


Multiplexed genome engineering by Cas12a and CRISPR arrays encoded on single transcripts

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Campa, Carlo C.; Weisbach, Niels R.; Santinha, António J.; Incarnato, Danny; [Platt, Randall](#) 

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2 **single transcripts**

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4 Carlo C. Campa^{1*}, Niels R. Weisbach^{1*}, António J. Santinha¹, Danny Incarnato², Randall J. Platt^{1,3†}
5
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7
8

9 ¹Department of Biosystems Science and Engineering
10 ETH Zurich
11 4058 Basel, Switzerland
12

13 ²Department of Molecular Genetics
14 Groningen Biomolecular Sciences and Biotechnology Institute
15 University of Groningen
16 Nijenborgh 7, 9747 AG Groningen, Netherlands
17

18 ³Department of Chemistry
19 University of Basel
20 4003 Basel, Switzerland
21
22
23
24
25
26
27

28 *These authors contributed equally to this work.

29 †To whom correspondence should be addressed: rplatt@ethz.ch.
30

31 **Abstract**

32 The ability to modify multiple genetic elements simultaneously would help to elucidate and control
33 the gene interactions and networks underlying complex cellular functions. However, current
34 genome engineering technologies are limited in both the number and the type of perturbations that
35 can be performed simultaneously. Here, we demonstrate that both Cas12a and large CRISPR array
36 can be encoded in a single transcript by adding a stabilizer tertiary RNA structure. By leveraging
37 this system, we illustrate constitutive, conditional, inducible, orthogonal and highly multiplexed
38 genome engineering using a single plasmid as a delivery agent. Our method provides a powerful
39 platform to investigate and orchestrate the sophisticated genetic programs underlying complex cell
40 behaviours.

41

42 **Introduction**

43 Clustered Regularly Interspaced Short Palindromic Repeats (CRISPR)-associated (Cas)
44 nucleases are versatile RNA-guided enzymes that facilitate a wide-range of genome engineering
45 applications^{1,2}. Through the binding of short RNA molecules, known as CRISPR RNAs
46 (crRNAs), members of the Cas nuclease family are programmed to edit specific genomic loci
47 thereby facilitating systematic investigation of gene function^{3,4}. In addition, nuclease-inactive
48 Cas enzymes fused to transcriptional effectors enable fine control of gene expression⁵⁻¹⁰.
49 Remarkably, use of either different Cas enzymes or engineered crRNAs enable distinct gene
50 perturbations, including gene knockout, gene activation, and gene repression¹¹⁻¹⁶. Despite this
51 versatility, either the heterologous expression of different proteins (e.g. Csy4, scFV) or the
52 presence of long RNA-based regulatory regions (e.g. ribozymes, aptamers) are required to
53 permit regulation of multiple genes using current platforms¹⁷⁻¹⁹. This inevitably limits the
54 scalability of CRISPR-based multiplexed genome engineering approaches and consequently the
55 possibility to investigate complex cell behaviours.

56
57 In order to coordinate a myriad of different processes using a limited number of cellular elements,
58 cells evolved to maximize the efficiency of their molecular components. This principle is adopted
59 by distinct organisms to rapidly match different environmental contexts, thereby supporting the
60 notion that maximizing the performance of each molecular element is a general principle on which
61 biological systems based their own survival²⁰. As an example, protection against bacteriophages
62 and other foreign genetic elements is mediated by Cas12a, a class II type V CRISPR-Cas
63 system^{21,22}. Functioning as a both RNase and DNase, Cas12a controls both processing and
64 maturation of its own crRNA as well as DNA target cleavage²¹⁻²³. CRISPR arrays associated
65 with Cas12a are transcribed as a long RNA transcript, termed pre-crRNA, that contains a
66 succession of ~30-nucleotide (nt) spacer sequences, separated by 36-nt direct repeat sequences
67 (DRs)^{21,22}. While DRs direct Cas12a-mediated pre-crRNA processing into mature crRNAs,
68 recognition and cleavage of target DNA is controlled by Watson-Crick base pairing between the
69 spacer and target sequences^{21,22}. Transplantation of this microbial immune system into mammalian
70 cells enables genome engineering applications, including gene editing and transcriptional gene
71 control²⁴. Optimization of this genome editing system for mammalian cells was obtained by DR
72 length reduction and adoption of distinct promoters that differentially drive transcription of Cas12a
73 and pre-crRNA²⁵⁻²⁷. Although, the implementation of these strategies enables a wide range of
74 applications, they fail to harness the full potential of Cas12a in genome engineering.

75

76 Here, we leverage the dual RNase/DNase function of *Acidaminococcus sp.* Cas12a to develop a
77 robust system for multiplexed genome engineering using a single plasmid as a delivery agent. This
78 system, termed SiT-Cas12a, encodes Cas12a and dozens of crRNAs in a single transcript.
79 Stabilization of SiT-Cas12a transcripts through inclusion of a tertiary structural motif improves
80 both pre-crRNA processing as well as Cas12a production. When coupled with transcriptional
81 effectors and engineered spacer sequences, SiT-Cas12a enables multiplexed orthogonal gene
82 transcriptional regulation and editing thereby providing a scalable way to elucidate and control the
83 gene networks underlying cellular functions.

84

85 **Results**

86 ***Compact encoding of Cas12a and CRISPR arrays in a single Pol II-derived transcript***

87 In mammalian cells, distinct promoters control transcription of different RNA molecules. While
88 Pol II promoters are mainly used for transcription of coding genes characterized by long RNA
89 sequences, Pol III promoters are employed for production of small non-coding RNAs including
90 CRISPR RNAs (crRNAs)²⁸. To assess whether transcripts derived from Pol II promoters could
91 facilitate Cas12a-based genome engineering applications²⁶, we expressed a crRNA targeting
92 *DNMT1* from either a U6 (Pol III) or EF1a (Pol II) promoter along with ectopically expressed
93 Cas12a followed by quantification of insertions and deletions (indels) 72 hours after transient
94 transfection. Expression of crRNA from both Pol II and Pol III promoters resulted in comparable
95 gene editing efficiencies (Figure 1A, B, conditions I, II), indicating that Cas12a processes and
96 utilizes crRNAs derived from both promoter types.

97

98 To determine whether Pol II promoters facilitate simultaneous protein and crRNA expression, we
99 cloned a CRISPR array containing a spacer targeting *DNMT1* within the 3'UTR of the enhanced
100 green fluorescence protein (*EGFP*) gene and assessed both gene editing efficiency at the
101 endogenous *DNMT1* locus and EGFP expression 72 hours after transient transfection in HEK 293T
102 cells (Figure 1B, condition III). In cells harbouring *EGFP* transcripts containing CRISPR arrays, we
103 observed complete loss of EGFP expression (Figure 1B, condition III), suggesting destabilization of
104 *EGFP* transcripts mediated by the RNase activity of Cas12a. Consistently, expression of an RNase
105 dead Cas12a (rdCas12a), but not DNase dead Cas12a (ddCas12a), rescued EGFP expression
106 (Figure 1B, condition IV, V). Taken together, this suggests that Cas12a-mediated crRNA
107 processing via the RNase domain results in efficient cleavage and destabilization of protein-coding
108 mRNAs, which is likely a result of removal of the polyadenylation (poly(A)) tail.

109

110 To overcome mRNA destabilization and enable simultaneous expression of protein and crRNA on
111 the same transcript, we leveraged a 110-nt structure derived from the 3' end of the mouse non-
112 coding RNA Metastasis-associated lung adenocarcinoma transcript 1 (*Malat1*), previously
113 described to stabilize transcripts lacking poly(A) tails through the formation of a tertiary structure
114 element termed “Triplex”²⁹. We cloned the Triplex sequence between the *EGFP* coding sequence
115 and the CRISPR array (Figure 1B, condition VI), which effectively rescued EGFP expression
116 without affecting gene editing efficiency (Figure 1B, condition VI). These results indicate that a
117 Triplex sequence positioned at the 3' of a protein coding gene stabilizes mRNAs after Cas12a-
118 mediated RNA processing, enabling concomitant protein expression and gene editing.

119 120 ***Constitutive, conditional and inducible gene editing with SiT-Cas12a***

121 To determine whether Cas12a and CRISPR arrays could be compactly encoded on single Pol II-
122 driven mRNA, we developed single-transcript Cas12a (SiT-Cas12a). SiT-Cas12a is composed of a:
123 i) Pol II promoter EF1a; ii) Cas12a derived from *Acidaminococcus sp.*; iii) Triplex sequence; iv)
124 CRISPR array containing spacers targeting a set of mammalian genomic loci; and v) poly(A) signal
125 (Figure 1A). We evaluated the platform using a CRISPR array containing a spacer targeting
126 *DNMT1* and observed consistent and efficient gene editing at the *DNMT1* locus (Figure S1A).

127
128 Utilization of a single Pol II promoter offers the unique potential for more sophisticated and
129 simultaneous control of both Cas12a and crRNAs expression. To evaluate this possibility, we
130 generated conditional and inducible SiT-Cas12a platforms, termed SiT-Cas12a-[Cond] (Figure
131 S1B) and SiT-Cas12a-[Ind], respectively (Figure S1C). SiT-Cas12a-[Cond] relies on a Lox-Stop-
132 Lox (LSL) cassette positioned downstream of an EF1a promoter and upstream of the SiT-Cas12a
133 coding region. To demonstrate conditional genome editing, we co-transfected HEK 293T cells with
134 SiT-Cas12a-[Cond] and either a Cre-recombinase encoding plasmid or a control plasmid. At 72
135 hours post-transfection, we detected genome editing events exclusively in Cre-recombinase
136 expressing cells (Figure S1B). SiT-Cas12a-[Ind] relies on a Tetracycline responsive element (TRE)
137 positioned upstream of a minimal CMV promoter (minCMV) that, differently from the constitutive
138 promoter EF1a, resulted in gene editing efficiencies proportional to inducer concentration
139 (doxycycline) both in a Tet-on and Tet-off configuration (Figure S1C). Taken together, SiT-Cas12a
140 enables either constitutive or conditional and inducible gene editing through fine temporal control
141 of Cas12a and crRNA expression.

142 143 ***Multiplexed gene editing with SiT-Cas12a***

144 We evaluated the potential of the SiT-Cas12a platform for multiplexed gene editing using a
145 CRISPR array containing 5 distinct spacers targeting different genomic loci (*FANCF1*, *EMX1*,
146 *GRIN2B*, *VEGF*, *DNMT1*) and quantified both mature crRNAs and gene editing efficiency (Figure
147 2A). In the SiT-Cas12a context, we observed expression and processing of each mature crRNA,
148 albeit with varying efficiencies as observed in previous studies²⁵ (Figure 2A). In addition,
149 transcripts stabilized by the Triplex sequence (Figure 2A, SiT-Cas12a) increased both Cas12a
150 expression (Figure S2A) and processed crRNA abundance compared to transcripts without the
151 Triplex (Figure 2A, Cas12a). In line with these results, the gene editing efficiency was higher in
152 cells expressing SiT-Cas12a compared to a control lacking the Triplex structure between Cas12a
153 and the CRISPR array (Figure 2B). The increased crRNA production facilitated by the Triplex
154 sequence disappeared upon mutation of the RNase domain of Cas12a (SiT-rdCas12a) (Figure 2A).
155 Consistently, gene editing efficiencies were negligible for the SiT-Cas12a RNase inactive mutant
156 and found to be comparable to the DNase inactive mutant of Cas12a (SiT-ddCas12a) (Figure 2B).

157
158 Next, we compared the gene editing efficiency of SiT-Cas12a with previously reported Cas12a
159 platforms based on independent transcription of Cas12a and a CRISPR array from distinct
160 promoters^{25,26}. Expression of SiT-Cas12a resulted in gene editing efficiencies equal to or higher
161 than other tested platforms (Figure 2C). Taken together, these data demonstrate that compact
162 encoding of Cas12a and a CRISPR array on Pol II transcripts mediates efficient multiplexed gene
163 editing.

164
165 In contrast to Pol III promoters, Pol II promoters express transcripts of seemingly unlimited
166 length³⁰. To leverage this property, we cloned a CRISPR array harbouring 10 distinct spacer
167 sequences targeting the *CD47* locus in the SiT-Cas12a context, either singularly or jointly, and
168 performed gene editing quantification (Figure S2B-G). While the gene editing efficiency using
169 single-crRNAs ranged from 2 to 17%, simultaneous expression of all crRNAs increased the gene
170 editing efficiency up to 60% (Figure S2C), indicating that the targeting of multiple crRNAs in the
171 same coding gene introduced more loss of function mutations. We obtained similar gene editing
172 efficiency in cells infected with a SiT-Cas12a-based lentiviral vector enabling stable expression and
173 delivery to cells that are difficult to transfect (Figures S3A-E). In agreement, multiple independent
174 indels at each target site as well as large fragment deletions between target sites were detected using
175 deep sequencing in cells expressing such large CRISPR array (Figure S2D-G), confirming the
176 targeting of all crRNAs used. Consistently, single cell analysis of CD47 protein expression showed

177 a 4-fold increase in CD47-negative cells, reaching 37% of the total population when using multiple
178 crRNAs (Figure S4A).

179

180 Finally, we evaluated whether this strategy could be employed to increase the efficiency of
181 generating multi-gene knockouts after a simple transient transfection experiment. We cloned
182 distinct spacers targeting different coding genes (*CD47*, *CD166* and *CD97*) (Figure S4B) in the
183 SiT-Cas12a context and measured the rate of either single (either CD47 or CD166 or CD97) double
184 (CD47/CD166, CD47/CD97, CD166/CD97) or triple (CD47/CD166/CD97) knockout by single cell
185 flow cytometry analysis. Targeting multiple sites per gene using SiT-Cas12a increased the rate of
186 single, double and triple multiplexed gene knockout generation by 2- to 3-fold compared to single
187 crRNA conditions (Figure S4B). Overall, these results demonstrate that SiT-Cas12a facilitates
188 scalable and multiplexed genome engineering.

189

190 ***Multiplexed transcriptional regulation with SiT-Cas12a***

191 By leveraging on the simultaneous transcription of both Cas12a and a large CRISPR array under a
192 single Pol II promoter, we set out to develop SiT-Cas12a-based platforms to control the expression
193 of endogenous genes. We fused either one, two, or three copies of the Krüppel associated box
194 (KRAB) domain of the transcriptional repressor ZNF10³¹ to the C-terminus of ddCas12a, thus
195 generating ddCas12a-KRAB₁, ddCas12a-KRAB₂, and ddCas12a-[Repr], respectively (Figure S5A).
196 To assess the efficiency of the transcriptional repression conferred by multiple KRAB domains, we
197 co-transfected the SiT-Cas12a repressor variants along with different concentrations of CRISPR
198 arrays containing spacers targeting 4 distinct genomic loci (*RAB5A*, *RAB7A*, *EEA1*, *PIK3C3*). We
199 found that three tandem KRAB domains fused at the C-terminus of ddCas12a (ddCas12a-[Repr])
200 conferred potent transcriptional repression (Figure 3A, S5B) with differential efficiencies according
201 to the target gene and distance from the TSS (Figure S5C). Next, we cloned a CRISPR array
202 harbouring 20 different spacers targeting 10 distinct genes (*RAB5A*, *RAB7A*, *PIK3C3*, *EEA1*,
203 *RAB9A*, *CHRM3*, *PLCB1*, *PRKCI*, *FZD1*) within SiT-ddCas12a-[Repr] and quantified mature
204 crRNAs and transcriptional repression. Small-RNA-seq analysis confirmed generation of mature
205 crRNAs leading to efficient gene repression ranging from 40 to 80% reduction (Figures 3B, C).

