt ctccagatGCCTTGATATAGCCAGATGCCTTGATATAGCCAGATGCCTTGAT ATAGCCAGATGCCTTGATATAGCCAGATGCCTTGATATAGCCAGATGCCTTG ATATAGCCAGATGCCTTGATATAGCCAGATGCCT TGATATAGCCAGATGCCTTGATATAGCCAGATGCCTTGATATAGCCAGATGC CTTGATATAGCCAGATGCCTTGATATAGCCAGATGCCTTGATATAGCCAGAT GCCTTGATATAGCCAGATGCCTTGATATAGCCAGATGCCTTGATATAGCCAG ATGCCTTGATATAGCCAGATCCTTGATATAGCCAGATGCCTTGATATAGCCa gatCTTTTTTqttttaqaqctaqaaataqcaaqttaaaataaqqctaqtcc gtagegegtgegeeaattetgeagaeaaatgge

U6::5×PBS32-sgRNA
expression cassette
containing the target
sequences as Ns and 5
copies of PBS32
(UGUAUGUA) separated
by GCCAGATGCC spacer
sequence attached at 5'
region of the sgRNA

U6::sgRNA-2×[PBS32-PBS6272] expression cassette containing the target sequences as Ns and 2 copies of PBS32(UGUAUGUA)-PBS6272 (UUGAUAUA) clusters attached at 3' region of the sgRNA

Name and Description	DNA sequence
U6::sgRNA-8×[PBS32-	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
PBS6272] expression	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca
cassette containing the	aaatacgtgacgtagaaagtaataatttcttgggtagttttgcagttttaaaa ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcg
target sequences as Ns and 8	atttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNN
copies of	NNNNNNNNgtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag
PBS32(UGUAUGUA)-	gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt
PBS6272 (UUGAUAUA)	ctccagatTGTATGTAGTCTATTGATATAGTCTTGTCTATGTATGTAGTCTA TTGATATAAGATTGTATGTAGTCTATTGATATAGTCTTGTCTATGTATG
clusters attached at 3' region	TCTATTGATATAAGATTGTATGTAGTCTATTGATATAGTCTTGTCTATGTAT
of the sgRNA	GTAGTCTATTGATATAAGATTGTATGTAGTCTATTGATATAGTCTTGTCTAT
	GTATGTAGTCTATTGATATAagatCTTTTTTTgttttagagctagaaatagc
	aagttaaaataaggctagtccgtagcgcgtgcgccaattctgcagacaaatg gc
U6::sgRNA-4×[PBS32-	gagggcctatttcccatgattccttcatatttgcatatacgatacaaggctg
PBS6272] expression	ttagagagataattggaattaatttgactgtaaacacaaagatattagtaca
cassette containing the	aaatacgtgacgtagaaagtaataatttcttgggtagtttgcagttttaaaa
target sequences as Ns and 4	ttatgttttaaaatggactatcatatgcttaccgtaacttgaaagtatttcgatttcttggctttatatatcttGTGGAAAGGACGAAACACCNNNNNNNNNNN
copies of	NNNNNNNNgtttAagagctaTGCTGGAAACAGCAtagcaagttTaaataag
PBS32(UGUAUGUA)-	gctagtccgttatcaacttgaaaaagtggcaccgagtcggtgcCAATTGggt
PBS6272 (UUGAUAUA)	ctccagatTGTATGTAGTCTATTGATATAGTCTTGTCTATGTATGTAGTCTA
clusters attached at 3' region	TTGATATAAGATTGTATGTAGTCTATTGATATAGTCTTGTCTATGTATG
of the sgRNA	TCTATTGATATAagatCTTTTTTTgttttagagctagaaatagcaagttaaa ataaggctagtccgtagcgcgtgcgccaattctgcagacaaatggc
	etimes an additional G is prepended to increase U6
transcriptional efficiency):	1 1
Control Sequence	GTTCTCTTGCTGAAAGCTCGA
TetO promoter	GCTTTTCTCTATCACTGATA
SV40P1	GCATACTTCTGCCTGCGGGAGCCTG
SV40P2	GAAAGTCCCCAGGCTCCCCAGC
SV40P3	GCATCTCAATTAGTCAGCAACC
Telomere	GTTAGGGTTAGGGTTA
Centromere	GTTGAGGCCTTCGTTGGAAAC
MUC4-Nonrepeat-1	GAAGAGTGGAGGCCGTGCGCGG
MUC4-Nonrepeat-2	GCAAGCAAGGGACAAGG
MUC4-Nonrepeat-3	GATGTTTCAGGACTAGGCTGA
MUC4-Nonrepeat-4	GAGCTGGGCCAGGAGAGA
MUC4-Nonrepeat-5	GAGGGGTCTGTGGAGAGTTT
MUC4-Nonrepeat-6	GGCTTGGTGTATTCAGAATG
MUC4-Nonrepeat-7	GTAGAGATGCCGCCCCCCCC
OCT4-PP-1	GGCCCGCCCCTGGATGGG
OCT4-PP-2	GGGGGAGAAACTGAGGCGA
OCT4-PP-3	GGTGGTGGCAATGGTGTCTG
OCT4-PP-4	GACACAACTGGCGCCCCTCC
OCT4-PE-1	GGCCCCTACTTCCCCTTCAA
OCT4-PE-2	GAGTGATAAGACACCCGCTT
OCT4-PE-3	GCCTGGGAGGACTGGGGGA
OCT4-PE-4	GGACAATCCCGGTCCCCAGA
OCT4-DE-1	GGTCTGCCGGAAGGTCTACA
OCT4-DE-2	GGCAGGTAGATTATGGGGCC
OCT4-DE-3	GAAGACGGCCTCTCAGAGGA

Name and Description	DNA sequence
OCT4-DE-4	GTATTTCTGGCCTGGGCAAG
SOX2-PP-1	GCATGTGACGGGGGCTGTCA
SOX2-PP-2	GCTGCCGGGTTTTGCATGAA
SOX2-PP-3	GCCGGCCGCGGGGAGGC
SOX2-PP-4	GGCAGGCGAGGAGG
SV40-P1	GCATACTTCTGCCTGCTGGGGAGCCTG

Name	Peptide sequence
S. pyrogene NLS-	MYPYDVPDYASPKKKRKVEASDKKYSIGLAIGTNSVGWAVITDEYKVPSKKFKVLGNT
dCas9-NLS	DRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDS
deasy-NES	FFHRLEESFLVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLI
	YLALAHMIKFRGHFLIEGDLNPDNSDVDKLFIQLVQTYNQLFEENPINASGVDAKAIL
	SARLSKSRRLENLIAQLPGEKKNGLFGNLIALSLGLTPNFKSNFDLAEDAKLQLSKDT
	YDDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEH
	HQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGT
	EELLVKLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYPFLKDNREKIEKIL
	TFRIPYYVGPLARGNSRFAWMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNL
	PNEKVLPKHSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVT
	VKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILED
	IVLTLTLFEDREMIEERLKTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSG
	KTILDFLKSDGFANRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIK
	KGILQTVKVVDELVKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEGIKELG
	SQILKEHPVENTQLQNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDAIVPQSFLKDD
	SIDNKVLTRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGL
	SELDKAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSKLVSDF
	RKDFQFYKVREINNYHHAHDAYLNAVVGTALIKKYPKLESEFVYGDYKVYDVRKMIAK
	SEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFAT
	VRKVLSMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVA
	YSVLVVAKVEKGKSKKLKSVKELLGITIMERSSFEKNPIDFLEAKGYKEVKKDLIIKL
	PKYSLFELENGRKRMLASAGELOKGNELALPSKYVNFLYLASHYEKLKGSPEDNEOKO
	LFVEQHKHYLDEIIEQISEFSKRVILADANLDKVLSAYNKHRDKPIREQAENIIHLFT
	LTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHOSITGLYETRIDLSOLGGDSPKK
	KRKVEAS
S. pyogenes NLS-	MYPYDVPDYASPKKKRKVEASDKKYSIGLDIGTNSVGWAVITDEYKVPSKKFKVLGNT
Cas9WT-NLS	DRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDS
Casy W 1-IVES	FFHRLEESFLVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLI
	YLALAHMIKFRGHFLIEGDLNPDNSDVDKLFIQLVQTYNQLFEENPINASGVDAKAIL
	SARLSKSRRLENLIAQLPGEKKNGLFGNLIALSLGLTPNFKSNFDLAEDAKLQLSKDT
	YDDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEH
	HQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGT
	EELLVKLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYPFLKDNREKIEKIL
	TFRIPYYVGPLARGNSRFAWMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNL
	PNEKVLPKHSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVT
	VKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILED
	IVLTLTLFEDREMIEERLKTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSG
	KTILDFLKSDGFANRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIK
	KGILQTVKVVDELVKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEGIKELG
	SQILKEHPVENTQLQNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDD
	SIDNKVLTRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGL
	SELDKAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSKLVSDF
	RKDFQFYKVREINNYHHAHDAYLNAVVGTALIKKYPKLESEFVYGDYKVYDVRKMIAK
	SEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFAT
	VRKVLSMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVA
	YSVLVVAKVEKGKSKKLKSVKELLGITIMERSSFEKNPIDFLEAKGYKEVKKDLIIKL
	PKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLKGSPEDNEQKQ
	LFVEQHKHYLDEIIEQISEFSKRVILADANLDKVLSAYNKHRDKPIREQAENIIHLFT
	LTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQSITGLYETRIDLSQLGGDSPKK
	KRKVEAS