206 To further strengthen the finding that SiT-Cas12a enables simplified multi-gene transcriptional gene
207 control, we replaced the transcriptional repression domain KRAB with either the VPR activator³² or
208 a combination of the p65 activation domain (p65) together with the Heat Shock Factor 1 (HSF1)³ to
209 generate two RNA-guided transcriptional activators: ddCas12a-VPR and ddCas12a-p65-HSF1
210 (Figure S6A). Both of these Cas12a-based chimeric proteins induced gene activation at different

211 efficiencies (Figure S6B). Comparative analyses of these gene activators (Figure 4A, B) showed a
212 10-fold increase in activation efficiency of ddCas12a-[Activ] compared to ddCas12a-VPR (Figure
213 S6B) also in the presence of limiting spacer concentrations (Figure 4A, S6C). We also observed that
214 transcriptional activation efficiency varies according to the target gene and distance from the TSS
215 (Figure S6D). Next, to explore the potential of multiplexed transcriptional activation in the SiT-
216 Cas12a context, we combined SiT-ddCas12a-[Activ] with a CRISPR array harbouring 20 spacers
217 targeting 10 distinct genes (*ASCL1*, *ZFP42*, *CCR4*, *CCR10*, *IL1B*, *IL1R2*, *HBB*, *CHRM4*, *HTR6*,
218 *ADRB2*) and quantified mature crRNAs and transcriptional activation. Small RNA-seq analysis
219 confirmed generation of all 20 mature crRNAs (Figure 4B). In addition, we measured robust gene
220 activation (10-1000-fold) for all target genes (Figure 4C, S6E). Overall, these data indicate that both
221 repressor and activator transcriptional domains when combined with SiT-Cas12a enable multi-gene
222 transcriptional control.

223 224 ***Orthogonal gene editing and transcriptional control with SiT-Cas12a***

225 The Cas12a endonuclease, similarly to Cas9, is characterized by unique DNA binding kinetics,
226 which enables binding while avoiding cleavage in the presence of truncated crRNAs^{13-15,33}. To fully
227 harness the potential of Cas12a for genome engineering, we set out to explore these unique
228 properties for generating a SiT-Cas12a-based platform that could facilitate orthogonal gene editing
229 and transcriptional gene control.

230
231 Towards developing such a platform, we assessed *Acidaminococcus sp.* Cas12a (*AsCas12a*)
232 processing efficiency using CRISPR arrays containing both long (20 bp) and short (15 bp) spacers
233 (Figure S7A). We measured 3- to 5-fold higher amounts of mature crRNAs in cells expressing
234 arrays containing short spacers compared to those with long spacers (Figure S7B). Furthermore,
235 both short and long spacers generated comparable transcriptional gene control when combining
236 SiT-Cas12a-[Repr], SiT-Cas12a-p65-HSF1, and SiT-Cas12a-[Activ] with 2 distinct CRISPR arrays
237 (Figure S7C, D). This strategy does not extend with a similar efficiency to *Lachnospiraceae*
238 *bacterium* Cas12a (*LbCas12a*), whereby crRNAs containing short spacers did not induce significant
239 gene activation (Figure S7E). Next, we evaluated the SiT-Cas12a platform in an orthogonal
240 transcriptional control and gene editing context (Figure 5A). We combined both active and inactive
241 DNase versions of SiT-Cas12a-based transcriptional repressor and activator (SiT-ddCas12a-[Repr],
242 SiT-Cas12a-[Repr], SiT-ddCas12a-[Activ] and SiT-Cas12a-[Activ]) with 2 sets of CRISPR arrays
243 harbouring spacers targeting 2 distinct promoters using either short or long spacers (Figure 5B, C).
244 Subsequently, we quantified gene expression and genome editing and determined that only DNase

245 active SiT-Cas12a effectors combined with 20 bp spacers facilitated gene editing. In contrast, SiT-
246 Cas12a effectors combined with 15 bp spacers induced either gene repression (Figure 5B) or gene
247 activation (Figure 5C) with comparable efficiencies and without any detectable gene editing events.
248 Lastly, large CRISPR arrays containing both short and long spacers enable coordinated and highly
249 multiplexed regulation of 10 distinct genes simultaneously with gene editing of another 5 distinct
250 genes (Figure 6A, B). Taken together, SiT-Cas12a effectors, based on *AsCas12a*, facilitate
251 orthogonal transcriptional control and gene editing simply by altering spacer length.
252

253 **Discussion**

254 In this work we have described SiT-Cas12a, a platform for constitutive, conditional, inducible,
255 orthogonal and highly multiplexed genome engineering. We have demonstrated its potential in
256 numerous multiplexed genome engineering applications, including both multi-gene transcriptional
257 regulation, multi-gene knockout and coordinated multiplexed orthogonal gene transcriptional
258 control and editing. By encoding dozens of crRNAs in a single transcript, SiT-Cas12a provides a
259 powerful and highly flexible tool for highly multiplexed genome engineering thereby exceeding
260 previous published reports²⁵⁻²⁷.

261
262 In contrast to most CRISPR-Cas gene editing expression strategies, where Cas enzymes and guide
263 RNAs are expressed from distinct promoters, in our platform a single Pol II promoter expresses a
264 single transcript harbouring both Cas12a and a CRISPR array. Consequently, the ratio of Cas12a
265 and CRISPR array is fixed, which introduces distinct disadvantages and advantages. A potential
266 disadvantage is that the fixed ratio of the two components may be suboptimal, especially in
267 expression-limited conditions. Such theoretical disadvantage could be overcome by either
268 increasing SiT-Cas12a transcript abundance or encoding multiple crRNAs per gene. On the other
269 hand, the advantage of the fixed ratio is that it enables tight control of both Cas12a and crRNA
270 expression, facilitating conditional and inducible genome engineering applications. Lastly, as the
271 extent of genome editing reflects the underlying expression of the components, derived from either
272 the intrinsic variability among different cell types or the cellular environment (e.g. Cre
273 recombinase, rtTA, tTA in this work), the SiT-Cas12a platform empowers further applications in
274 DNA-writing, molecular recording and synthetic biology³⁴⁻³⁶.

275
276 Considering the mean natural length of protein coding transcripts found within mammalian cells
277 (13.5 kb), the potential for expressing multiple crRNAs in the SiT-Cas12a context is profound. In
278 the future this could theoretically be used to enable massively multiplex expression of hundreds to
279 thousands of independent crRNAs, opening up avenues for large-scale genome engineering
280 efforts^{30,37}. While SiT-Cas12a offers a seemingly unlimited potential for crRNA expression, longer
281 pre-crRNA transcripts will inevitably be challenging to synthesize and clone. Furthermore, DRs and
282 spacers containing complementary sequences that could generate complex secondary RNA
283 structures affecting the maturation of crRNAs in cells^{38,39}. Consequently, complementary regions in
284 pre-crRNA must be considered in order to improve crRNAs maturation. Future work overcoming
285 these limitations will open up numerous applications for highly multiplexed genome engineering.

286

287 In the control of cell behaviour, genetic elements act in concert. Deciphering such complexity
288 requires fine modulation of multiple genetic elements. Simultaneous encoding of dozens of crRNAs
289 in a single plasmid simplifies both guide testing and validation of gene function. Inspired by design
290 principles that embody biological efficiency, our genome engineering platform harnesses the full
291 potential of the Cas12a enzyme, providing an easy and customizable way to allow highly
292 multiplexed gene editing and transcriptional control making it possible, in the future, to
293 systematically interrogate complex genetic interactions and cellular behaviours.
294

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302

303 **Author Contributions**

304 C.C.C., N.R.W., and R.J.P. conceived of and designed the experiments. C.C.C. and N.R.W.
305 performed the experiments and analysed the data. A.J.S analysed the deep sequencing data. D.I.
306 analysed the RNA-seq data. C.C.C., N.R.W., and R.J.P. wrote the manuscript. All authors reviewed
307 the paper and provided comments.

308

309 **Competing financial interests statement**

310 The authors declare that they have no conflicts.

311

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407

408 **Figure legends**

409

410 **Figure 1:** Simultaneous control of protein and crRNA expression from Pol II promoters.

411

412 **A.** Schematic of a single transcript architecture containing both a protein coding sequence and a
413 CRISPR array. The transcript encodes for: Protein/Cas12a (grey/yellow rectangle); Triplex, a
414 tertiary RNA structural motif (small purple rectangle); direct repeat (DR, grey square); spacer (blue
415 square); and a polyadenylation sequence (Poly(A), red rectangle). After transcription and Cas12a-
416 mediated crRNA processing, the Triplex sequence stabilizes the transcript enabling concomitant
417 Protein/Cas12a expression and gene editing.

418

419 **B.** Validation of Triplex-mediated mRNA stabilization for concomitant gene editing and
420 protein expression. Combinations of crRNA expression constructs and Cas12a proteins used (I to
421 VI). Representative EGFP fluorescent images after co-transfection of different plasmid
422 combinations (I to VI). Scale bar = 1.00 mm. Quantification of gene editing efficiencies
423 (combinations I to VI). Values represent mean \pm s.e.m, n = 3 independent experiments.

424

425 **Figure 2:** Multiplexed genome editing by SiT-Cas12a.

426

427 **A.** Schematic of SiT-Cas12a, which includes the Pol II promoter EF1a, Cas12a (yellow
428 rectangle), Triplex (purple rectangle), CRISPR array (coloured and grey squares) and
429 polyadenylation sequence (Poly(A), red rectangle). Alignments and quantification of mature
430 crRNAs. n = 2 independent experiments.

431

432 **B.** Schematics of the Cas12a, SiT-Cas12a, rdSiT-Cas12a and ddSiT-Cas12a constructs used.
433 Cas12a (yellow rectangle), Triplex (purple rectangle), CRISPR array (coloured square) are
434 indicated. The Poly(A) signal is present on the constructs but not displayed. Quantification of
435 multiplexed gene editing efficiencies. Values represent mean \pm s.e.m., n = 3 independent
436 experiments.

437

438 **C.** Schematics of the plasmid combinations used. Quantification of multiplexed gene editing
439 efficiencies in cells transfected with either EF1a-Cas12a and U6-crRNA on the same plasmid
440 (Cas12a/U6), EF1a-Cas12a and EF1a-crRNA on 2 different plasmids (Cas12a+EF1a) or SiT-
441 Cas12a on a single plasmid. Values represent mean \pm s.e.m., n = 3 independent experiments.

442

443 **Figure 3:** Multiplexed transcriptional repression using SiT-Cas12a-[Repr].

444

445 **A.** Quantification of relative mRNA expression for 4 distinct genes (*EEA1*, *RAB5*, *PIK3C3*,
446 *RAB7A*) in cells co-transfected with ddCas12a fused with either 1 (KRAB), 2 (KRAB₂) or 3
447 ([Repr]) KRAB domains in combination with different crRNA array concentrations. Values
448 represent mean \pm s.e.m., n = 3 independent experiments.

449

450 **B.** Alignments of mature crRNAs in cells transfected with SiT-ddCas12a-[Repr] containing a
451 CRISPR array with 20 distinct spacers.

452

453 **C.** Quantification of relative RNA expression for 10 distinct genes after multiplexed expression
454 of 20 distinct crRNAs from SiT-ddCas12a-[Repr]. Values represent mean \pm s.e.m., n = 3
455 independent experiments.

456

457 **Figure 4:** Multiplexed transcriptional activation using SiT-Cas12a.

458

459 **A.** Quantification of relative RNA expression for 4 distinct genes (*IL1B*, *ASCL1*, *ZFP42*,
460 *IL1R2*) in cells co-transfected with ddCas12a fusion proteins (VP128, VPR, [Activ]) in combination
461 with different crRNA concentrations. Values represent mean \pm s.e.m., n = 3 independent
462 experiments.

463

464 **B.** Alignments of mature crRNAs in cells transfected with SiT-Cas12a-[Activ] containing a
465 CRISPR array with 20 distinct spacers.

466

467 **C.** Quantification of relative RNA expression for 10 distinct genes after multiplexed expression
468 of 20 distinct crRNAs from SiT-Cas12a-[Activ]. Values represent mean \pm s.e.m., n = 3 independent
469 experiments.

470 **Figure 5:** Orthogonal transcriptional gene regulation and editing.

471

472 **A.** Schematic of the SiT-Cas12a platform for orthogonal gene editing and transcriptional
473 regulation with the Pol II promoter EF1a, Cas12a (yellow rectangle), [ED] (Effector Domain, grey
474 rectangle), Triplex (purple rectangle), CRISPR array (coloured and grey squares) and
475 polyadenylation sequence (Poly(A), red rectangle) indicated. Delivery of SiT-Cas12a-[ED] coupled
476 to a CRISPR array consisting of either long (20 bp) or short (15 bp) spacer sequences enables
477 orthogonal gene editing and transcriptional gene regulation.

478

479 **B.** Quantification of relative *RAB5A* or *PIK3C3* RNA expression and gene editing efficiencies
480 in cells expressing either long (20 bp) or short (15 bp) spacers in combination with either SiT-
481 Cas12a-[Repr] or ddCas12a-[Repr]. Values represent mean \pm s.e.m., n = 3 independent experiments.

482

483 **C.** Quantification of relative RNA expression and gene editing efficiencies for 2 distinct genes
484 (*ASCL1* and *IL1B*) in cells expressing long (20 bp) or short (15 bp) spacers in combination with
485 either SiT-Cas12a-[Activ] or ddCas12a-[Activ]. Values represent mean \pm s.e.m., n = 3 independent
486 experiments.

487

488

489 **Figure 6:** Multiplexed orthogonal gene editing and transcriptional activation.

490

491 **A.** Quantification of relative RNA expression and gene editing efficiencies in cells expressing a
492 large CRISPR array (25 crRNAs) harbouring both short (15 bp) and long (20 bp) spacers using SiT-
493 Cas12a-[Repr]. Values represent mean \pm s.e.m., n = 3 independent experiments.

494

495 **B.** Quantification of relative RNA expression and gene editing efficiencies in cells expressing a
496 large CRISPR array (25 crRNAs) harbouring both short (15 bp) and long (20 bp) using SiT-Cas12a-
497 [Activ]. Values represent mean \pm s.e.m., n = 3 independent experiments.

498

499 **Methods:**

500

501 ***Mammalian cell culture.***

502 Human embryonic kidney 293T (HEK 293T) cell line (SIGMA-Aldrich) was maintained in
503 Dulbecco's modified Eagle's Medium (DMEM) (SIGMA) supplemented with 10% FBS (HyClone)
504 at 37°C with 5% CO₂.

505

506 ***Transient transfection.***

507 HEK 293T cells were transfected using Lipofectamine 2000 (Invitrogen) according to
508 manufacturer's instructions. Transient transfections were performed using either 0.6 µg or 1 µg of
509 plasmid DNA per well.