Name	Peptide sequence
S. pyogenes NLS-	MYPYDVPDYASPKKKRKVEASDKKYSIGLAIGTNSVGWAVITDEYKVPSKKFKVLGNT
Cas9Nickase(D10A)-	DRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDS
` '	FFHRLEESFLVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLI
NLS	YLALAHMIKFRGHFLIEGDLNPDNSDVDKLFIQLVQTYNQLFEENPINASGVDAKAIL
	SARLSKSRRLENLIAQLPGEKKNGLFGNLIALSLGLTPNFKSNFDLAEDAKLQLSKDT
	YDDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEH
	HQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGT
	EELLVKLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYPFLKDNREKIEKIL
	TFRIPYYVGPLARGNSRFAWMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNL
	PNEKVLPKHSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVT
	VKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILED
	IVLTLTLFEDREMIEERLKTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSG
	KTILDFLKSDGFANRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIK
	KGILQTVKVVDELVKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEGIKELG
	SQILKEHPVENTQLQNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDHIVPQSFLKDD
	SIDNKVLTRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGL
	SELDKAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSKLVSDF
	RKDFQFYKVREINNYHHAHDAYLNAVVGTALIKKYPKLESEFVYGDYKVYDVRKMIAK
	SEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFAT
	VRKVLSMPQVNIVKKTEVQTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVA
	YSVLVVAKVEKGKSKKLKSVKELLGITIMERSSFEKNPIDFLEAKGYKEVKKDLIIKL
	PKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLKGSPEDNEQKQ
	LFVEQHKHYLDEIIEQISEFSKRVILADANLDKVLSAYNKHRDKPIREQAENIIHLFT
	LTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQSITGLYETRIDLSQLGGDSPKK
	KRKVEAS
S. pyogenes NLS-	MYPYDVPDYASPKKKRKVEASDKKYSIGLDIGTNSVGWAVITDEYKVPSKKFKVLGNT
Cas9Nickase(H840A)	DRHSIKKNLIGALLFDSGETAEATRLKRTARRRYTRRKNRICYLQEIFSNEMAKVDDS
-NLS	FFHRLEESFLVEEDKKHERHPIFGNIVDEVAYHEKYPTIYHLRKKLVDSTDKADLRLI
-INL/S	YLALAHMIKFRGHFLIEGDLNPDNSDVDKLFIQLVQTYNQLFEENPINASGVDAKAIL
	SARLSKSRRLENLIAQLPGEKKNGLFGNLIALSLGLTPNFKSNFDLAEDAKLQLSKDT
	YDDDLDNLLAQIGDQYADLFLAAKNLSDAILLSDILRVNTEITKAPLSASMIKRYDEH
	HQDLTLLKALVRQQLPEKYKEIFFDQSKNGYAGYIDGGASQEEFYKFIKPILEKMDGT
	EELLVKLNREDLLRKQRTFDNGSIPHQIHLGELHAILRRQEDFYPFLKDNREKIEKIL
	TFRIPYYVGPLARGNSRFAWMTRKSEETITPWNFEEVVDKGASAQSFIERMTNFDKNL
	PNEKVLPKHSLLYEYFTVYNELTKVKYVTEGMRKPAFLSGEQKKAIVDLLFKTNRKVT
	VKQLKEDYFKKIECFDSVEISGVEDRFNASLGTYHDLLKIIKDKDFLDNEENEDILED
	IVLTLTLFEDREMIEERLKTYAHLFDDKVMKQLKRRRYTGWGRLSRKLINGIRDKQSG
	KTILDFLKSDGFANRNFMQLIHDDSLTFKEDIQKAQVSGQGDSLHEHIANLAGSPAIK
	KGILQTVKVVDELVKVMGRHKPENIVIEMARENQTTQKGQKNSRERMKRIEEGIKELG
	SQILKEHPVENTQLQNEKLYLYYLQNGRDMYVDQELDINRLSDYDVDAIVPQSFLKDD
	SIDNKVLTRSDKNRGKSDNVPSEEVVKKMKNYWRQLLNAKLITQRKFDNLTKAERGGL
	SELDKAGFIKRQLVETRQITKHVAQILDSRMNTKYDENDKLIREVKVITLKSKLVSDF
	RKDFQFYKVREINNYHHAHDAYLNAVVGTALIKKYPKLESEFVYGDYKVYDVRKMIAK
	SEQEIGKATAKYFFYSNIMNFFKTEITLANGEIRKRPLIETNGETGEIVWDKGRDFAT
	VRKVLSMPOVNIVKKTEVOTGGFSKESILPKRNSDKLIARKKDWDPKKYGGFDSPTVA
	YSVLVVAKVEKGKSKKLKSVKELLGITIMERSSFEKNPIDFLEAKGYKEVKKDLIIKL
	PKYSLFELENGRKRMLASAGELQKGNELALPSKYVNFLYLASHYEKLKGSPEDNEQKQ
	LFVEQHKHYLDEIIEQISEFSKRVILADANLDKVLSAYNKHRDKPIREQAENIIHLFT
	LTNLGAPAAFKYFDTTIDRKRYTSTKEVLDATLIHQSITGLYETRIDLSQLGGDSPKK
	KRKVEAS
	TOTAL VILLO

Name	Peptide sequence
Ruby::PUF(3-2)	MVRGSHHHHHHGMASMTGGQQMGRDLYDDDDKDPMVSKGEELIKENMRMKVVMEGSVN
	GHQFKCTGEGEGNPYMGTQTMRIKVIEGGPLPFAFDILATSFMYGSRTFIKYPKGIPD
	FFKQSFPEGFTWERVTRYEDGGVVTVMQDTSLEDGCLVYHVQVRGVNFPSNGPVMQKK
	TKGWEPNTEMMYPADGGLRGYTHMALKVDGGGHLSCSFVTTYRSKKTVGNIKMPGIHA
	VDHRLERLEESDNEMFVVQREHAVAKFAGLGGGMDELYKGGGGSGPAGILPPKKKRKV
	SRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATPAERQLVFN
	EILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIEKA
	LEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFA
	LSTHPYGCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGNYVIQHVLEHGRPED
	KSKIVAEIRGNVLVLSQHKFASNVVEKCVTHASRTERAVLIDEVCTMNDGPHSALYTM
	MKDQYANYVVQKMIDVAEPGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGV
	DLG
Ruby::PUF(6-2/7-2)	MVRGSHHHHHHGMASMTGGQQMGRDLYDDDDKDPMVSKGEELIKENMRMKVVMEGSVN
, , , , ,	GHQFKCTGEGEGNPYMGTQTMRIKVIEGGPLPFAFDILATSFMYGSRTFIKYPKGIPD
	FFKQSFPEGFTWERVTRYEDGGVVTVMQDTSLEDGCLVYHVQVRGVNFPSNGPVMQKK
	TKGWEPNTEMMYPADGGLRGYTHMALKVDGGGHLSCSFVTTYRSKKTVGNIKMPGIHA
	VDHRLERLEESDNEMFVVQREHAVAKFAGLGGGMDELYKGGGGSGPAGILPPKKKRKV
	SRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATPAERQLVFN
	EILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGCRVIQKA
	LEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFA
	LSTHPYGCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGSYVIEHVLEHGRPED
	KSKIVAEIRGNVLVLSQHKFANNVVQKCVTHASRTERAVLIDEVCTMNDGPHSALYTM
	MKDQYANYVVQKMIDVAEPGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGV
	DLG
Clover::PUF(3-2)	MVSKGEELFTGVVPILVELDGDVNGHKFSVRGEGEGDATNGKLTLKFICTTGKLPVPW
	PTLVTTFGYGVACFSRYPDHMKQHDFFKSAMPEGYVQERTISFKDDGTYKTRAEVKFE
	GDTLVNRIELKGIDFKEDGNILGHKLEYNFNSHNVYITADKQKNGIKANFKIRHNVED
	GSVQLADHYQQNTPIGDGPVLLPDNHYLSHQSALSKDPNEKRDHMVLLEFVTAAGITH
	GMDELYKSRGPYSIVSPKCGGGGGSGPAGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQ LREIAGHIMEFSQDQHGSRFIQLKLERATPAERQLVFNEILQAAYQLMVDVFGNYVIQ
	KFFEFGSLEQKLALAERIRGHVLSLALQMYGSRVIEKALEFIPSDQQNEMVRELDGHV
	LKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPYGCRVIQRILEHCLP
	DOTLPILEELHOHTEOLVODOYGNYVIOHVLEHGRPEDKSKIVAEIRGNVLVLSOHKF
	ASNVVEKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAEPG
	ORKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLG
Classem DLIE(6, 2/7, 2)	MVSKGEELFTGVVPILVELDGDVNGHKFSVRGEGEGDATNGKLTLKFICTTGKLPVPW
Clover::PUF(6-2/7-2)	PTLVTTFGYGVACFSRYPDHMKQHDFFKSAMPEGYVQERTISFKDDGTYKTRAEVKFE
	GDTLVNRIELKGIDFKEDGNILGHKLEYNFNSHNVYITADKQKNGIKANFKIRHNVED
	GSVQLADHYQQNTPIGDGPVLLPDNHYLSHQSALSKDPNEKRDHMVLLEFVTAAGITH
	GMDELYKSRGPYSIVSPKCGGGGGSGPAGILPPKKKRKVSRGRSRLLEDFRNNRYPNLO
	LREIAGHIMEFSQDQHGSRFIQLKLERATPAERQLVFNEILQAAYQLMVDVFGNYVIQ
	KFFEFGSLEQKLALAERIRGHVLSLALQMYGCRVIQKALEFIPSDQQNEMVRELDGHV
	LKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPYGCRVIQRILEHCLP
	DOTLPILEELHOHTEOLVODOYGSYVIEHVLEHGRPEDKSKIVAEIRGNVLVLSOHKF
	ANNVVQKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQKMIDVAEPG
	QRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLG
PUF(3-2)::VP64	MGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLE
	RATPAERQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLA
	LQMYGSRVIEKALEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQ
	FIIDAFKGQVFALSTHPYGCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGNYV
	IQHVLEHGRPEDKSKIVAEIRGNVLVLSQHKFASNVVEKCVTHASRTERAVLIDEVCT
	MNDGPHSALYTMMKDQYANYVVQKMIDVAEPGQRKIVMHKIRPHIATLRKYTYGKHIL
	AKLEKYYMKNGVDLGGPAGSGRADALDDFDLDMLGSDALDDFDLDMLGSDALDDFDLD
	MLGSDALDDFDLDMLYID

Name	Peptide sequence
PUF(6-2/7-2)::VP64	MGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLE
	RATPAERQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLA
	LQMYGCRVIQKALEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQ
	FIIDAFKGQVFALSTHPYGCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGSYV
	IEHVLEHGRPEDKSKIVAEIRGNVLVLSQHKFANNVVQKCVTHASRTERAVLIDEVCT
	MNDGPHSALYTMMKDQYANYVVQKMIDVAEPGQRKIVMHKIRPHIATLRKYTYGKHIL
	AKLEKYYMKNGVDLGGPAGSGRADALDDFDLDMLGSDALDDFDLDMLGSDALDDFDLD
	MLGSDALDDFDLDMLYID
PUF(6-2/7-	MGILPPKKKRKVSRGRSRLLEDFRNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLE
2)::p65_HSF1	RATPAERQLVFNEILQAAYQLMVDVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLA
2)pes_rist 1	LQMYGCRVIQKALEFIPSDQQNEMVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQ
	FIIDAFKGQVFALSTHPYGCRVIQRILEHCLPDQTLPILEELHQHTEQLVQDQYGSYV
	IEHVLEHGRPEDKSKIVAEIRGNVLVLSQHKFANNVVQKCVTHASRTERAVLIDEVCT
	MNDGPHSALYTMMKDQYANYVVQKMIDVAEPGQRKIVMHKIRPHIATLRKYTYGKHIL
	AKLEKYYMKNGVDLGGPAGGGGSGGGGGGGGGGGFKKKRKVAAAGSPSGQISNQALAL
	APSSAPVLAQTMVPSSAMVPLAQPPAPAPVLTPGPPQSLSAPVPKSTQAGEGTLSEAL
	LHLQFDADEDLGALLGNSTDPGVFTDLASVDNSEFQQLLNQGVSMSHSTAEPMLMEYP
	EAITRLVTGSQRPPDPAPTPLGTSGLPNGLSGDEDFSSIADMDFSALLSQISSSGQGG
	GGSGFSVDTSALLDLFSPSVTVPDMSLPDLDSSLASIQELLSPQEPPRPPEAENSSPD
	SGKQLVHYTAQPLFLLDPGSVDTGSNDLPVLFELGEGSYFSEGDGFAEDPTISLLTGS
	EPPKAKDPTVSID
KRAB::PUF(6-2/7-2)	MGSPKKKRKVEASMDAKSLTAWSRTLVTFKDVFVDFTREEWKLLDTAQQIVYRNVMLE
, , ,	NYKNLVSLGYQLTKPDVILRLEKGEEPWLVSRGSIVGILPPKKKRKVSRGRSRLLEDF
	RNNRYPNLQLREIAGHIMEFSQDQHGSRFIQLKLERATPAERQLVFNEILQAAYQLMV
	DVFGNYVIQKFFEFGSLEQKLALAERIRGHVLSLALQMYGCRVIQKALEFIPSDQQNE
	MVRELDGHVLKCVKDQNGNHVVQKCIECVQPQSLQFIIDAFKGQVFALSTHPYGCRVI
	QRILEHCLPDQTLPILEELHQHTEQLVQDQYGSYVIEHVLEHGRPEDKSKIVAEIRGN
	VLVLSQHKFANNVVQKCVTHASRTERAVLIDEVCTMNDGPHSALYTMMKDQYANYVVQ
	KMIDVAEPGQRKIVMHKIRPHIATLRKYTYGKHILAKLEKYYMKNGVDLG

List of vectors and their Addgene accession numbers

pAC	Descriptive name	Description
number		
pAC164	pmax-dCas9Master_VP64	dCas9-VP64 driven by CAGGS promoter in
		expression vector pmax (Clontech)
pAC1119	PB3-neo(-)-	PB gateway destination vector with neo selectable
	pmaxDEST(+)	marker and pmax cassette (Clonetech)
pAC1355	pmax-NLSPUFa_VP64	NLSPUFa_VP64 in transient expression vector pmax
pAC1356	pmax-NLSPUFb_VP64	NLSPUFb_VP64 in expression vector pmax
pAC1357	pmax-NLSPUFw_VP64	NLSPUFw_VP64 in expression vector pmax
pAC1358	pmax-NLSPUFc_VP64	NLSPUFc_VP64 in expression vector pmax
pAC1360	PB3-neo(-)-pmax-	Clover_NLSPUFa in pAC1119
	Clover_NLSPUFa	
pAC1362	PB3-neo(-)-pmax-	mRuby2_NLSPUFa in pAC1119
	mRuby2_NLSPUFa	
pAC1364	pmax-	dCas9Master_mCBPHAT in pmax expression vector
	dCas9Master_mCBPHAT	
pAC1371	pX-sgRNA-5xPBSa	Cloning vector for expression of sgRNA-5xPBSa
pAC1372	pX-sgRNA-15xPBSa	Cloning vector for expression of sgRNA-15xPBSa
pAC1373	pX-sgRNA-25xPBSa	Cloning vector for expression of sgRNA-25xPBSa
pAC1374	pX-sgRNA-5xPBSb	Cloning vector for transient expression of sgRNA-
		5xPBSb

pAC	Descriptive name	Description
number		
pAC1375	pX-sgRNA-15xPBSb	Cloning vector for expression of sgRNA-15xPBSb
pAC1376	pX-sgRNA-25xPBSb	Cloning vector for expression of sgRNA-25xPBSb
pAC1379	pX-sgRNA-5xPBSw	Cloning vector for expression of sgRNA-5xPBSw
pAC1380	pX-sgRNA-5xPBSc	Cloning vector for expression of sgRNA-5xPBSc
pAC1381	PB3-neo(-)-pmax-	Clover_NLSPUFc in pAC1119
1.01202	Clover_NLSPUFc	NI ODIJE (SHOEL)
pAC1393	pmax- NLSPUFa_p65HSF1	NLSPUFa_p65HSF1 in pmax expression vector
pAC1394	pX-sgRNA-0xPBS	Cloning vector for expression of sgRNA without PBS. It contains extra sequences for BsaI digestion for insertion of PBS
pAC1399	pX-sgRNA-20xPBSc	Cloning vector for expression of sgRNA-20xPBSc
pAC1402	pCR8-Clover_NLSPUFa	Clover_NLSPUFa in pCR8 gateway donor vector
pAC1403	pCR8-Clover_NLSPUFc	Clover_NLSPUFc in pCR8 gateway donor vector
pAC1404	pCR8- mRuby2_NLSPUFa	mRuby2_NLSPUFa in pCR8 gateway donor vector
pAC1405	pCR8- 4xNLS_PUFa_2xNLS	NLSPUFa pCR8 gateway donor vector for insertion of N-terminal domain (SgrAI or AgeI with ClaI) and C-terminal domain (FseI PacI). Grow in dcm- cells (e.g., NEB C3040) to prepare vector for ClaI digestion
pAC1406	pCR8- 4xNLS_PUFb_2xNLS	NLSPUFb pCR8 gateway donor vector for insertion of N-terminal domain (SgrAI or AgeI with ClaI) and C-terminal domain (FseI PacI). Grow in dcm- cells (e.g., NEB C3040) to prepare vector for ClaI digestion
pAC1407	pCR8- 4xNLS_PUFw_2xNLS	NLSPUFw pCR8 gateway donor vector for insertion of N-terminal domain (SgrAI or AgeI with ClaI) and C-terminal domain (FseI PacI). Grow in dcm- cells (e.g., NEB C3040) to prepare vector for ClaI digestion
pAC1408	pCR8- 4xNLS_PUFc_2xNLS	NLSPUFc pCR8 gateway donor vector for insertion of N-terminal domain (SgrAI or AgeI with ClaI) and C-terminal domain (FseI PacI). Grow in dcm- cells (e.g., NEB C3040) to prepare vector for ClaI digestion
pAC1410	pmax- dCas9Master_p65HSF1	dCas9Master_p65HSF1 in pmax expression vector
pAC1411	pmax- NLSPUFc_p65HSF1	NLSPUFc_p65HSF1 in pmax expression vector
pAC1412	PB3-neo(-)-pmax- NLSKRAB_NLSPUFa	NLSKRAB_NLSPUFa in pAC1119
pAC1413	PB3-neo(-)-pmax- NLSPUFb_p65HSF1	NLSPUFb_p65HSF1 in pAC1119
pAC1414	PB3- NLSPUFb_p65HSF1- neo(-)-	Dual expression vector for NLSPUFb_p65HSF1 and BFPKRAB_NLSPUFa