510

511 ***Plasmids.***

512 All plasmids were generated using either restriction enzyme-based cloning, Golden Gate Cloning or
513 Gibson Assembly. SiT-Cas12a constructs for multiplexed genome editing were generated by
514 replacing the U6-sgRNA from pY026 (Addgene: 84741) with the Triplex sequence and 2 DR
515 sequences, separated by 2 BsmBI restriction sites. SiT-Cas12a constructs for transcriptional control
516 were generated by replacing the U6-gRNA and Cas9-Puromycin resistance from lentiCRISPR v2
517 (Addgene: 52961) with AsCas12a, Triplex sequence and 2 DR sequences, separated by 2 SapI
518 restriction sites. Constructs for production of lentivirus were generated by replacing the U6-gRNA,
519 Cas9-Puromycin, and WPRE element from lentiCRISPR v2 (Addgene: 52961) with a CMV
520 promoter together with either *EGFP* or AsCas12a, Puromycin, Triplex sequence, 2 DR sequences,
521 separated by 2 BsmBI restriction sites and/or a WPRE element and/or a polyadenylation sequence
522 in inverted orientation. Depending on the experiment setup, either a CMV or an EF1a promoter was
523 used. The ddCas12a and rdCas12a mutants were generated by site-directed mutagenesis through the
524 substitution E993A or H800A, respectively. Transcriptional control elements were amplified from
525 gene fragments (IDT) and integrated 3' of AsCas12a using a BamHI restriction site. Assembly of
526 CRISPR arrays was performed by Golden Gate Cloning via BsmBI or SapI using either annealed
527 oligonucleotides (IDT), for single spacers or small arrays (2 to 4 spacers), or gene fragments (IDT)
528 with flanking Type IIS restriction sites for medium (10 spacers) and large (20-25 spacers) arrays.
529 The tetracycline/doxycycline inducible plasmid was constructed by exchanging the constitutive
530 promoter of SiT-Cas12a with tetracycline response element consisting of 5 repeats of bacterial TetO
531 sequence upstream of a minCMV promoter by Gibson assembly. SiT-Cas12a-[Cond] was generated

532 by adding a Lox-Stop-Lox (LSL) cassette between the CMV promoter and Cas12a by Gibson
533 assembly. DNA and spacer target sequences were listed in Supplemental File. Relevant plasmids
534 used in this study will be deposited to AddGene.

535

536 ***Large CRISPR array (10-20-25 spacers) assembly.***

537 Large CRISPR arrays containing either 10, 20 or 25 spacer sequences were assembled based on
538 previous procedures⁴⁰. Double stranded DNA fragments encoding for 4-6 crRNAs were purchased
539 from IDT Technologies (gBlock or custom gene synthesis services) and amplified by primers
540 containing type IIS restriction site (BsmBI or SapI). Type IIS recognition sites were designed to
541 allow cleavage, at different positions, on the direct repeat (DR) sequence. This procedure enables
542 both generation of non-identical 5' or 3' ends and reduction of repetitive elements (in our case DR)
543 which prohibit the chemical synthesis of double-stranded crRNAs fragments by IDT Technologies.
544 Overhangs generated by type IIS restriction enzyme digestion were intended to be complementary,
545 enabling assembly of a large CRISPR array with defined directionality. DNA fragments encoding
546 for Cas12a crRNAs and the vector backbone were assembled by standard Golden Gate Cloning. In
547 brief, simultaneous type II enzymatic digestion using either BsmBI (Thermo Scientific) or SapI
548 (Thermo Scientific) and ligation using T7 DNA ligase (NEB) in 1xT4 Ligase buffer (NEB) reaction
549 was performed. The reaction was incubated for 30 cycles (37°C for 5 min; 16°C for 5 min)
550 followed by 55°C for 10 min. Ligation reaction was transformed into chemical competent E. coli
551 Stbl3 bacteria strain to avoid potential plasmid recombination. Bacteria cells were plated on agar
552 plates supplemented with ampicillin (100 mg/l). Single colonies were cultivated overnight in liquid
553 LB supplemented with ampicillin (100 mg/l) and DNA were isolated. Correct assembly of CRISPR
554 array was verified by SANGER sequencing.

555

556 ***Inducible gene editing.***

557 HEK 293T cells were transfected using Lipofectamine 2000 (Invitrogen) according to
558 manufacturer's instructions. Transient transfections were performed using a total of 0.6 µg (0.3 µg
559 of SiT-Cas12a-[Ind] and 0.3 µg of either rtTA or tTA-Advanced (Clontech) plasmid DNA per well
560 and Doxycycline (Sigma) was added (final concentration: 0, 0.01, 0.10, 1.00 µg/ml). Quantification
561 of gene editing was performed 72 hours post-transfection.

562

563 ***Quantification of gene editing.***

564 5 x 10⁴ HEK 293T cells per well were seeded in 24-well-plates and transfected with 1 µg of DNA
565 plasmid. 72 hours post-transfection, cells were harvested and lysed in QE buffer (1 mM CaCl₂, 3
566 mM MgCl₂, 1 mM EDTA, 1% Triton X-100, 10 mM Tris pH 7.5, 0.2 mg/ml Proteinase K) using
567 the following temperature conditions: 65°C (15 min), 68°C (15 min) and 98°C (10 min). Genomic
568 DNA was used as a template for PCR-based amplification of targeted genomic regions using
569 Phusion flash polymerase (Thermo Scientific) and primers specific for each target site
570 (Supplementary material file (Indel analyses primer list)). PCR amplicons were purified using the
571 Zymo PCR purification Kit (Zymo Research) and quantified using Nanodrop 3000 (Thermo
572 Scientific).

573 In order to generate heteroduplex DNA fragments, 250 ng purified PCR amplicons were mixed with
574 10x annealing buffer (500 mM NaCl₂, 100 mM MgCl₂, 100 mM Tris-HCl) and incubated for 10
575 min at 95°C followed by ramping 95°C to 85°C with 1.34°C/s, 85°C to 75°C with 0.2°C/s, 75°C to
576 65°C with 0.2°C/s, 65°C to 55°C with 0.2°C/s, 55°C to 45°C with 0.2°C/s, 45°C to 35°C with
577 0.2°C/s and 35°C to 25°C with 0.2°C/s. Heteroduplex DNA were treated with Surveyor enhancer
578 and Surveyor nuclease from Surveyor Mutation Detection Kit (IDT) in presence of 15 mM MgCl₂
579 according to manufacturer's protocol and separated on 2% E-Gel (Thermo Scientific). Separated
580 cleavage products were imaged using Gel DOC EZ imager (Biorad) and quantified using Image Lab
581 software (Biorad). The percentage of heteroduplex DNA formation was quantified using the
582 formula: $(1-(1-(b/(a+b))*0.5))*100$ with a being equivalent to the integrated intensity of uncut DNA
583 fragments and b being equivalent to the sum of the integrated intensity of all cleavage products.

584

585 ***FACS analyses.***

586 1.2 x 10⁵ HEK 293T cells were plated in 24-well-plate and transfected with 600 ng plasmid. 48 hr
587 after transfection, cells were split and 120 hours post-transfection, cells were harvested (using PBS
588 supplemented with 0.5 mM EDTA) and fixed in 1.8% PFA (Electron Microscopy Sciences). Fixed
589 cells were stained using conjugated antibodies CD166 conjugated PE (Miltenyi Biotec 130-106-
590 575, flow cytometry dilution (FC) 1:11); CD47 conjugated FITC (Miltenyi Biotec 130-101-344, FC
591 1:11); CD97 conjugated APC-Vio770 (Miltenyi Biotec 130-105-526, FC 1:11). Cytometric analysis
592 was performed using 5-color Fortessa (BD). Data from 10000 events was collected and analysed by
593 FlowJo (BD). Cell debris was removed by SSC-A/FSC-A gating and fluorescent intensity (FITC,
594 PE or APC-Vio770) was measured. Cells transfected with empty plasmids were used as positive
595 controls. Non-transfected cells were used as a negative control. Percentage of negative cells was
596 calculated by gating on the positive control cell population. Quantification of EGFP-positive cells
597 and EGFP expression of infected cells (72 hours post infection) were performed by harvesting HEK

598 293T cells (using PBS supplemented with 0.5 mM EDTA) and fixed in 1.8% PFA (Electron
599 Microscopy Sciences). Data from 10000 events was collected and analysed by FlowJo (BD). Cell
600 debris was removed by SSC-A/FSC-A gating and fluorescent intensity (EGFP) was measured. Non-
601 infected cells were used as a negative control. Both percentage of positive cells and EGFP
602 expression (median fluorescence intensity) were calculated.

603

604 ***Quantification of mRNA expression.***

605 Gene expression analyses were conducted 48 hours after transfection according to a previously
606 established protocol³. In brief, RNA was isolated using Quick RNA Miniprep Plus kit (Zymo
607 Research), followed by reverse transcription of 500 ng RNA using Qscript cDNA supermix
608 (Quantabio). A qPCR reaction was performed using Fast Plus EvaGreen qPCR master mix
609 (Biotium) according to manufactory protocol (primers used are indicated in the Supplementary
610 material file (qPCR primer list). Quantification of RNA expression was normalized based on
611 expression of GAPDH and calculated using $\Delta\Delta Ct^{41}$. For samples with a nearly undetectable
612 amount of mRNA, Ct values exceeding 45, an arbitrary cycle number of 45 was assigned.

613

614 ***Small RNA-seq library preparation.***

615 1.2×10^7 HEK 293T cells were plated in a 150 mm dish (Thermo Scientific) and transfected with
616 30 μg of plasmid. After 48 hours, purification of small RNAs was conducted using Quick RNA
617 Miniprep Plus kit (Zymo Research) according to the manufacturer's instructions. Small RNA
618 library preparation was performed in line with a previous published protocol⁴². In brief, residual
619 genomic DNA was removed by DNase I digestion. 20 μg of small RNAs were dissolved in 39.5 μl
620 water and denatured for 5 minutes at 65°C. After cooling on ice for 5 minutes, 5 μl 10X DNase I
621 buffer including MgCl_2 (NEB), 10 units Superase-In RNase Inhibitor (Thermo Scientific) and 5
622 units DNaseI (NEB) were added and incubated for 45 minutes at 37°C. Purification of this pure
623 small RNA fraction was conducted using Quick RNA Miniprep Plus kit (Zymo Research) and
624 quantified using Nanodrop 3000 (Thermo Scientific). To capture 5' phosphorylated crRNA, an
625 additional 5' phosphorylation step was performed. RNA samples, treated with 10 units DNaseI
626 (NEB), were denatured for 2 minutes at 90°C and stored on ice for 5 minutes. Subsequently, 20
627 units of T4 PNK (Thermo Scientific) together with 10 units of Superase-In RNase Inhibitor
628 (Thermo Scientific) were added in the presence of T4 PNK buffer (Thermo Scientific) to a final
629 volume of 50 μl . After 6 hours of incubation at 37°C, an additional 10 units of T4 PNK (Thermo
630 Scientific) and 10 mM ATP (2 mM final concentration) (Thermo Scientific) were added to the
631 samples and incubated for 1 hour at 37°C. Purification of 5' phosphorylated RNA fractions were

632 performed using Quick RNA Miniprep Plus kit (Zymo Research) and quantified. As T4 PNK
633 treatment enriched transcript with 5' pyrophosphates (PP) thereby interfering with the following
634 step of small RNA library preparations, 5' PP were removed by Tobacco Acid Pyrophosphatase
635 (TAP) treatment (Epicentre). In a total volume of 20 μ l, 10 units of TAP, 2 μ l TAP buffer, 10 units
636 of Superase-In RNase Inhibitor (Thermo Scientific) and 10 μ g of PNK-treated small RNA were
637 mixed and incubated for 1 hour at 37°C and purified using Quick RNA Miniprep Plus kit (Zymo
638 Research). These enriched pre-crRNA transcripts were used as template for the preparation of small
639 RNA library. Library preparation was performed using NEBNext multiplex small rna library prep
640 set (set 1) (NEB), separated on a 6% polyacrylamide gel and purified using SpinX columns
641 (Costar). This step ensured enrichment of mature crRNAs (<60nt). 5 Million reads were sequenced
642 for each sample using Illumina NextSeq 500 (Illumina) and analysed as described below.

643

644 ***Mature crRNA quantification.***

645 Adapter sequences (described in NEBNext multiplex small rna library prep set (set 1)) were clipped
646 from sequencing reads using CutAdapt v1.10⁴³. Only reads ≥ 10 nt were retained. Reads were
647 mapped to the reference sequence of the entire crRNA array, using Bowtie v1.2.2⁴⁴, with
648 parameters "--norc -l 28 -n 2 -m 1 --best --strata". For the quantification of mature crRNAs, a BED
649 file containing the coordinates of each crRNA within the array was created and used to extract the
650 number of reads mapping onto each crRNA using the intersectBed utility from BEDTools v2.27.0
651⁴⁵. Raw counts were then normalized to the number of non-crRNA reads mapping on human
652 snoRNAs, through the following formula: $N_i = (c_i/S) \cdot 10^6$; where N_i and c_i are respectively the
653 normalized count and the raw read count for crRNA i , and S is the count of reads mapping on
654 human snoRNAs, a small RNA population largely recognized for performing housekeeping
655 function and used to normalize human microRNA (miRNA) expression^{46,47}. Finally, a fold change
656 over control condition was calculated to quantify changes across distinct experiments.

657

658 ***Deep sequencing-based CD47 editing analyses.***

659 Analyses of deep sequencing data derived from HEK 293T cells stably expressing SiT-Cas12a
660 harbouring 10 distinct crRNAs were performed using CRISPResso2 tool⁴⁸ *CD47* edited region was
661 amplified, size selected between 200 bp and 500 bp and sequenced. Analysis was performed by
662 CRISPResso2 tool⁴⁸ using the following parameters: -w 10 -cleavage offset 1 -S 20; -w 10 -
663 cleavage offset 1 -g.

664

665 ***Statistics.***

666 Unless otherwise noted, experiments in this study were performed using three independent
667 biological replicates. Tests for determination of statistical significance were not implemented.

668

669 **Data availability statement**

670 The data that support the findings of this study are available from the corresponding author upon
671 reasonable request. The datasets generated during the current study are available in the NCBI
672 Sequence Read Archive (BioProject ID PRJNA530879).

673

674 **Methods-only References:**

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695

Figure 1

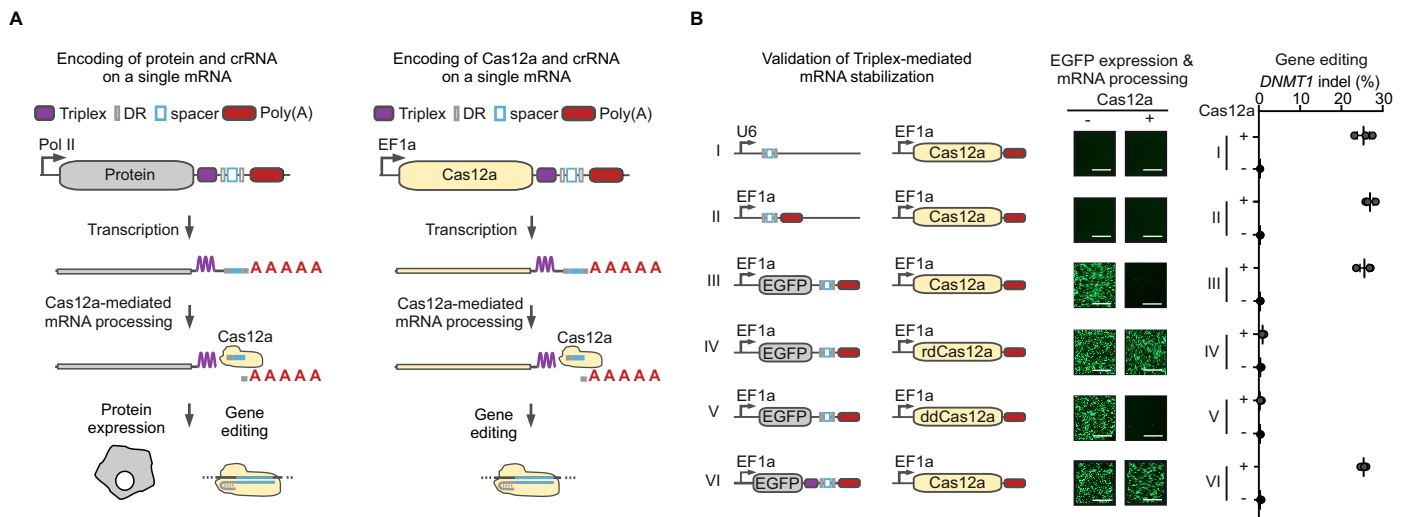


Figure 2

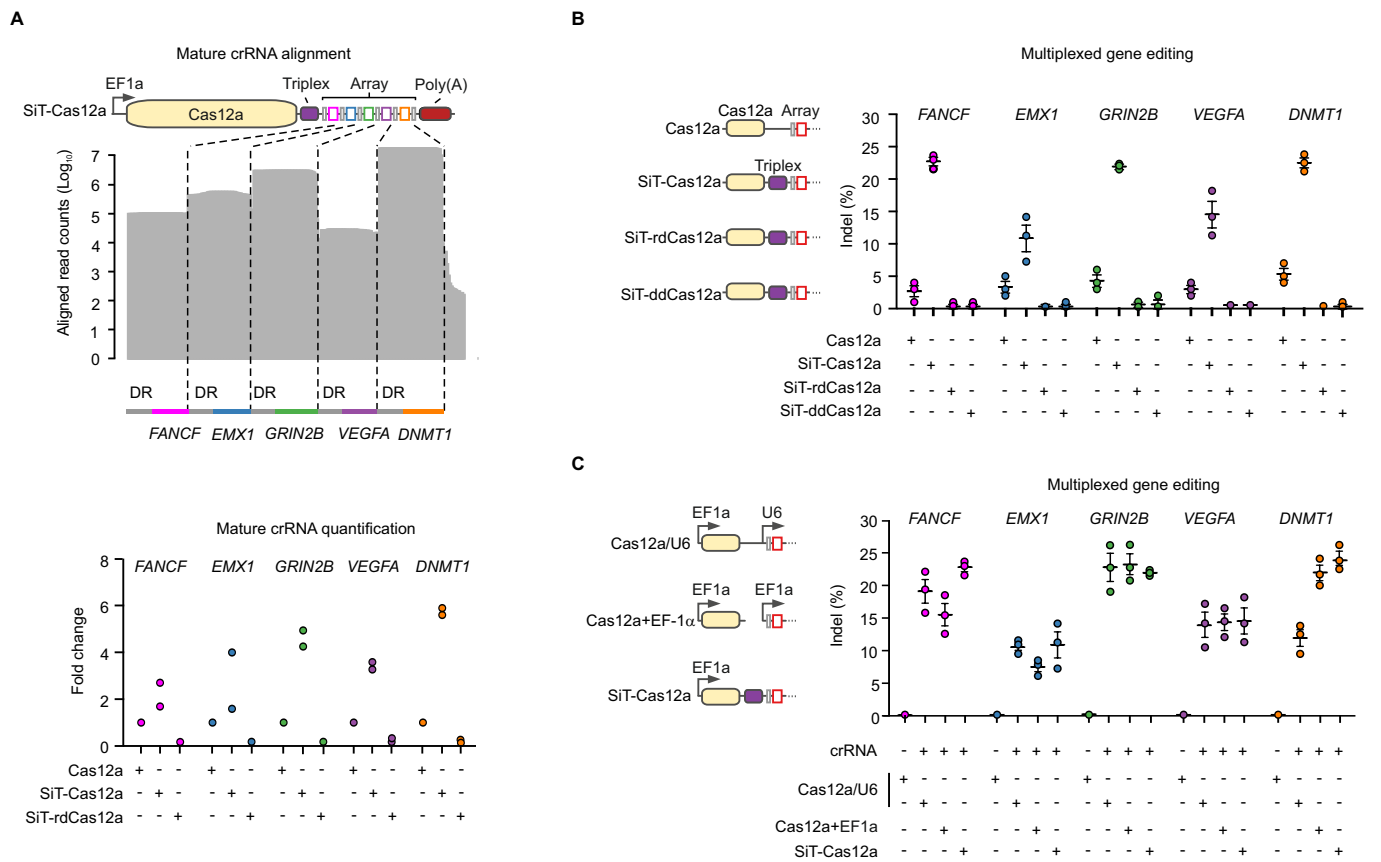


Figure 3

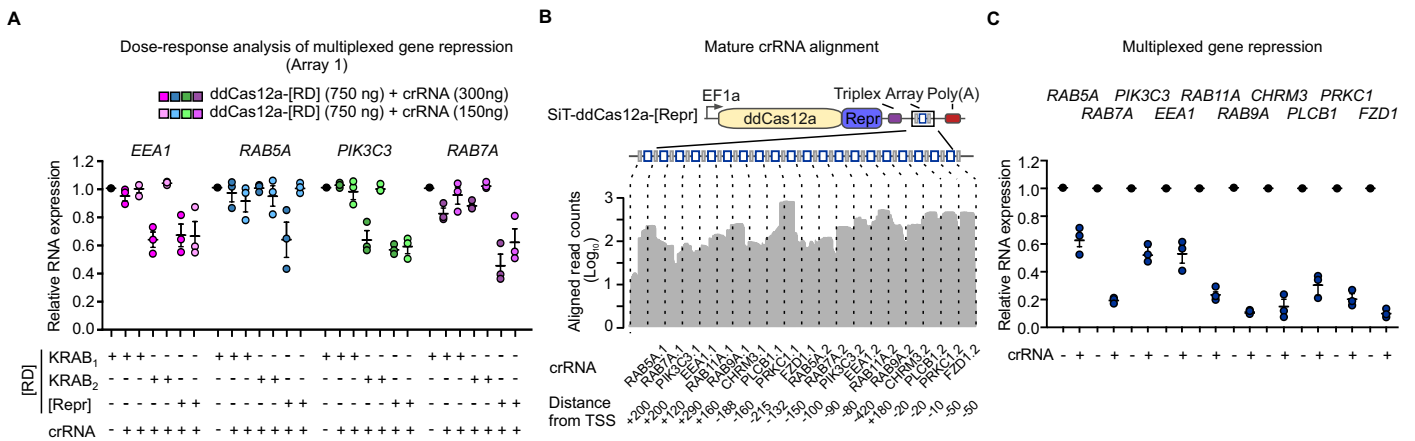


Figure 4

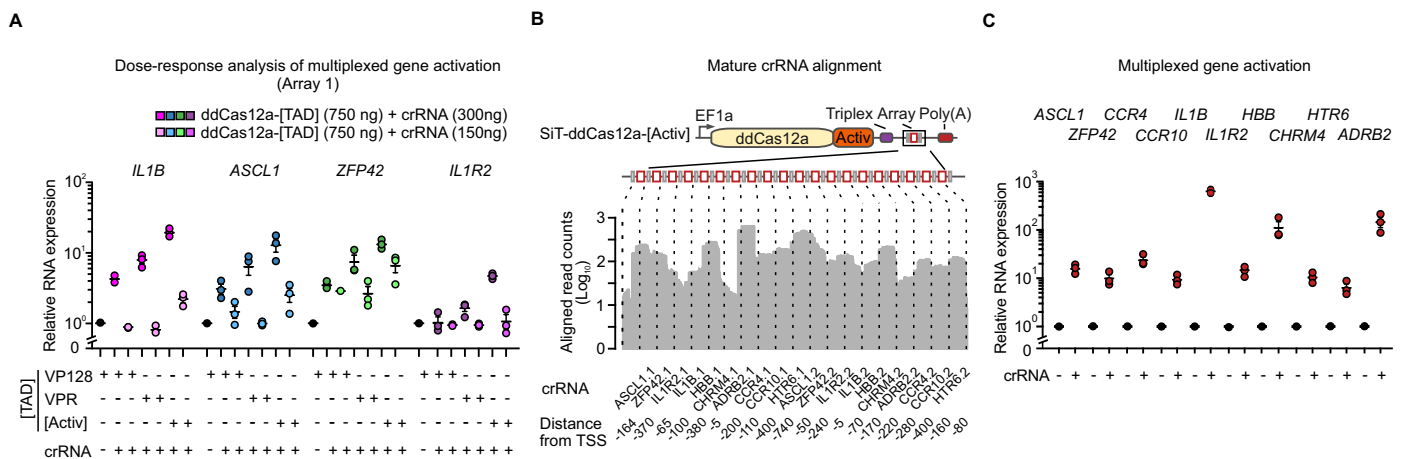


Figure 5

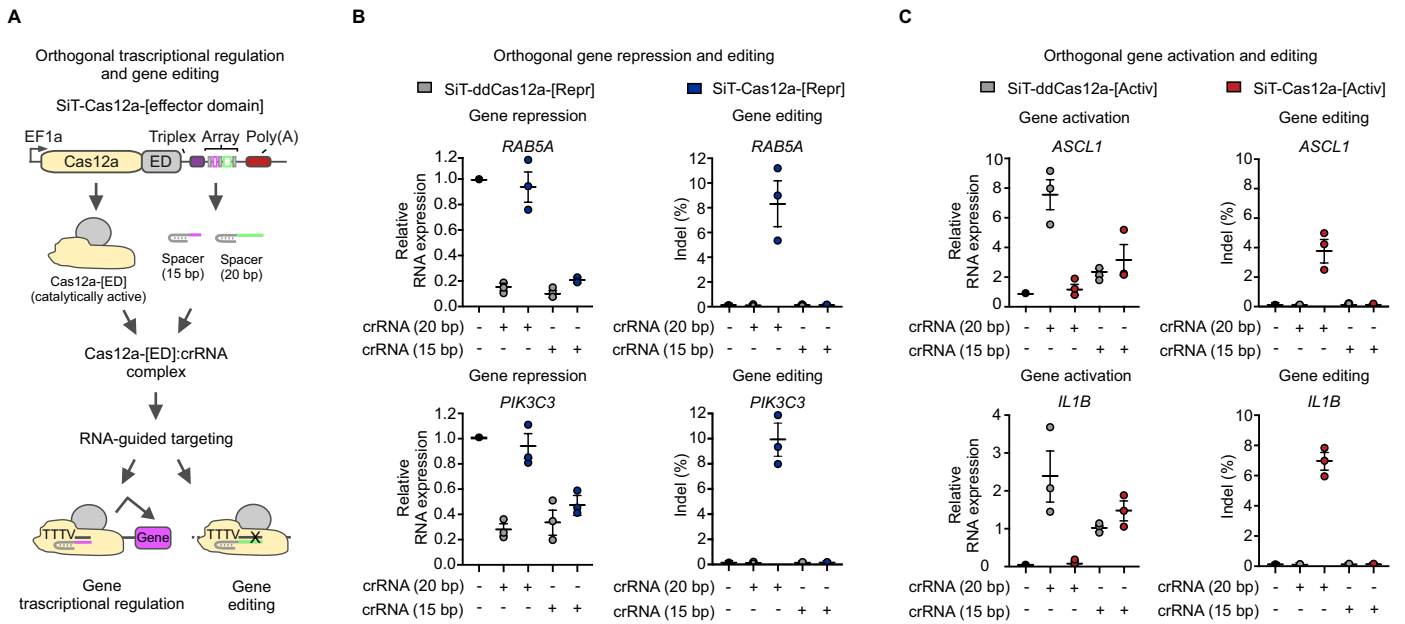
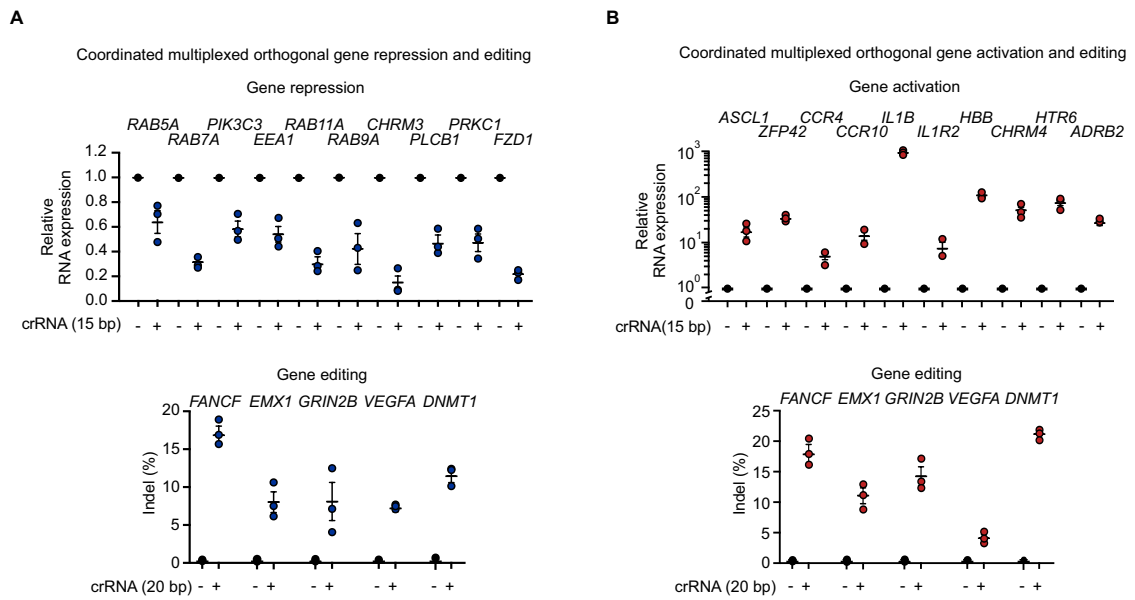
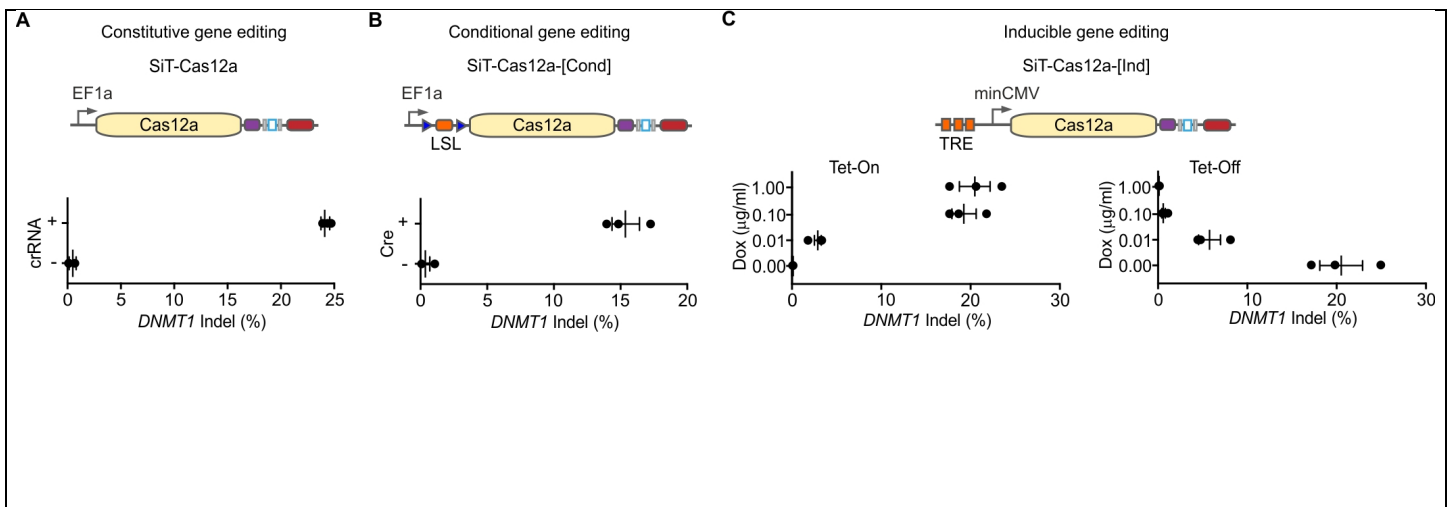