pAC number	Descriptive name	Description
	BFPKRAB_NLSPUFa	
pAC1415	pCR8-	4xNLS_PUFa_2xNLS_mCBPHAT in pCR8
1	4xNLS_PUFa_2xNLS_m	Gateway donor vector
	СВРНАТ	·
pAC1416	pCR8-	mCBPHAT_4xNLS_PUFa_2xNLS in pCR8
	mCBPHAT_4xNLS_PUF	Gateway donor vector
	a_2xNLS	
pAC1417	pmax-	4xNLS_PUFa_2xNLS_mCBPHAT in pmax
	4xNLS_PUFa_2xNLS_m	expression vector
	СВРНАТ	
pAC1418	pmax-	mCBPHAT_4xNLS_PUFa_2xNLS in pmax
	mCBPHAT_4xNLS_PUF	expression vector
1 01 110	a_2xNLS	10.015
pAC1419	PB3-neo(-)-pmax-	dCas9Master_mCherry in pAC1119
- A C1 420	dCas9Master_mCherry	Classic and Community of a DNA 1 DDC
pAC1420	pX-sgRNA-1xPBSa	Cloning vector for expression of sgRNA-1xPBSa
pAC1421	pX-sgRNA-2xPBSa	Cloning vector for expression of sgRNA-2xPBSa
pAC1422	pX-sgRNA-1xPBSb	Cloning vector for expression of sgRNA-1xPBSb
pAC1423	pX-sgRNA-2xPBSb	Cloning vector for expression of sgRNA-2xPBSb
pAC1424	pX-sgRNA-10xPBSb	Cloning vector for expression of sgRNA-20vPRSh
pAC1425	pX-sgRNA-20xPBSb	Cloning vector for expression of sgRNA-20xPBSb
pAC1426	pX-sgRNA-47xPBSb pX-sgRNA-10xPBSw	Cloning vector for expression of sgRNA-47xPBSb Cloning vector for expression of sgRNA-10xPBSw
pAC1427	pX-sgRNA-10xPBSw	Cloning vector for expression of sgRNA-10xFBSw
pAC1428	pX-sgRNA-13xFBSc	Cloning vector for expression of sgRNA-13xr BSw Cloning vector for expression of sgRNA-10xPBSc
pAC1429	pX-sgRNA-10x1BSc pX-sgRNA-15xPBSc	Cloning vector for expression of sgRNA-15xPBSc
pAC1430	PB3-LGFPL(-)-	Vector for expression of sgSOX2PP1-5xPBSa with a
p/101431	sgSOX2PP1-5xPBSa	GFP marker flanked by loxP sites
pAC1432	PB3-LGFPL(-)-	Vector for expression of sgSOX2PP2-5xPBSa with a
pricrisz	sgSOX2PP2-5xPBSa(-)	GFP marker flanked by loxP sites
pAC1433	PB3-LGFPL(-)-	Vector for expression of sgSOX2PP3-5xPBSa with a
Priorite	sgSOX2PP3-5xPBSa	GFP marker flanked by loxP sites
pAC1434	PB3-LGFPL(-)-	Vector for expression of sgSOX2PP4-5xPBSa with a
	sgSOX2PP4-5xPBSa	GFP marker flanked by loxP sites
pAC1435	PB3-LGFPL(-)-	Vector for expression of sgOCT4PP1-5xPBSb with a
_	sgOCT4PP1-5xPBSb	GFP marker flanked by loxP sites
pAC1436	PB3-LGFPL(-)-	Vector for expression of sgOCT4PP4-5xPBSb with a
	sgOCT4PP4-5xPBSb	GFP marker flanked by loxP sites
pAC1437	PB3-LGFPL(-)-	Vector for expression of sgOCT4PP3-5xPBSb with a
	sgOCT4PP3-5xPBSb	GFP marker flanked by loxP sites
pAC1438	PB3-LGFPL(-)-	Vector for expression of sgOCT4PP2-5xPBSb with a
	sgOCT4PP2-5xPBSb	GFP marker flanked by loxP sites

List of sgRNA-PBS expression vectors by number and type of PBS.

#PBS\PBSType	PUFa	PUFb	PUFw	PUFc
1x	pAC1420	pAC1422		
2x	pAC1421	pAC1423		
5x	pAC1371	pAC1374	pAC1379	pAC1380
10x		pAC1424	pAC1427	pAC1429
15x	pAC1372	pAC1375	pAC1428	pAC1430
20x		pAC1425		pAC1399
25x	pAC1373	pAC1376		
47x		pAC1426		

CLAIMS

- 1. A polynucleotide comprising:
 - a DNA-targeting sequence that is complementary to a target polynucleotide sequence;
 - (2) a Cas9-binding sequence; and,
 - (3) one or more copies of a PUF domain-Binding Sequence (PBS), wherein each of said one or more copies of the PBS binds to the same or a different PUF domain; wherein a Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) is capable of forming a complex with the polynucleotide by binding to the Cas9-binding sequence.
- 2. The polynucleotide of claim 1, wherein the Cas9 protein is a nuclease-deficient dCas9 protein that retains DNA-binding ability when complexed with the polynucleotide.
- 3. The polynucleotide of claim 1, wherein the DNA-targeting sequence base-pairs with the target polynucleotide sequence when the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) is complexed with the polynucleotide.
- 4. The polynucleotide of claim 1, wherein the target polynucleotide sequence comprises or is adjacent to a transcription regulatory element.
- 5. The polynucleotide of claim 4, wherein the transcription regulatory element comprises one or more of: core promoter, proximal promoter element, enhancer, silencer, insulator, and locus control region.
- 6. The polynucleotide of claim 1, wherein the target polynucleotide sequence comprises or is adjacent to a telomere sequence, a centromere, or a repetitive genomic sequence.
- 7. The polynucleotide of claim 1, wherein the target polynucleotide sequence comprises or is adjacent to a genomic marker sequence (or a genomic locus of interest).
- 8. The polynucleotide of claim 1, wherein the target polynucleotide sequence is immediately 3' to a PAM (protospacer adjacent motif) sequence of the complementary strand that is 5'-CCN-3', wherein N is any DNA nucleotide.
- 9. The polynucleotide of claim 1, wherein the DNA-targeting sequence is complementary to the target polynucleotide sequence over about 12-22 nucleotides (nts), about 14-20 nts, about 16-20 nts, about 18-20 nts, or about 12, 14, 16, 18, or 20 nts (preferably, the complementary region comprises a continuous stretch of 12-22 nts, preferably at the 3' end of the DNA-binding sequence).

10. The polynucleotide of claim 9, wherein the DNA-binding sequence is 50, 60, 70, 80, 90, or 95-100% complementary to the target polynucleotide sequence.

- 11. The polynucleotide of claim 1, wherein the DNA-binding sequence has a 5' end nucleotide G.
- 12. The polynucleotide of claim 1, further comprising a linker sequence linking the DNA-targeting sequence to the Cas9-binding sequence.
- 13. The polynucleotide of claim 1, wherein the Cas9-binding sequence forms a hairpin structure.
- 14. The polynucleotide of claim 1, wherein the Cas9-binding sequence is about 37-47 nt, or about 42 nt.
- 15. The polynucleotide of claim 1, wherein the Cas9 protein is a Cas9 nickase or a dCas9 protein that lacks endonuclease activity due to point mutations at one or both endonuclease catalytic sites (RuvC and HNH) of wild type Cas9.
- 16. The polynucleotide of claim 15, wherein the point mutations are D10A and H840A.
- 17. The polynucleotide of claim 1, wherein each of said one or more copies of the PBS has about 8 nucleotides.
- 18. The polynucleotide of claim 1, comprising 1, 2, 3, 4, 5, 10, 15, 20, 25, 30, 35, 40, 45, 46, 47, 48, 49, or 50 copies, or 1-50, 2-45, 3-40, 5-35, 5-10, 10-20 copies of identical or different PBS.
- 19. The polynucleotide of claim 1, comprising a PBS of the sequence 5'-UGUAUGUA-3' that can be bound by the PUF domain PUF(3-2).
- 20. The polynucleotide of claim 1, comprising a PBS of the sequence 5'-UUGAUAUA-3' that can be bound by the PUF domain PUF(6-2/7-2).
- 21. A vector encoding the polynucleotide of any one of claims 1-20.
- 22. The vector of claim 21, wherein transcription of the polynucleotide is under the control of a constitutive promoter, or an inducible promoter.
- 23. The vector of claim 20, wherein the vector is active in a cell from a mammal (a human; a non-human primate; a non-human mammal; a rodent such as a mouse, a rat, a hamster, a Guinea pig; a livestock mammal such as a pig, a sheep, a goat, a horse, a camel, cattle; or

- a pet mammal such as a cat or a dog); a bird, a fish, an insect, a worm, a yeast, or a bacterium.
- 24. A plurality of vectors of any one of claims 21-23, wherein two of the vectors differ in the encoded polynucleotides in their respective DNA-targeting sequences, Cas9-binding sequences, and/or the copy number, identity, or relative order of the PBS.
- 25. A complex comprising the polynucleotide of any one of claims 1-20, and the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein).
- 26. The complex of claim 25, further comprising one or more PUF domain(s) bound to said one or more PBS(s).
- 27. The complex of claim 26, wherein each of said PUF domains is fused to an effector domain.
- 28. The complex of claim 27, wherein the effector domain is independently a transcription repressor, a transcription activator, a fluorescent protein, an enzyme, or a chromatin remodeling protein (HDAC/HAT).
- 29. The complex of claim 27, wherein at least two of the PUF domains are fused to different effector domains.
- 30. The complex of any one of claims 25-29, wherein the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), the PUF domain, and/or the effector domain further comprises a nuclear localization sequence (NLS).
- 31. The complex of any one of claims 25-30, which is bound to the target polynucleotide sequence through the DNA-targeting sequence.
- 32. A host cell comprising the vector of any one of claims 21-23, or the plurality of vectors of claim 24.
- 33. The host cell of claim 32, further comprising a second vector encoding the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein).
- 34. The host cell of claim 33, wherein the second vector further encodes an effector domain fused to the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein).
- 35. The host cell of claim 33, wherein expression of the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) is under the control of a constitutive promoter or an inducible promoter.
- 36. The host cell of any one of claims 32-35, further comprising a third vector encoding said

- one or more PUF domains, each fused to an effector domain.
- 37. The host cell of claim 36, wherein expression of said one or more PUF domains is independently under the control of a constitutive promoter or an inducible promoter.
- 38. The host cell of any one of claims 34-37, wherein the effector domain is a transcription repressor, a transcription activator, a fluorescent protein, an enzyme, or a chromatin remodeling protein (HDAC/HAT).
- 39. The host cell of any one of claims 33-38, wherein the second vector further encodes a nuclear localization signal fused to the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein) or the effector domain, and/or the third vector further encodes a nuclear localization signal fused to the PUF domain or the effector domain.
- 40. The host cell of any one of claims 33-39, wherein the second vector is the same as the vector, and/or wherein the third vector is the same as the vector or the second vector.
- 41. The host cell of any one of claims 32-40, which is in a live animal.
- 42. The host cell of any one of claims 32-40, which is a cultured cell.
- 43. A method of assembling the complex of any one of claims 26-31 at the target polynucleotide sequence, the method comprising contacting or bringing to the vicinity of the target polynucleotide sequence:
 - (1) any one of the polynucleotide of claims 1-20, or any one of the vector of claims 21-23, or the plurality of vector of claim 24;
 - (2) the Cas9 protein (*e.g.*, wt, nickase, or dCas9 protein), or any one of the second vector of claims 33-35 and 38-39; and,
 - (3) one or more of the PUF domains, each fused to an effector domain, or any one of the third vector of claims 36-38 and 40.
- 44. The method of claim 43, wherein the complex is assembled inside a cell, the target polynucleotide sequence is a part of the genomic DNA of the cell, and wherein the vector of claims 21-23, the second vector of claims 33-35 and 38-39, and the third vector of claims 36-38 and 40 are introduced into the cell.
- 45. The method of claim 43, wherein the target polynucleotide sequence is at or near a genomic locus rich in heterochromatin, and wherein the effector domain is a detectable marker (*e.g.*, a fluorescent protein).
- 46. The method of claim 43, wherein the target polynucleotide sequence is at or near a

transcription regulatory element of a target gene, and wherein the effector domain is a transcription modulator (*e.g.*, activator, suppressor).

- 47. The method of claim 46, wherein transcription of the target gene affects cell fate determination, cell differentiation, metabolic flux, or a biologically or biochemically determinable outcome.
- 48. A method of modulating transcription or epigenetic state, respectively, of a plurality of target genes in a cell, the method comprising: introducing into the cell the plurality of the vector of claim 24, a coding sequence for a dCas9 protein or a wt Cas9, respectively, and a coding sequence for one or more PUF domains, wherein each of said target genes comprises a target polynucleotide sequence that permits (1) the assembly, at the target polynucleotide sequence, of a tripartite complex of a polynucleotide encoded by one of said plurality of the vector, the dCas9 protein or the wt Cas9, and a PUF domain; and (2) transcription or epigenetic state modulation of the target gene comprising the target polynucleotide sequence.
- 49. The method of claim 48, wherein the transcription of at least one target gene is enhanced/stimulated, while the transcription of at least another target gene is inhibited.
- 50. A kit comprising:
 - (1) a polynucleotide of any one of claims 1-20, or a vector of any one of claims 21-23;
 - (2) a second vector encoding the Cas9 protein (e.g., wt, nickase, or dCas9 protein); and
 - (3) a third vector encoding one or more PUF domains, each fused to an effector domain.
- 51. The kit of claim 50, further comprising transformation, transfection, or infection reagents to facilitate the introduction of said vectors into a cell.

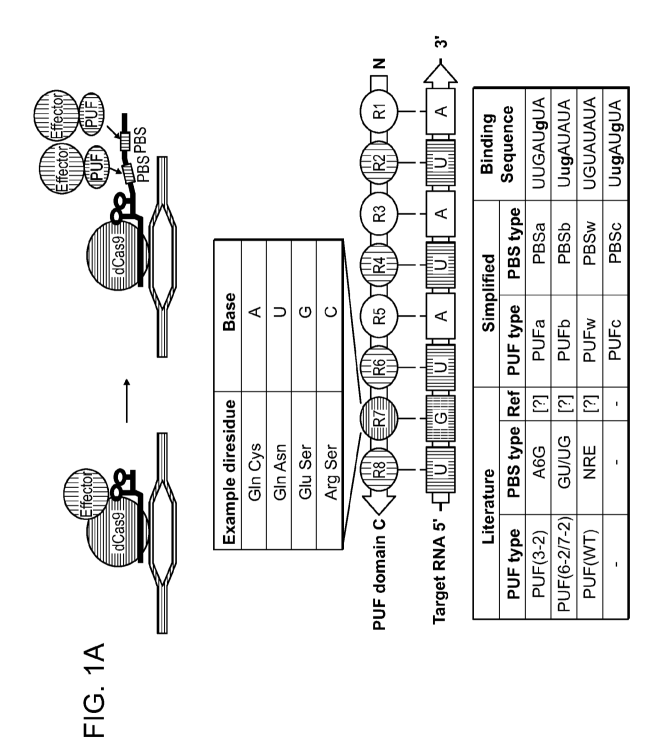
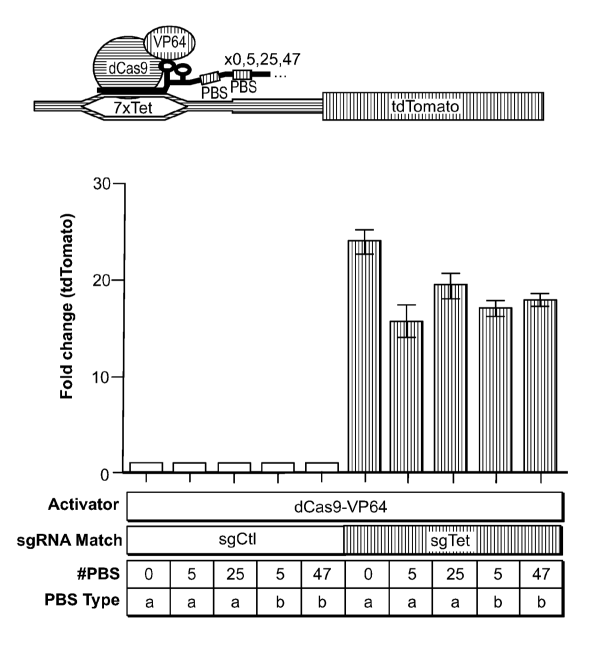
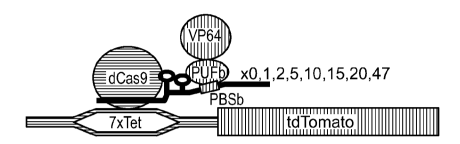