Figure 6





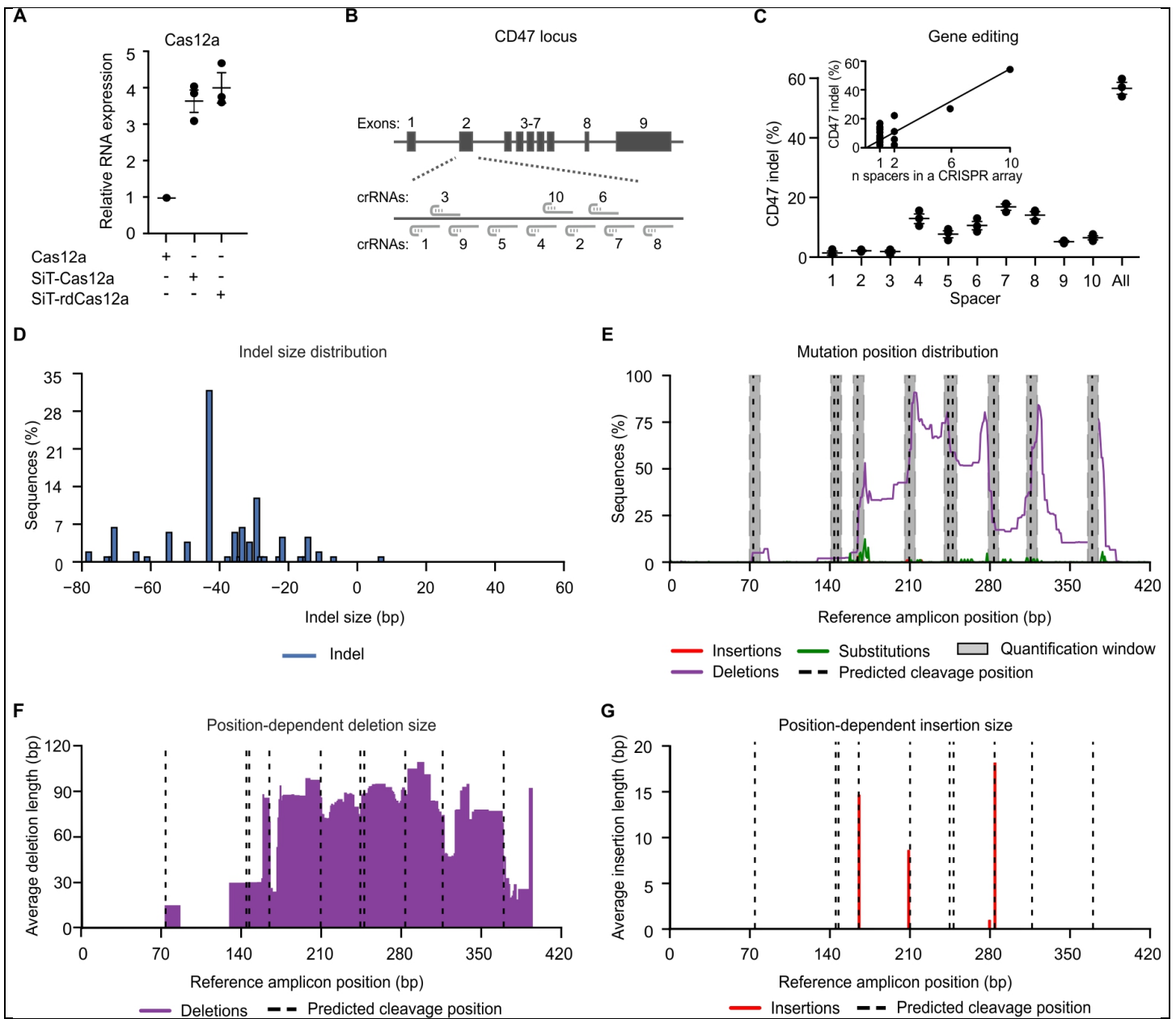
Supplementary Figure 1

Constitutive, conditional and inducible gene editing with SiT-Cas12a.

A. Quantification of gene editing efficiencies by constitutive SiT-Cas12a construct. Values represent mean \pm s.e.m., n = 3 independent experiments.

B. Quantification of gene editing efficiencies by conditional SiT-Cas12a (SiT-Cas12a-[Cond]) construct. Values represent mean \pm s.e.m., n = 3 independent experiments.

C. Quantification of gene editing efficiencies by inducible SiT-Cas12a (SiT-Cas12a-[Ind]) construct in either Tet-Off or Tet-On configuration. Values represent mean \pm s.e.m., n = 3 independent experiments.



Supplementary Figure 2

Quantification of SiT-Cas12a-induced gene mutations

A. Quantification of relative Cas12a mRNA expression in either Cas12a, SiT-Cas12a or SiT-rdCas12a expressing cells. Values represent mean \pm s.e.m., $n = 3$ independent experiments.

B. Schematic of the *CD47* locus and the target position of crRNA used.

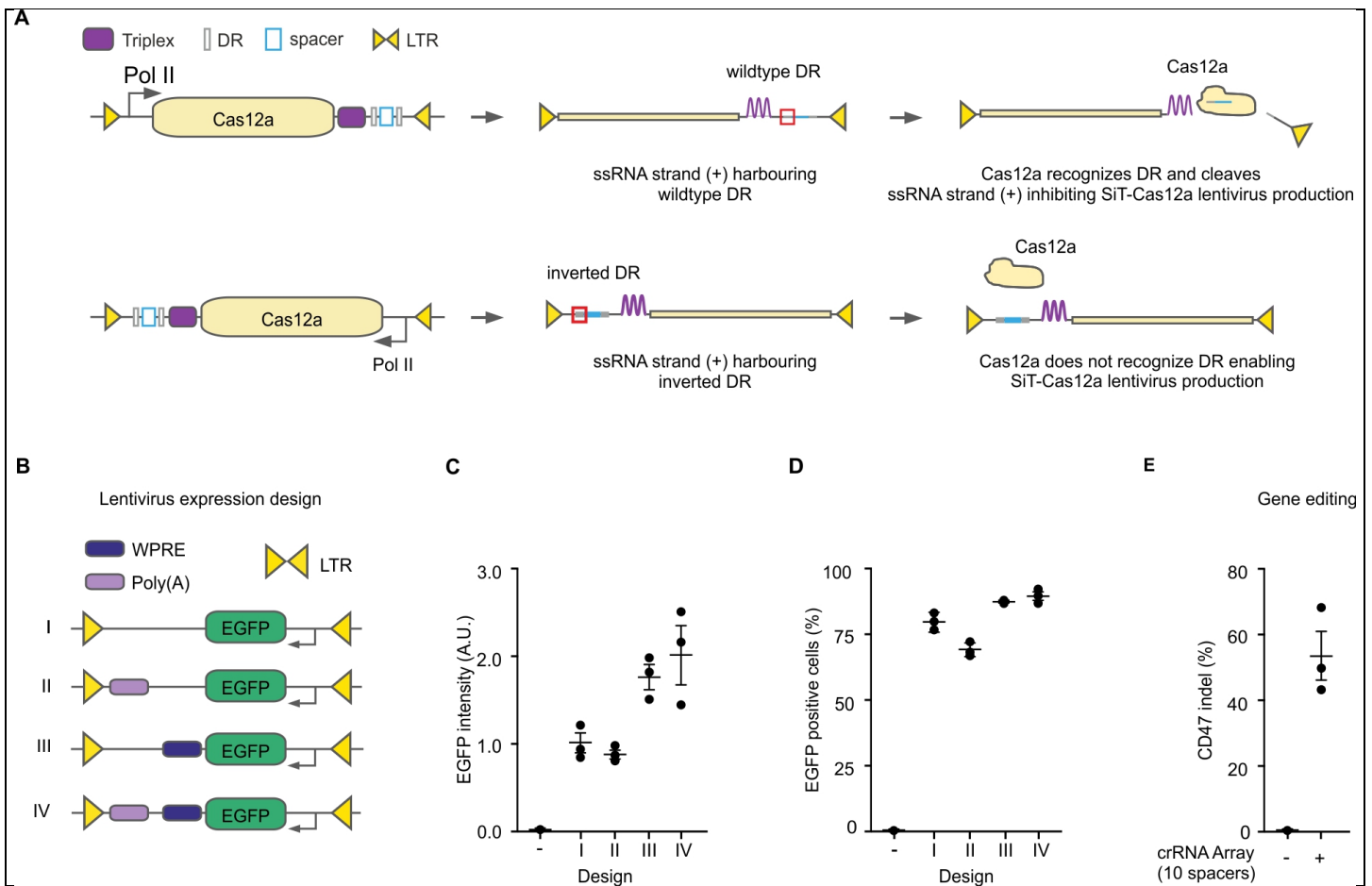
C. Quantification of SiT-Cas12a gene editing efficiencies using either single (crRNA 1-10), double (crRNAs 1-2, 3-4, 5-6, 7-8, 9-10), or multiple (comprising crRNA from 4 to 9 or from 1 to 10) spacer sequences targeting the *CD47* gene. Values represent mean \pm s.e.m., $n = 3$ independent experiments.

D. Quantification of indel size distribution generated in *CD47* locus using a CRISPR array containing 10 distinct spacer sequences. Similar results for n = 3 independent experiments.

E. Quantification of mutation position distribution generated in *CD47* locus using a CRISPR array containing 10 distinct spacer sequences. Similar results for n = 3 independent experiments.

F. Quantification of position-dependent deletion size generated in *CD47* locus using a CRISPR array containing 10 distinct spacer sequences. Similar results for n = 3 independent experiments.

G. Quantification of position-dependent insertion size generated in *CD47* locus using a CRISPR array containing 10 distinct spacer sequences. Similar results for n = 3 independent experiments.



Supplementary Figure 3

Generation of SiT-Cas12a-based lentiviral vector

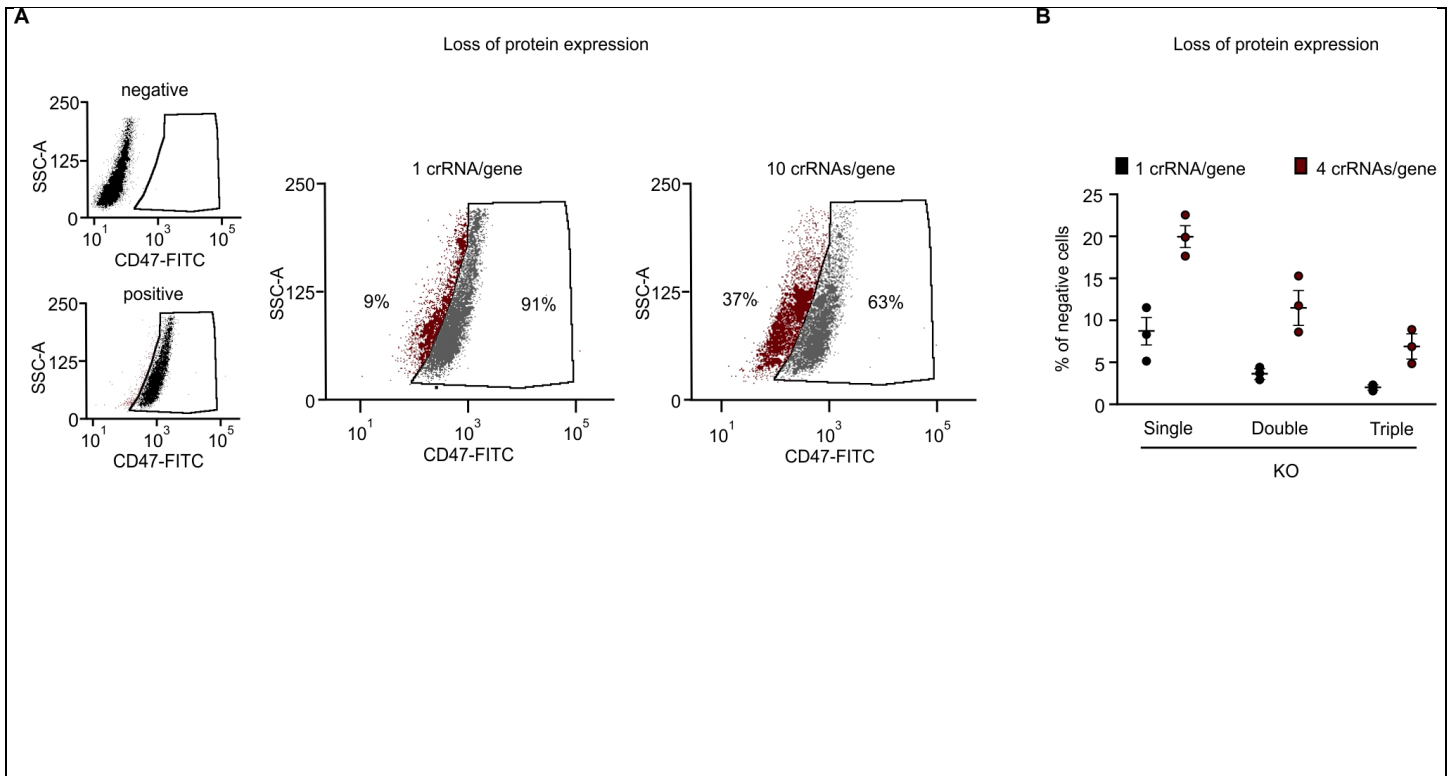
A. Schematic of a single transcript lentivirus architecture containing both Cas12a and a CRISPR array in different orientations and its effect on lentivirus production. The transcript encodes for: Cas12a (yellow rectangle); Triplex, a tertiary RNA structural motif (small purple rectangle); direct repeat (DR, grey square); spacer (blue square). The (+)-strand RNA contains a recognizable direct repeat prone to Cas12a-mediated cleavage, preventing functional assembly of virions. On the contrary, direct repeats positioned in inverted orientation are not recognized by Cas12a enabling lentivirus production.

B. Schematic of a single transcript lentivirus architecture containing both EGFP and mRNA stabilizer sequences in inverted orientation. The transcript encodes for: EGFP (green rectangle); Woodchuck Hepatitis Virus Posttranscriptional Regulatory Element (WPRE) (small blue rectangle); and a polyadenylation sequence (Poly(A), lilac rectangle).

C. Validation of single transcript lentivirus architecture in the inverted orientation. Values represent mean \pm s.e.m., $n = 3$ independent experiments.

D. Quantification of percentage of EGFP-positive cells by single transcript lentivirus architectures in the inverted orientation. Values represent mean \pm s.e.m., $n = 3$ independent experiments.

E. Quantification of gene editing efficiencies by SiT-Cas12a-based lentivirus harbouring 10 distinct spacer sequences targeting the *CD47* gene. Values represent mean \pm s.e.m., n = 3 independent experiments.

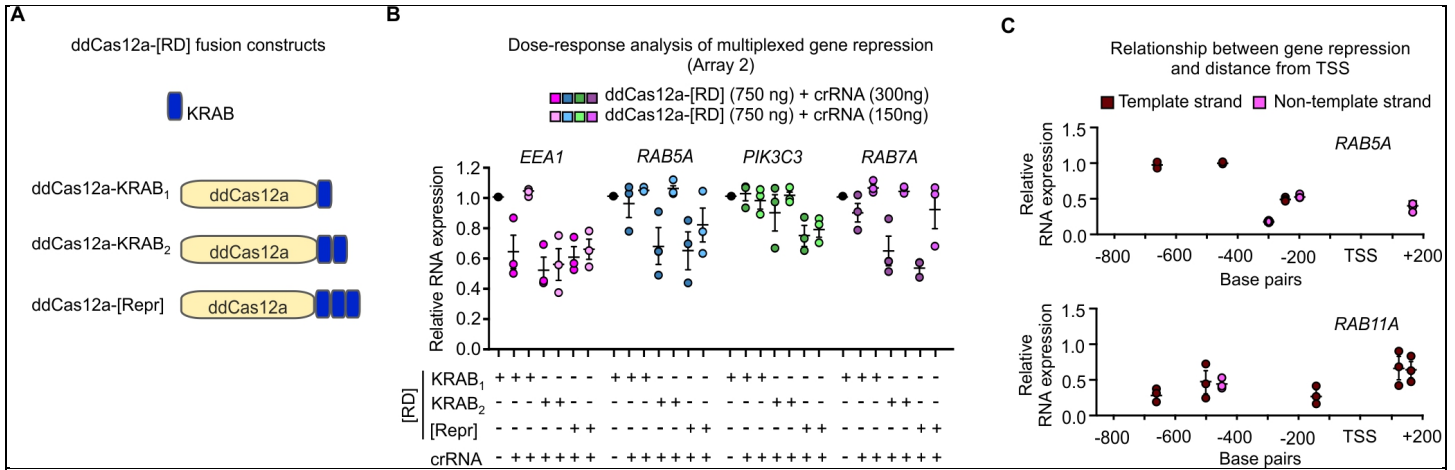


Supplementary Figure 4

Multiplexed gene editing with SiT-Cas12a.

A. Quantification of residual CD47-positive cells after SiT-Cas12a-mediated gene editing using single or multiple (10) crRNAs targeting the *CD47* gene. Similar results for n = 3 independent experiments.

B. Quantification of single, double and triple-negative cells after SiT-Cas12a-mediated gene editing using single or multiple (4) crRNAs targeting the *CD47*, *CD97*, and *CD166* genes. Values represent mean \pm s.e.m., n = 3 independent experiments.



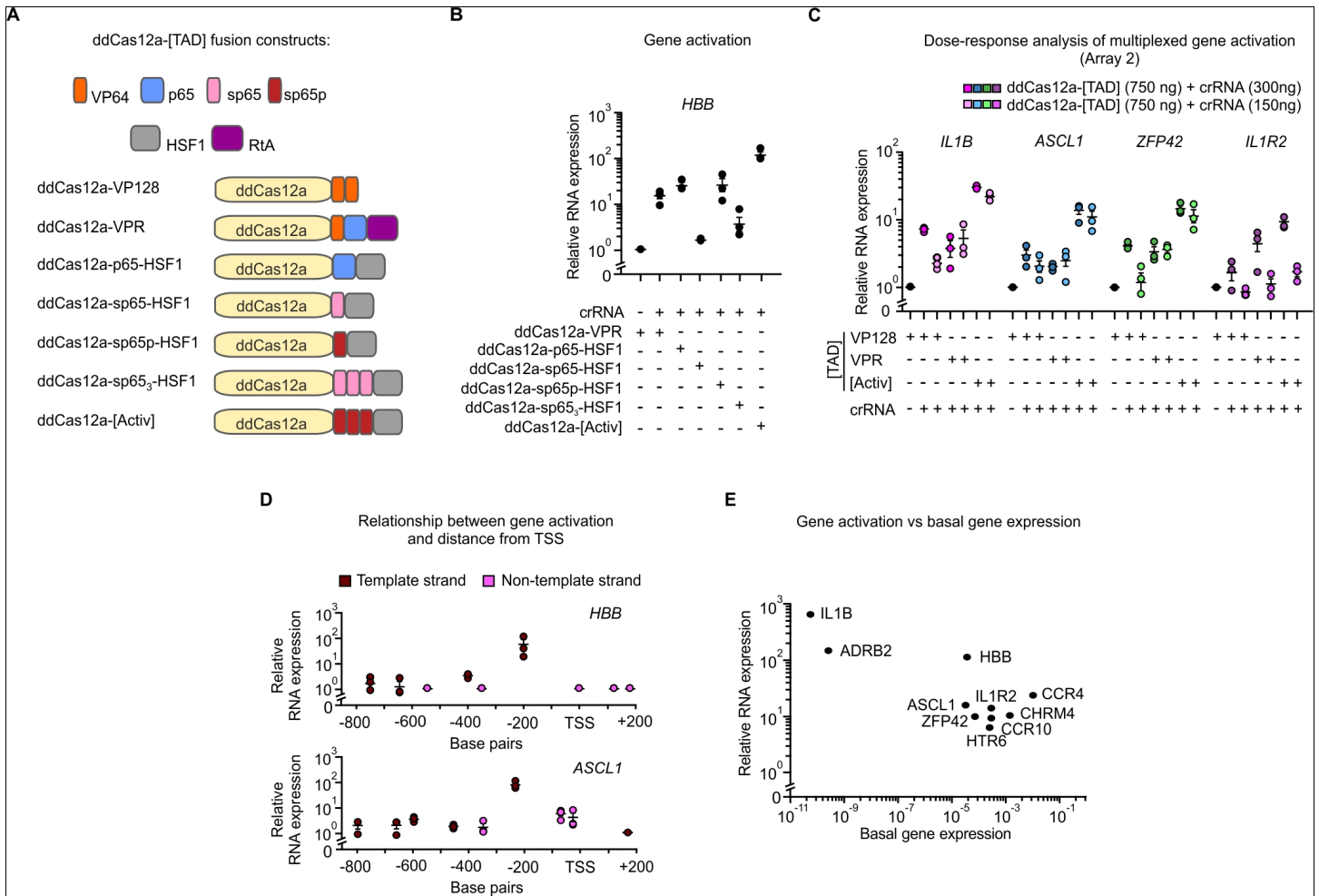
Supplementary Figure 5

Optimization and characterization of a SiT-Cas12a transcriptional repressor.

A. Schematic of Cas12a transcriptional repressors. Cas12a (yellow rectangle) and repression domain fused (RD, blue rectangle) are indicated.

B. Quantification of relative RNA expression for 4 distinct genes (*EEA1*, *RAB5*, *PIK3C3*, *RAB7A*) using ddCas12a fused with either 1 (*KRAB₁*), 2 (*KRAB₂*) or 3 (*[Repr]*) KRAB domains and in combination with different crRNA array concentrations. Values represent mean \pm s.e.m., n = 3 independent experiments.

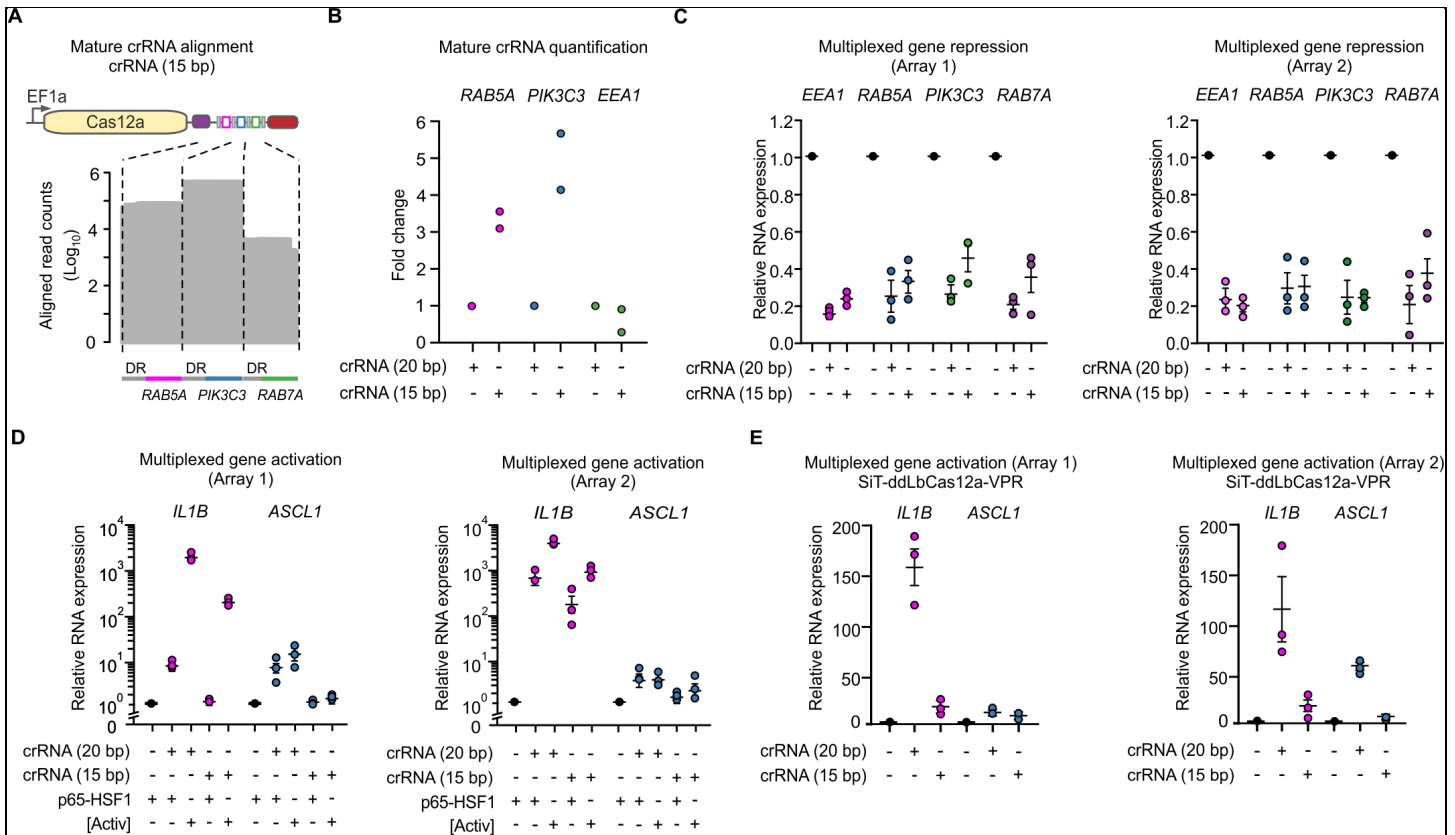
C. Quantification of relative RNA expression for 2 distinct genes (*RAB5A*, *RAB11A*) as a function of the distance between the target and the transcription start site (TSS). Values represent mean \pm s.e.m., n = 3 independent experiments.



Supplementary Figure 6

Optimization and characterization of a SiT-Cas12a transcriptional activator.

- A. Schematic of Cas12a transcriptional activators. Cas12a (yellow rectangle) and transactivation domains fused (TAD) are indicated.
- B. Quantification of relative *HBB* RNA expression using different Cas12a transcriptional activation domain fusion protein. Values represent mean \pm s.e.m., n = 3 independent experiments.
- C. Quantification of relative RNA expression for 4 distinct genes (*IL1B*, *ASCL1*, *ZFP42*, *IL1R2*) using ddCas12a fusion proteins (VP128, VPR, [Activ]) in combination with different crRNA array concentrations. Values represent mean \pm s.e.m., n = 3 independent experiments.
- D. Quantification of relative RNA expression of 2 distinct genes (*HBB*, *ASCL1*) as a function of the distance between the target and the transcription start site (TSS). Values represent mean \pm s.e.m., n = 3 independent experiments.
- E. Quantification of relative RNA expression as a function of basal gene expression for 10 distinct genes (*IL1B*, *ADRB2*, *HBB*, *IL1R2*, *CCR4*, *ASCL1*, *ZFP42*, *CHRM4*, *CCR10*, *HTR6*) induced by SiT-Cas12a-[Activ] and 20 distinct spacers targeting those 10 genes. Similar results for n = 3 independent experiments.



Supplementary Figure 7

Orthogonal genome engineering in the SiT-Cas12a context.

A. Small RNA-seq alignment showing mature short crRNAs (15 bp).

B. Quantification of mature crRNAs. n = 2 independent experiments.

C. Quantification of relative RNA expression of 4 distinct genes (*EEA1*, *RAB5*, *PIK3C3*, *RAB7A*) using SiT-ddCas12a-[Repr]. Results from 2 distinct CRISPR arrays are represented. Values represent mean ± s.e.m., n = 3 independent experiments.

D. Quantification of relative RNA expression of 2 distinct genes (*IL1B*, *ASCL1*) using either SiT-ddCas12a-p65-HSF1 or SiT-ddCas12a-[Activ]. Results from 2 distinct CRISPR arrays are represented. Values represent mean ± s.e.m., n = 3 independent experiments.

E. Quantification of relative RNA expression of 2 distinct genes (*IL1B*, *ASCL1*) using SiT-ddLbCas12a-VPR. Results from 2 distinct CRISPR arrays are represented. Values represent mean ± s.e.m., n = 3 independent experiments.