FIG. 1B



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FIG. 1C



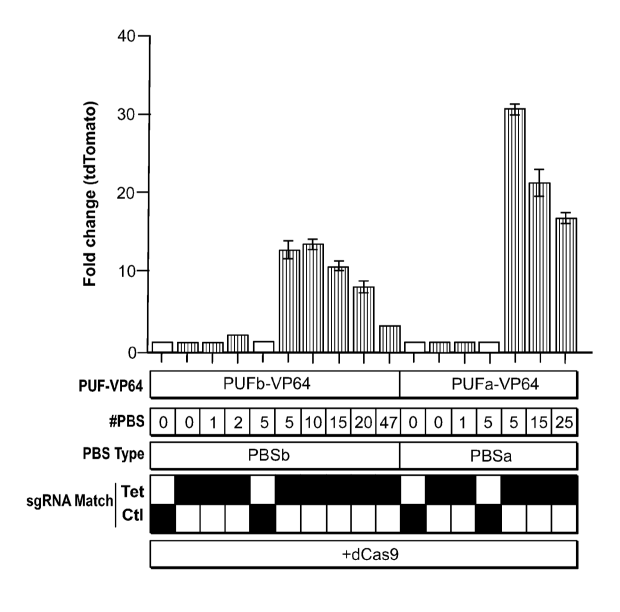
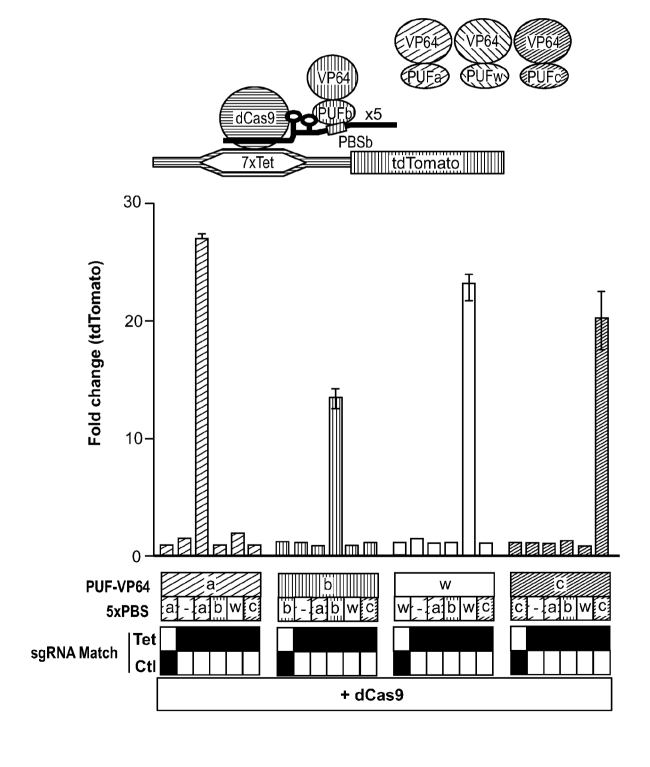


FIG. 1D



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FIG. 2A

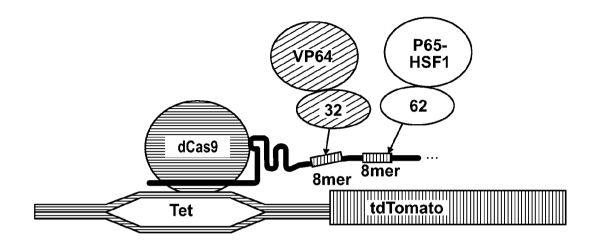
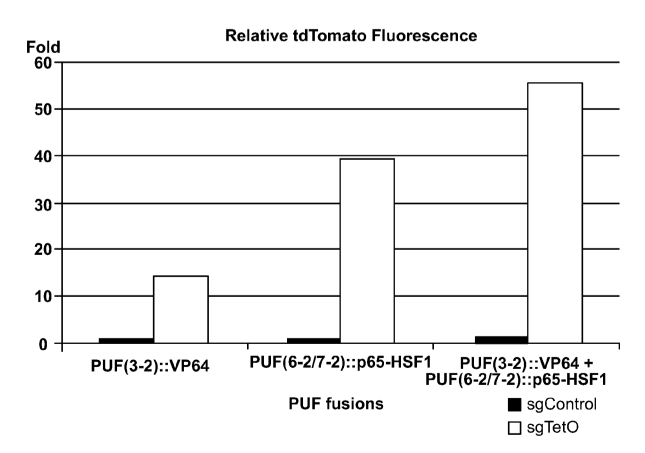
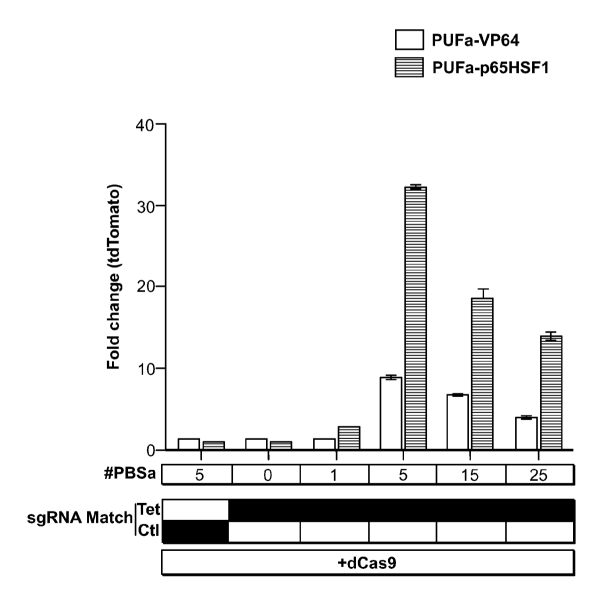


FIG. 2B



dCas9 + PUF fusion(s) + sgX-4x(PBS32-PBS6272)

FIG. 2C



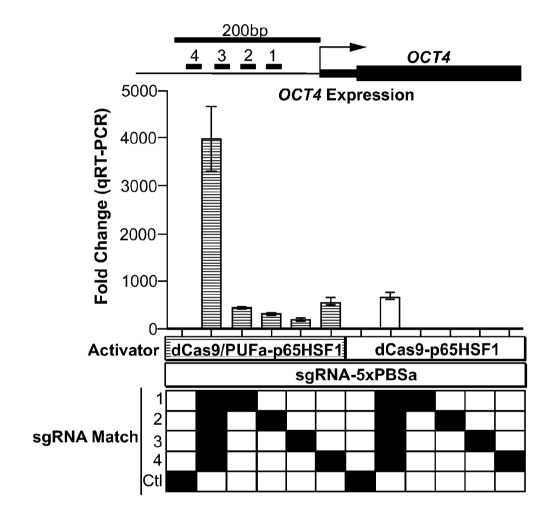


FIG. 3A

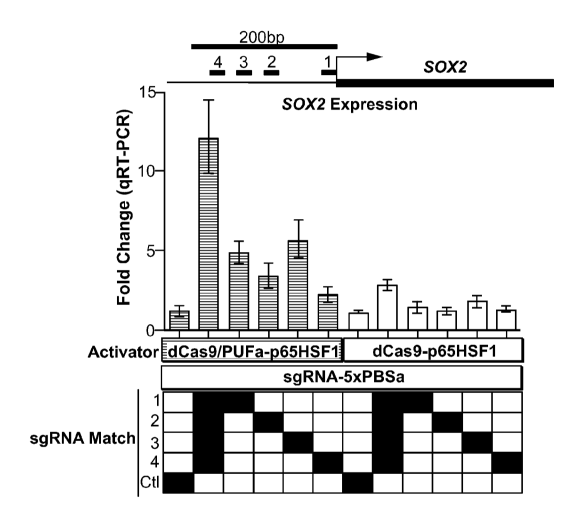
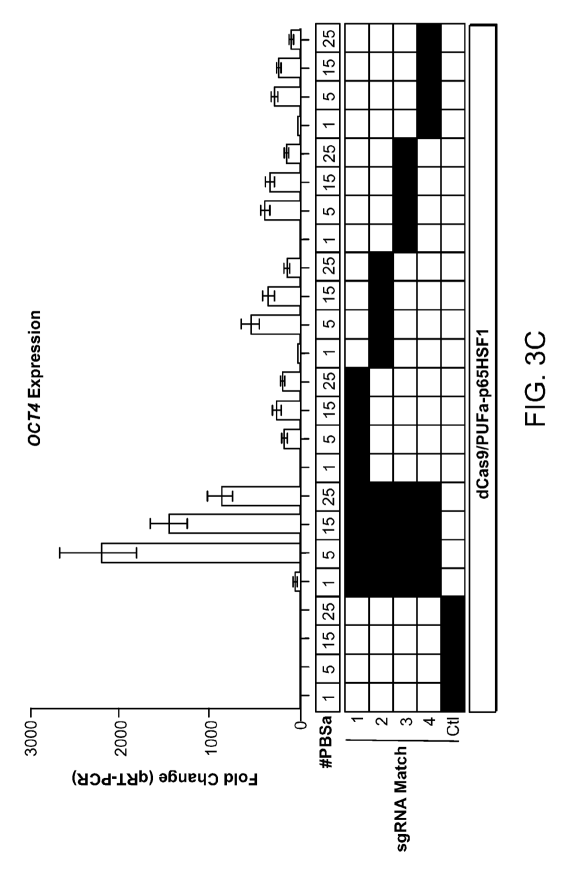
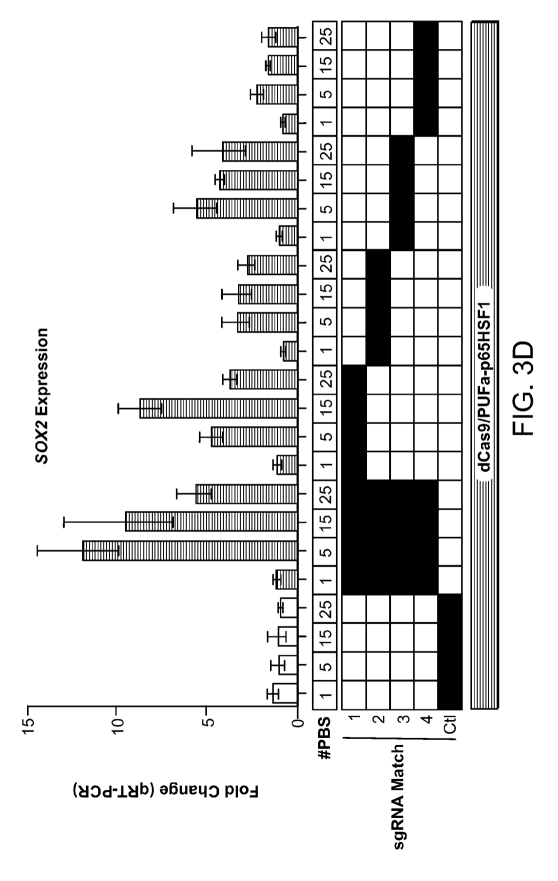