Supplementary Note 1

Single-transcript Cas12a construct:

AsCas12a-NLS-HAx3-Triplex-DR-BsmBI-BsmBI-DR

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Single-transcript Cas12a construct for Lentivirus production:

[CMV-AsCas12a-NLS-P2A-Puro-HAx3-Triplex-DR-BsmBI-BsmBI-DR-WPRE-Poly(A)] inverted

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G

Cas12a variants:

ddCas12a (E993A)

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rdCas12a (H800A)

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ddLbCas12a (D832A)

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Single-transcript Cas12a - [TAD] construct:

AsCas12a-NLS-[Aktiv]-HAX3-Triplex-DR-SapI-SapI-DR

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Single-transcript Cas12a – [Repr] construct:

AsCas12a-NLS-[Repr]-HAx3-Triplex-DR-SapI-SapI-DR

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Transcriptional control elements:

KRAB

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KRAB₂

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[Repr]

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VP64₂

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p65

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sp65

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sp65₃

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sp65p₃

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HSF1

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RtA

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[Activ]

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Direct Repeat (DR)

AATTTCTACTCTTGATAGAT

Gene editing

Gene	Target Sequence
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CD47.2	GCACTACTAAAGTCAGTGGGGAC
CD47.3	GTAATGACACTGTCGTCATTCCA
CD47.4	GAGCTCCATCAAAGGTGTAATA
CD47.5	CGTATACTTCAGTAGTGTTTTGT
CD47.6	GTAGTGCAAAAATTGAAGTCTCA
CD47.7	CTGTGTGTGAGACAGCATCACTC
CD47.8	GCTCGATGATCGTTTCACCTTCT
CD47.9	GTGCCTCCATATTAGTAACAAAG
CD47.10	ATGGAGCTCTAAACAAGTCCACT
CD97.1	AGCAGGGCTTCCTCTGGAGCTTC
CD97.2	TTGTGGTGCGCGTGTTCCAAGGC
CD97.3	CTGGCCGCCTTCTGCTGGATGAG
CD97.4	CGTGACTIONCGTCTGGAAGCTCA
CD166.1	TAGAGGATCTGAAGGCAATAAAT
CD166.2	AGGCACCTACAATAGTCAAGGTG
CD166.3	TGTGCATGCTAGTAACTGAGGAC
CD166.4	ATCACTGATCCTTGCATTACTGA
DNMT1	CTGATGGTCCATGTCTGTTACTC
FANCF	GGCGGGGTCCAGTTCGGGATTA
EMX1	TGGTTGCCACCCTAGTCATTGG
GRIN2B	GTGCTCAATGAAAGGAGATAAGG
VEGFA	CTAGGAATATTGAAGGGGGCAGG

Transcriptional activator

Gene	Target Sequence
ZFP42.1	TTGAGCGCTCACCAC
ZFP42.2	CGTGCGGGCCGGGTG
ASCL1.1	GGGAGTGGGTGGGAG
ASCL1.2	CAATGGGACACCAG
IL1B.1	CATGGTGATACATTT
IL1B.2	CTACTCCTTGCCCTT
IL1R2.1	CTTGGCCACTTCCCC
IL1R2.2	CCTATTTTTCTGTGA
ADRB2.1	GCAGTAAAGTCACAT
ADRB2.2	GTTACACTTCATGAA
CCR4.1	GCACATCTTCTTGGC
CCR4.2	ATTTTTGGGGAGATA
CCR10.1	CCCTACTCCACTTTG
CCR10.2	GTAAAATCCAGATCC
CHRM4.1	TCTCCCCTTCCTCCC
CHRM4.2	CATGTCTCCCCCAT
HBB.1	TCTACCATAATTCAG
HBB.2	AAGTCCAACCTCTAA
HTR6.1	CCAACCTCCTGGCTCC
HTR6.2	CCAGGGGCGGCTTTG

Transcriptional activator

Gene	Target Sequence
ASCL1.3 (15 bp)	GGGAGTGGGTGGGAG
ASCL1.3(20 bp)	GGGAGTGGGTGGGAGGAAGA
ASCL1.4(15 bp)	CAATGGGACACCCAG
ASCL1.4 (20 bp)	CAATGGGACACCCAGCCCCA
IL1B.3 (15 bp)	CATGGTGATACATTT
IL1B.3 (20 bp)	CATGGTGATACATTTGCAA
IL1B.4 (15 bp)	CTACTCCTTGCCCTT
IL1B.4 (20 bp)	CTACTCCTTGCCCTTCCATG
IL1R2.3 (15 bp)	CTTGGCCACTTCCCC
IL1R2.3 (20 bp)	CTTGGCCACTTCCCCATCTG
IL1R2.4 (15 bp)	CCTATTTTTCTGTGA
IL1R2.4 (20 bp)	CCTATTTTTCTGTGACTCGC
ZFP42.3 (15 bp)	TTGAGCGCTCACCAC
ZFP42.3 (20 bp)	TTGAGCGCTCACCACGTGCC
ZFP42.4 (15 bp)	CGTGCGGGCCGGGTG
ZFP42.4 (20 bp)	CGTGCGGGCCGGGTGCCTGG

Transcriptional repression

Gene	Target Sequence
CHRM3.1	AGCCAACCCACCCCA
CHRM3.2	GTCTCAAGCACGCAG
FZD1.1	TGCCTGAGTAGTGCC
FZD1.2	CGCCGGGAAGCCGGA
PLCB1.1	CCAGCCAGTTGGGAT
PLCB1.2	CACGCTGGGTCAGGC
PRKC1.1	ACAGCTCGCGTGAAA
PRKC1.2	TCTGCCCCCGTAAGG
RAB11A.1	AAGAGGTAGTCGTAC
RAB11A.2	AAGGTGAGGCCATGG
RAB9A.1	GTACTIONGCTGTCGCC
RAB9A.2	CCTGGCCCCGCCCCG
RAB7A.1	GTCTCCTCCTCGGCG
RAB7A.2	TAGCACGAGATCCAG
EEA1.1	ACAGAGGGTAAGAGA
EEA1.2	CAGCAGAAACTAGCA
PIK3C3.1	ACTACATCTATAGTT
PIK3C3.2	GGCGGGGAGTTCCGC
RAB5A.1	TCCTCCTCCGCCGCC
RAB5A.2	TTCGCGGGGCGGGGC

Transcriptional repression

Gene	Target Sequence
EEA1.3 (15 bp)	ACAGAGGGTAAGAGA
EEA1.3 (20 bp)	ACAGAGGGTAAGAGAGTGAA
EEA1.4 (15 bp)	ACCACCACCCGGCGC
EEA1.4 (20 bp)	ACCACCACCCGGCGCCCGC
PIK3C3.3 (15 bp)	ACTACATCTATAGTT
PIK3C3.3 (20 bp)	ACTACATCTATAGTTGTGAC
PIK3C3.4 (15 bp)	CAGACGGTGCGATGG
PIK3C3.4 (20 bp)	CAGACGGTGCGATGGGGGAA
RAB5A.3 (15 bp)	TCCTCCTCCGCCGCC
RAB5A.3 (20 bp)	TCCTCCTCCGCCGCCGCCGC
RAB5A.4 (15 bp)	TTCGCGGGGCGGGGC
RAB5A.4 (20 bp)	TTCGCGGGGCGGGGCGAGGC
RAB5A.5 (15 bp)	GGTTCGTGAAGGAGC
RAB5A.5 (20 bp)	GGTTCGTGAAGGAGCCGGCG
RAB7A.3 (15 bp)	GTCTCCTCCTCGGCG
RAB7A.3 (20 bp)	GTCTCCTCCTCGGCGGGAGC
RAB7A.4 (15 bp)	TGGCCAAGACTCCAG
RAB7A.4 (20 bp)	TGGCCAAGACTCCAGGCCCG

Distance from promoter TSS

Gene	Target Sequence
HBB.1	CCAAAACCTAATAAGTAACT
HBB.2	TTAGCATGCATGAGCAAATT
HBB.3	GATTA AACCTTCTGGTAAG
HBB.4	GTGCATCAACTTCTTATTTG
HBB.5	GAATCACAGCTTGGTAAGCA
HBB.6	AAGTCCA ACTCCTAAGCCAG
HBB.7	CTTCTGACACA ACTGTGTTC
HBB.8	TGATAGGCACTGACTCTCTC
HBB.9	TTGCCATGAGCCTTCACCTT
ASCL1.3	GAGCTGAATGGGACATTAGA
ASCL1.4	ACATAGTCCAGCACTTTTTT
ASCL1.5	CTCCAATTTCTAGGGTCACC
ASCL1.6	CTTCAAGTTCTTAGTAGAAT
ASCL1.7	GGAAGGGGGTGGGGGGCGTC
ASCL1.8	CAAGGAGCGGGAGAAAGGAA
ASCL1.9	AATGGGACACCCAGCCCCAC
ASCL1.10	ACTCGCCCTCCCTGGCCGGA
ASCL1.11	CTGCTGCTTCTGCTTTTTTT
RAB5A.1	TCCTCCTCCGCCGCCGCCGCCG
RAB5A.2	TTCGCGGGGCGGGGCGAGGCAGG
RAB5A.4	GTACAGTAAAGAGCGAAGGGAAA
RAB5A.5	GGCTGGGGGGTCTCTGGGCTCCT
RAB5A.6	GTTCGTGAAGGAGCCGGCGGCTG
RAB5A.7	GGGACTGACTGAGGGAGCGACGG
RAB11A.1	AAGAGGTAGTCGTA CTCTCGTC
RAB11A.2	AAGGTGAGGCCATGGGCTCTCGC
RAB11A.4	GTCACTAAGTAATTGAACA ACTA
RAB11A.5	CCCTTTGAGCCTCCTTTAGCGAC
RAB11A.6	GCGACTAAAGCTTGAAGCCCCAC
RAB11A.7	GCTCCTCGGCCGCGCAATGGGCA

qPCR Primer

Gene	sequence
ADRB2_forward	TTGCTGGCACCCAATAGAAGC
ADRB2_reverse	CAGACGCTCGAACTTGGCA
ASCL1_forward	CCCAAGCAAGTCAAGCGACA
ASCL1_reverse	AAGCCGCTGAAGTTGAGCC
CCR4_forward	TCTCGCCAAGACACTGAACAG
CCR4_reverse	GGCCCTGCATTCTCAAGAAG
CHRM3_forward	GGCCTGTGCCGATCTGATTAT
CHRM3_reverse	CGGCCTCGTGATGGAAAAG
CHRM4_forward	CAGCTCGGGCAATCAGTCC
CHRM4_reverse	GCCTATGATGAGATCAGCACAC
EEA1_forward	AGCAACTCCTATAAACACAGTGG
EEA1_reverse	AGCAAGATTAGACTCTCCTCCAT
FZD1_forward	AGCCATCCAGTTGCACGAG
FZD1_reverse	GAGTCGGGCCACTTGAAGTT
HBB_forward	AGGAGAAGTCTGCCGTTACTG
HBB_reverse	CCGAGCACTTTCTTGCCATGA
HTR6_forward	GCAACACGTCCAATTCTTCC
HTR6_reverse	TGCAGCACATCACGTCGAA
IL1B_forward	AAACAGATGAAGTGCTCCTTCC
IL1B_reverse	AAGATGAAGGGAAAGAAGGTGC
IL1R2_forward	ATGTTGCGCTTGTACGTGTTG
IL1R2_reverse	CCCGCTTGTAATGCCTCCC
PIK3C3_forward	CCTGGAAGACCCAATGTTGAAG
PIK3C3_reverse	CGGGACCATACACATCCCAT
PLCB1_forward	GGAAGCGGCAAAAAGAAGCTC
PLCB1_reverse	CGTCGTCTCACTTTCCGT
PRKC1_forward	GACAACGAACAGCTCTTCACC
PRKC1_reverse	CCAGGACGTTCTGGTACACA
RAB5A_forward	AGACCCAACGGGCCAAATAC
RAB5A_reverse	GCCCCAATGGTACTCTCTTGAA
RAB7A_forward	GTGTTGCTGAAGGTTATCATCCT
RAB7A_reverse	GCTCCTATTGTGGCTTTGTACTG
RAB9A_forward	AGGGACAACGGCGACTATC
RAB9A_reverse	TCTGACCTATCCTCGGTAGCA
RAB11A_forward	CAACAAGAAGCATCCAGGTTGA
RAB11A_reverse	GCACCTACAGCTCCACGATAAT
ZFP42_forward	AGAAACGGGCAAAGACAAGAC
ZFP42_reverse	GCTGACAGGTTCTATTTCCGC

NGS primer

CD47_forward	ACACTCTTTCCTACACGACGCTCTTCCGATCT NNAAGATCTTACAGTACAGACTTC
CD47_reverse	GTGACTGGAGTTCAGACGTGTGCTCTTCCGATC NNCAGGACAAATAAAAAAAGAAGC

Indel analysis primer

ASCL1_forward	CGAGGAACCCGAAGAGAATAACAGTGAG
ASCL1_reverse	CCGCTGGCGCCTTCTTGTTTCTA
CD47_forward	TATCAGTTCAGCAAGTTCTATTTAGCAGTGTGTT
CD47_reverse	AGTACATTACCCAGGACCAATCAGCCAAA
DNMT1_forward	CTGGGACTCAGGCGGGTCAC
DNMT1_reverse	CCTCACACAACAGCTTCATGTCAGC
EMX1_forward	TGGATGCCCGTGTCATTAAGAGAGAGACTTT
EMX1_reverse	CCCTTCTGTGAATGTTAGACCCATGGGAGC
FANCF_forward	GTATTAGGGCTTTTAAGTTGCCAGAGTCAAGGA
FANCF_reverse	GTCTGTTAGCAGACCCAGATAGACAGGAGAC
GRIN2B_forward	GAATGCAGGGCTTGTGTACTTATAGCCCC
GRIN2B_reverse	ACAAATGCATGGTTTAGTCCTCAGCACAAAC
IL1B_forward	CCATGAGATTGGCTAGGGTAACAGCA
IL1B_reverse	AGAGACAGAGAGACTCCCTTAGCACC
PIK3C3_forward	GTGGGCGCCTTGTGCACATGC
PIK3C3_reverse	CACCTCCCGTGCTAATACACCATGTGCTC
RAB5A_forward	CGCCCCGGAACAAACCTAGGC
RAB5A_reverse	CCAGGACGGAGACCAGGCGGAACC
VEGFA_forward	AAACTCCCCCACCCTTTCC
VEGFA_reverse	ATTCCAGCACCGAGCGCCC

Supplementary Note 1

Single-transcript Cas12a construct:

AsCas12a-NLS-HAx3-Triplex-DR-BsmBI-BsmBI-DR

ATGACACAGTTCGAGGGCTTTACCAACCTGTATCAGGTGAGCAAGACACTGCGGTTTGAGCT
GATCCCACAGGGCAAGACCCTGAAGCACATCCAGGAGCAGGGCTTCATCGAGGAGGACAAGG
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ACCAGTGCCTGCAGCTGGTGCAGCTGGATTGGGAGAACCTGAGCGCCGCCATCGACTCCTATA
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LSL-AsCas12a-NLS-HAx3-Triplex-DR-BsmBI-BsmBI-DR

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CTCACCAGGCAGTTCCATAGGATGGCAAGATCCTGGTATTGGTCTGCCAAATTTCTACTCT
TGTAGATGGAGACGCCAAATTCATCCAGCTTCAACAGCTATTTCAAGCTTTCATGATTT
CTCGTCTCTAATTTCTACTCTTGATAGAT

TRE-miniCMV-AsCas12a-NLS-HAx3-Triplex-DR-BsmBI-BsmBI-DR

TCGAGTTTACTCCCTATCAGTGATAGAGAACGTATGTTCGAGTTTACTCCCTATCAGTGATAG
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Single-transcript Cas12a construct for Lentivirus production:

[CMV-AsCas12a-NLS-P2A-Puro-HAx3-Triplex-DR-BsmBI-BsmBI-DR-WPRE-Poly(A)] inverted

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Cas12a variants:

ddCas12a (E993A)

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rdCas12a (H800A)