FIG. 3B

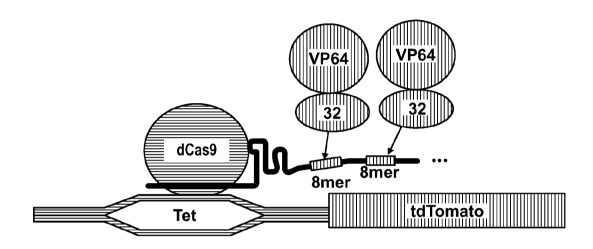


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FIG. 4A



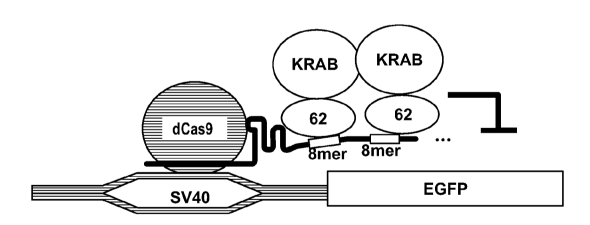
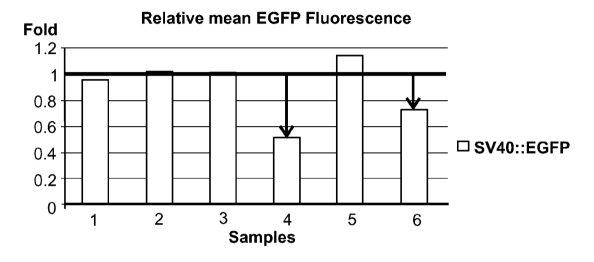
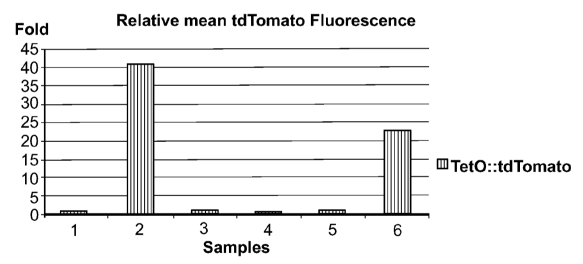
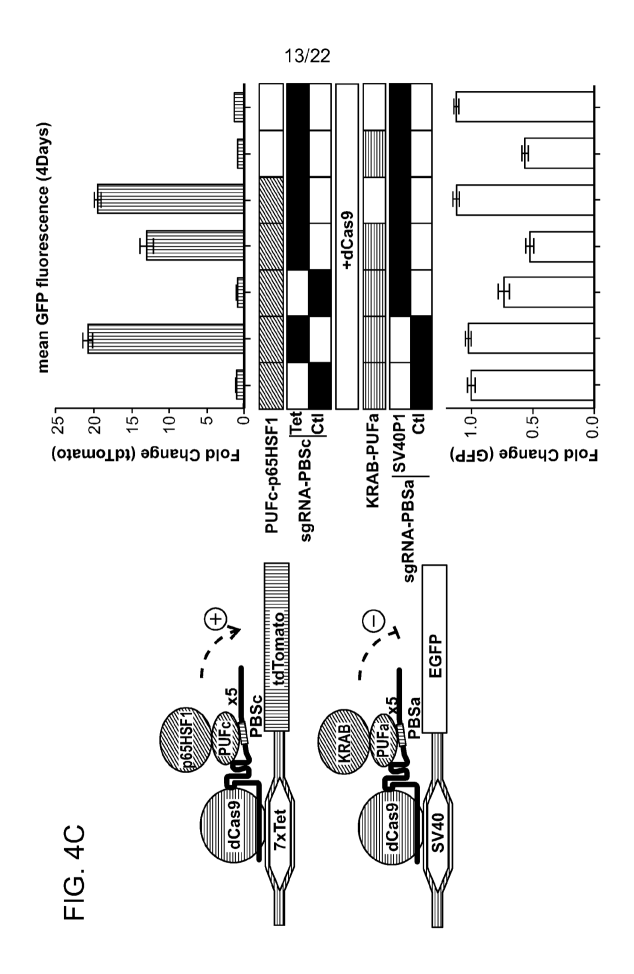


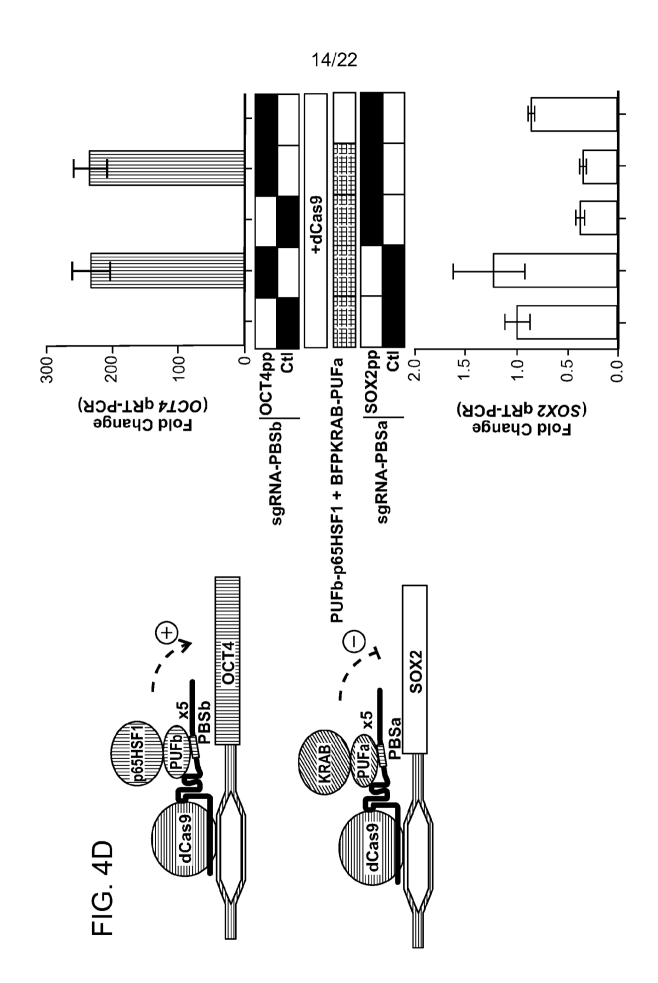
FIG. 4B

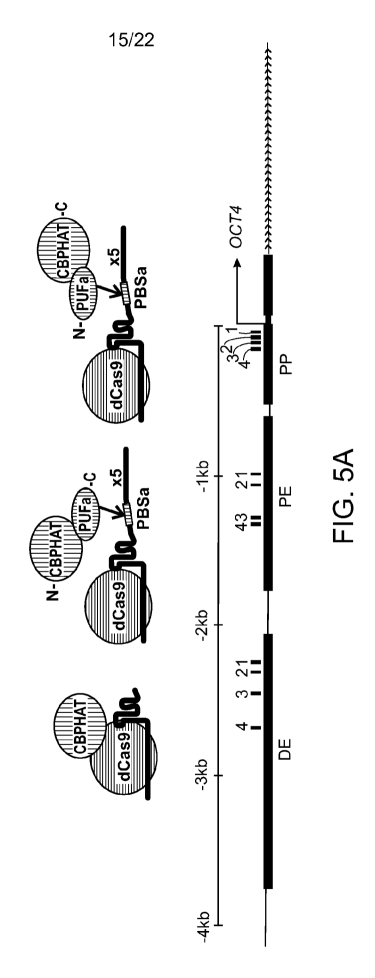




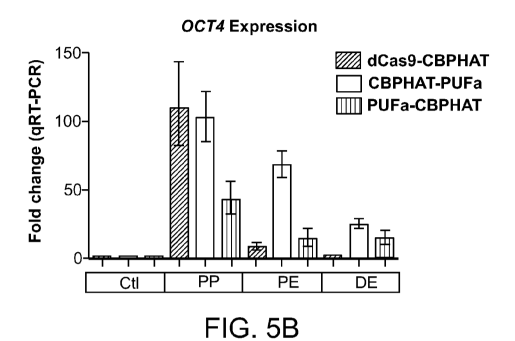
	Samples
1	dCas9/sgControl-5xPBS32/PUF(3-2)::VP64
2	dCas9/sgTetO-5xPBS32/PUF(3-2)::VP64
3	dCas9/sgControl-5xPBS6272/KRAB::PUF(6-2/7-2)
4	dCas9/sgSV40-5xPBS6272/KRAB::PUF(6-2/7-2)
5	dCas9/sgControl-5xPBS32/PUF(3-2)::VP64/sgControl-5xPBS6272/KRAB::PUF(6-2/7-2)
6	dCas9/sgTetO-5xPBS32/PUF(3-2)::VP64/sgSV40-5xPBS6272/KRAB::PUF(6-2/7-2)







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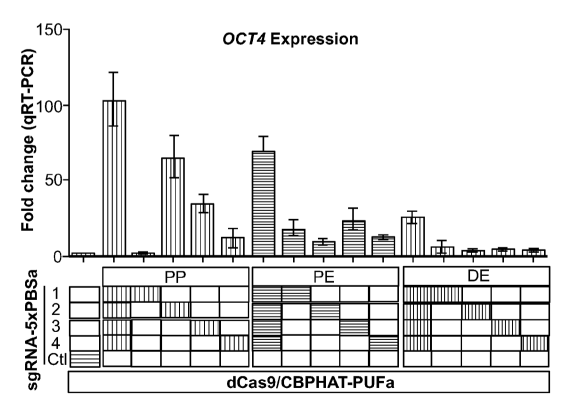


FIG. 5C

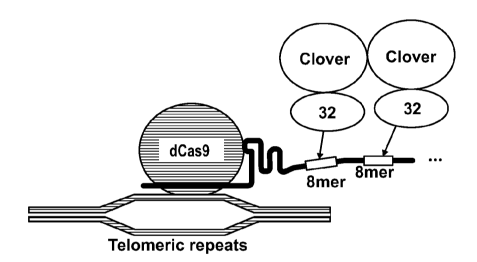
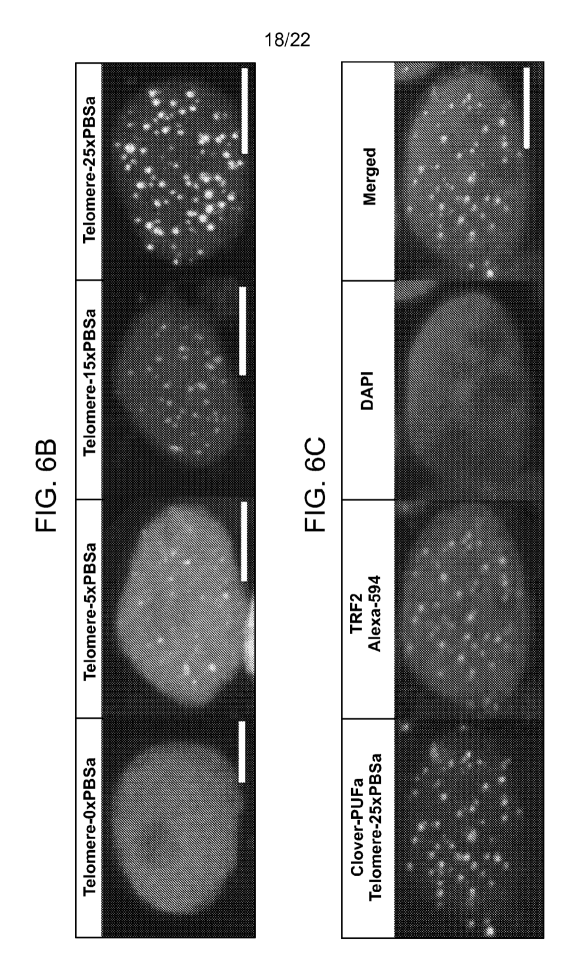


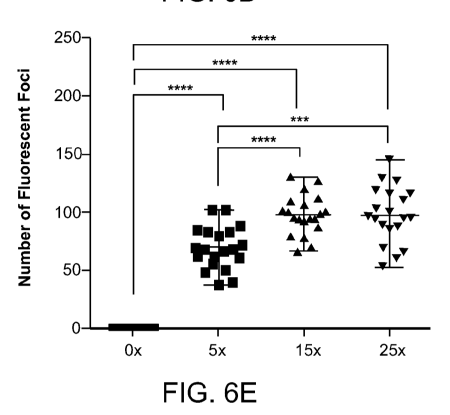
FIG. 6A

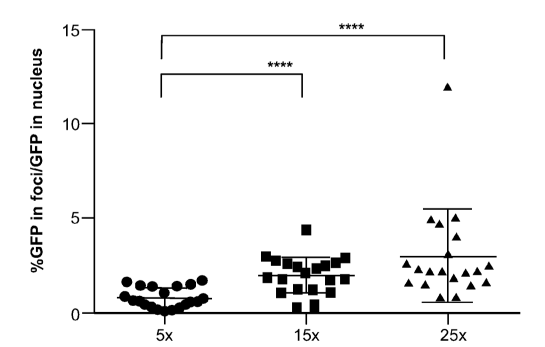


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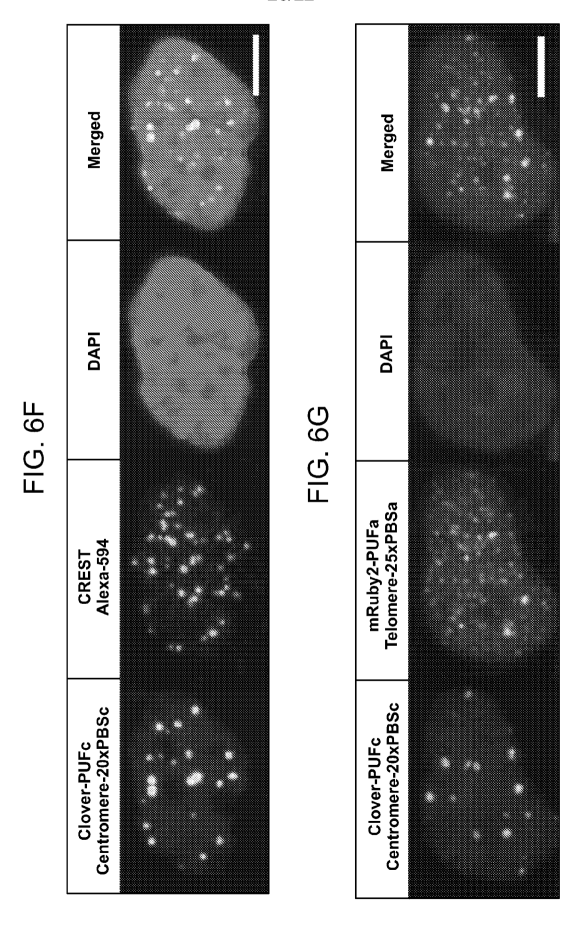


FIG. 6D









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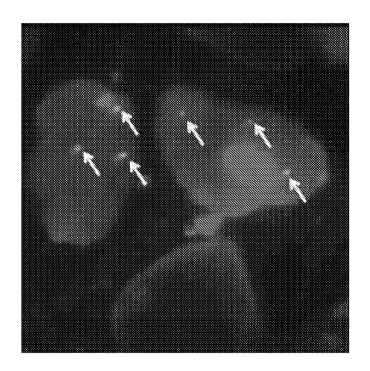


FIG. 7

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FIG. 8A

Multiplexing

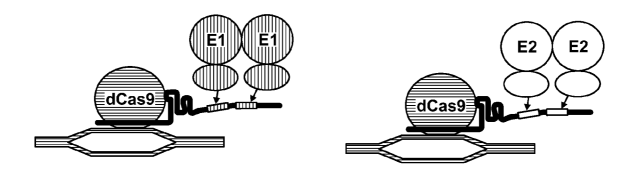


FIG. 8B

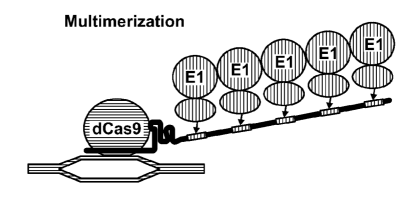
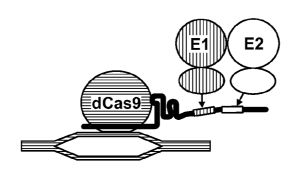


FIG. 8C

Complex formation



INTERNATIONAL SEARCH REPORT

International application No PCT/US2016/021491

A. CLASSIFICATION OF SUBJECT MATTER INV. C12N15/11

ADD.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols) C12N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

EPO-Internal, WPI Data, BIOSIS, EMBASE

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X, P	ALBERT W. CHENG ET AL.: "Casilio: a versatile CRISPR-Cas9-Pumilio hybrid for gene regulation and genomic labeling", CELL RESEARCH - XIBAO YANJIU, vol. 26, no. 2, 15 January 2016 (2016-01-15), pages 254-257, XP055278705, GB, CN ISSN: 1001-0602, DOI: 10.1038/cr.2016.3 the whole document -& Cheng: "Supplementary Information",	1-51 1-51
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X Further documents are listed in the continuation of Box C.	X See patent family annex.		
"A" document defining the general state of the art which is not considered to be of particular relevance "E" earlier application or patent but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "P" document published prior to the international filing date but later than the priority date claimed	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art "&" document member of the same patent family		
Date of the actual completion of the international search 13 June 2016	Date of mailing of the international search report $17/06/2016$		
Name and mailing address of the ISA/ European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016	Authorized officer Macchia, Giovanni		

INTERNATIONAL SEARCH REPORT

International application No
PCT/US2016/021491

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