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ddLbCas12a (D832A)

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Single-transcript Cas12a – [TAD] construct:

AsCas12a-NLS-[Aktiv]-HAx3-Triplex-DR-SapI-SapI-DR

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Single-transcript Cas12a - [Repr] construct:

AsCas12a-NLS-[Repr]-HAX3-Triplex-DR-SapI-SapI-DR

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Transcriptional control elements:

KRAB

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KRAB₂

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[Repr]

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VP64₂

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TGACATGCTTGGTTCGGATGCCCTTGATGACTTTGACCTCGACATGCTCGGCAGTGACGCCCT
TGATGATTTGACCTGGACATGCTGGGATCCGGTGGGGACGCATTGGACGATTTTGATCTGG
ATATGCTGGGAAGTGACGCCCTCGATGATTTTGACCTTGACATGCTTGGTTCGGATGCCCTTG
ATGACTTTGACCTCGACATGCTCGGCAGTGACGCCCTTGATGATTTGACCTGGACATGCTG

p65

CCTTCAGGGCAGATCAGCAACCAGGCCCTGGCTCTGGCCCCTAGCTCCGCTCCAGTGCTGGCC
CAGACTATGGTGCCCTCTAGTGCTATGGTGCTCTGGCCCAGCCACCTGCTCCAGCCCCTGTG
CTGACCCAGGACCACCCAGTCACTGAGCGCTCCAGTGCCCAAGTCTACACAGGCCGGCGAG
GGGACTCTGAGTGAAGCTCTGCTGCACCTGCAGTTCGACGCTGATGAGGACCTGGGAGCTCT
GCTGGGGAACAGCACCGATCCCGGAGTGTTACAGATCTGGCCTCCGTGGACAACCTCTGAGTT
TCAGCAGCTGCTGAATCAGGGCGTGTCCATGTCTCATAGTACAGCCGAACCAATGCTGATGG
AGTACCCCGAAGCCATTACCCGGCTGGTGACCGGCAGCCAGCGGCCCCCGACCCCGCTCCAA
CTCCCCTGGGAACCAGCGGCCTGCCTAATGGGCTGTCCGGAGATGAAGACTTCTCAAGCATCG
CTGATATGGACTTTAGTGCCCTGCTGTACAGATTTCTCTAGTGGGCAG

sp65

AGCGGCCTGCCTAATGGGCTGTCCGGAGATGAAGACTTCTCAAGCATCGCTGATATGGACTT
TAGTGCCCTGCTGTACAGATTTCTCT

sp65₃

AGCGGGTTGCCTAATGGGTTGAGTGGTGACGAGGATTTTAGCAGTATCGCCGATATGGACTT
TTCAGCTCTCCTCAGCCAGATAAGCTCTGGAAGTAGTGGGCTTCCAAATGGGCTTAGCGGTG
ACGAAGATTTCTCTAGTATAGCGGACATGGACTTCTCAGCTCTGCTCTCCAGATTTCTTCAG
GGTCCAGCGGCTTGCCCAACGGCCTCAGTGGCGACGAGGATTTTTCTAGCATAGCAGACATG
GACTTTTCCGCACTCCTCAGTCAGATTTCTTCT

sp65p3

AGCGGGTTGCCTAATGGGTTGGATGGTGACGAGGATTTTAGCGATATCGCCGATATGGACTT
TTCAGCTCTCCTCAGCCAGATAAGCTCTGGAAGTAGTGGGCTTCCAAATGGGCTTGACGGTG
ACGAAGATTTCTCTGATATAGCGGACATGGACTTCTCAGCTCTGCTCTCCAGATTTCTTCAG
GGTCCAGCGGCTTGCCCAACGGCCTCGATGGCGACGAGGATTTTTCTGACATAGCAGACATG
GACTTTTCCGCACTCCTCAGTCAGATTTCTTCT

HSF1

GGCTTCAGCGTGGACACCAGTGCCCTGCTGGACCTGTTTCAGCCCCTCGGTGACCGTGCCCGAC
ATGAGCCTGCCTGACCTTGACAGCAGCCTGGCCAGTATCCAAGAGCTCCTGTCTCCCCAGGAG
CCCCCAGGCCTCCCGAGGCAGAGAACAGCAGCCCGGATTCAGGGAAGCAGCTGGTGCCTAC
ACAGCGCAGCCGCTGTTCTGCTGGACCCCGGCTCCGTGGACACCGGGAGCAACGACCTGCCG
GTGCTGTTTGAGCTGGGAGAGGGCTCCTACTTCTCCGAAGGGGACGGCTTCGCCGAGGACCCC
ACCATCTCCCTGCTGACAGGCTCGGAGCCTCCCAAAGCCAAGGACCCCACTGTCTCC

RtA

CGGGATTCCAGGGAAGGGATGTTTTTGCCGAAGCCTGAGGCCGGCTCCGCTATTAGTGACGT
GTTTGAGGGCCGCGAGGTGTGCCAGCCAAAACGAATCCGGCCATTTTCATCCTCCAGGAAGTCC
ATGGGCCAACCGCCCACTCCCCGCCAGCCTCGCACCAACACCAACCGGTCCAGTACATGAGCC
AGTCGGGTCACTGACCCCGGCACCAGTCCCTCAGCCACTGGATCCAGCGCCCGCAGTGACTCC
CGAGGCCAGTCACCTGTTGGAGGATCCCGATGAAGAGACGAGCCAGGCTGTCAAAGCCCTTC
GGGAGATGGCCGATACTGTGATTCCCCAGAAGGAAGAGGCTGCAATCTGTGGCCAAATGGAC
CTTTCCCATCCGCCCCAAGGGGCCATCTGGATGAGCTGACAACCACACTTGAGTCCATGACC
GAGGATCTGAACCTGGACTCACCCCTGACCCCGGAATTGAACGAGATTCTGGATACCTTCTCTG
AACGACGAGTGCCTCTTGATGCCATGCATATCAGCACAGGACTGTCCATCTTCGACACATCT
CTGT

[Activ]

GGTGGGGTGGGTCCAGCGGGTTGCCTAATGGGTTGGATGGTGACGAGGATTTTAGCGATAT
CGCCGATATGGACTTTTCAGCTCTCCTCAGCCAGATAAGCTCTGGAAGTAGTGGGCTTCCAAA
TGGGCTTGACGGTGACGAAGATTTCTCTGATATAGCGGACATGGACTTCTCAGCTCTGCTCTC
CCAGATTTCTTCAGGGTCCAGCGGCTTGCCCAACGGCCTCGATGGCGACGAGGATTTTTCTGA
CATAGCAGACATGGACTTTTCCGCACTCCTCAGTCAGATTTCTTCTGGAGGCGGTGGAAGCGG
CTTCAGCGTGGACACCAGTGCCCTGCTGGACCTGTTTCAGCCCCTCGGTGACCGTGCCCGACAT
GAGCCTGCCTGACCTTGACAGCAGCCTGGCCAGTATCCAAGAGCTCCTGTCTCCCCAGGAGCC
CCCCAGGCCTCCCGAGGCAGAGAACAGCAGCCCGGATTCAGGGAAGCAGCTGGTGCCTACAC
AGCGCAGCCGCTGTTCTGCTGGACCCCGGCTCCGTGGACACCGGGAGCAACGACCTGCCGGT
GCTGTTTGAGCTGGGAGAGGGCTCCTACTTCTCCGAAGGGGACGGCTTCGCCGAGGACCCAC
CATCTCCCTGCTGACAGGCTCGGAGCCTCCCAAAGCCAAGGACCCCACTGTCTCC

Direct Repeat (DR)

AATTTCTACTCTTAGAT

Gene editing

Gene	Target Sequence
CD47.1	ATTAAATAGTAGCTGAGCTGATC
CD47.2	GCACTACTAAAGTCAGTGGGGAC
CD47.3	GTAATGACACTGTCGTCATTCCA
CD47.4	GAGCTCCATCAAAGGTGTAAATA
CD47.5	CGTATACTTCAGTAGTGTTTTGT
CD47.6	GTAGTGCAAAAATTGAAGTCTCA
CD47.7	CTGTGTGTGAGACAGCATCACTC
CD47.8	GCTCGATGATCGTTTCACCTTCT
CD47.9	GTGCCTCCATATTAGTAACAAAG
CD47.10	ATGGAGCTCTAAACAAGTCCACT
CD97.1	AGCAGGGCTTCCTCTGGAGCTTC
CD97.2	TTGTGGTGCGCGTGTTCCAAGGC
CD97.3	CTGGCCGCCTTCTGCTGGATGAG
CD97.4	CGTGACTACCGTCTGGAAGCTCA
CD166.1	TAGAGGATCTGAAGGCAATAAAT
CD166.2	AGGCACCTACAATAGTCAAGGTG
CD166.3	TGTGCATGCTAGTAACTGAGGAC
CD166.4	ATCACTGATCCTTGCATTACTGA
DNMT1	CTGATGGTCCATGTCTGTTACTC
FANCF	GGCGGGGTCCAGTTCGGGGATTA
EMX1	TGGTTGCCACCCTAGTCATTGG
GRIN2B	GTGCTCAATGAAAGGAGATAAGG
VEGFA	CTAGGAATATTGAAGGGGGCAGG

Transcriptional activator

Gene	Target Sequence
ZFP42.1	TTGAGCGCTCACCAC
ZFP42.2	CGTGCGGGCCGGGTG
ASCL1.1	GGGAGTGGGTGGGAG
ASCL1.2	CAATGGGACACCCAG
IL1B.1	CATGGTGATACATTT
IL1B.2	CTACTCCTTGCCCTT
IL1R2.1	CTTGGCCACTTCCCC
IL1R2.2	CCTATTTTTTCTGTGA
ADRB2.1	GCAGTAAAGTCACAT
ADRB2.2	GTTACACTTCATGAA
CCR4.1	GCACATCTTCTTGGC
CCR4.2	ATTTTTGGGGAGATA
CCR10.1	CCCTACTCCACTTTG
CCR10.2	GTAAAATCCAGATCC
CHRM4.1	TCTCCCCTTCCTCCC
CHRM4.2	CATGTCTCCCCCAT
HBB.1	TCTACCATAATTCAG
HBB.2	AAGTCCAACCTCCTAA
HTR6.1	CCAACCTCCTGGCTCC
HTR6.2	CCAGGGGCGGCTTTG

Transcriptional activator

Gene	Target Sequence
ASCL1.3 (15 bp)	GGGAGTGGGTGGGAG
ASCL1.3(20 bp)	GGGAGTGGGTGGGAGGAAGA
ASCL1.4(15 bp)	CAATGGGACACCCAG
ASCL1.4 (20 bp)	CAATGGGACACCCAGCCCCA
IL1B.3 (15 bp)	CATGGTGATACATTT
IL1B.3 (20 bp)	CATGGTGATACATTTGCAAA
IL1B.4 (15 bp)	CTACTCCTTGCCCTT
IL1B.4 (20 bp)	CTACTCCTTGCCCTTCCATG
IL1R2.3 (15 bp)	CTTGGCCACTTCCCC
IL1R2.3 (20 bp)	CTTGGCCACTTCCCCATCTG
IL1R2.4 (15 bp)	CCTATTTTTCTGTGA
IL1R2.4 (20 bp)	CCTATTTTTCTGTGACTCGC
ZFP42.3 (15 bp)	TTGAGCGCTCACCAC
ZFP42.3 (20 bp)	TTGAGCGCTCACCACGTGCC
ZFP42.4 (15 bp)	CGTGCGGGCCGGGTG
ZFP42.4 (20 bp)	CGTGCGGGCCGGGTGCCTGG

Transcriptional repression

Gene	Target Sequence
CHRM3.1	AGCCAACCCACCCCA
CHRM3.2	GTCTCAAGCACGCAG
FZD1.1	TGCCTGAGTAGTGCC
FZD1.2	CGCCGGGAAGCCGGA
PLCB1.1	CCAGCCAGTTGGGAT
PLCB1.2	CACGCTGGGTCAGGC
PRKC1.1	ACAGCTCGCGTGAAA
PRKC1.2	TCTGCCCCCGTAAGG
RAB11A.1	AAGAGGTAGTCGTAC
RAB11A.2	AAGGTGAGGCCATGG
RAB9A.1	GTACTCGCTGTCGCC
RAB9A.2	CCTGGCCCCGCCCCG
RAB7A.1	GTCTCCTCCTCGGCG
RAB7A.2	TAGCACGAGATCCAG
EEA1.1	ACAGAGGGTAAGAGA
EEA1.2	CAGCAGAACTAGCA
PIK3C3.1	ACTACATCTATAGTT
PIK3C3.2	GGCGGGGAGTTCCGC
RAB5A.1	TCCTCCTCCGCCGCC
RAB5A.2	TTCGCGGGGCGGGGC

Transcriptional repression

Gene	Target Sequence
EEA1.3 (15 bp)	ACAGAGGGTAAGAGA
EEA1.3 (20 bp)	ACAGAGGGTAAGAGAGTGAA
EEA1.4 (15 bp)	ACCACCACCCGGCGC
EEA1.4 (20 bp)	ACCACCACCCGGCGCCGCG
PIK3C3.3 (15 bp)	ACTACATCTATAGTT
PIK3C3.3 (20 bp)	ACTACATCTATAGTTGTGAC
PIK3C3.4 (15 bp)	CAGACGGTGCGATGG
PIK3C3.4 (20 bp)	CAGACGGTGCGATGGGGGAA
RAB5A.3 (15 bp)	TCCTCCTCCGCCGCC
RAB5A.3 (20 bp)	TCCTCCTCCGCCGCCGCCG
RAB5A.4 (15 bp)	TTCGCGGGGCGGGGC
RAB5A.4 (20 bp)	TTCGCGGGGCGGGGCGAGGC
RAB5A.5 (15 bp)	GGTTCGTGAAGGAGC
RAB5A.5 (20 bp)	GGTTCGTGAAGGAGCCGGCG
RAB7A.3 (15 bp)	GTCTCCTCCTCGGCG
RAB7A.3 (20 bp)	GTCTCCTCCTCGGCGGGAGC
RAB7A.4 (15 bp)	TGGCCAAGACTCCAG
RAB7A.4 (20 bp)	TGGCCAAGACTCCAGGCCCG

Distance from promoter TSS

Gene	Target Sequence
HBB.1	CCAAAACCTAATAAGTAACT
HBB.2	TTAGCATGCATGAGCAAATT
HBB.3	GATTAAAACCTTCTGGTAAG
HBB.4	GTGCATCAACTTCTTATTTG
HBB.5	GAATCACAGCTTGGTAAGCA
HBB.6	AAGTCCAACCTCCTAAGCCAG
HBB.7	CTTCTGACACAACTGTGTTC
HBB.8	TGATAGGCACTGACTCTCTC
HBB.9	TTGCCATGAGCCTTCACCTT
ASCL1.3	GAGCTGAATGGGACATTAGA
ASCL1.4	ACATAGTCCAGCACTTTTTT
ASCL1.5	CTCCAATTTCTAGGGTCACC
ASCL1.6	CTTCAAGTTCTTAGTAGAAT
ASCL1.7	GGAAGGGGGTGGGGGGCGTC
ASCL1.8	CAAGGAGCGGGAGAAAGGAA
ASCL1.9	AATGGGACACCCAGCCCCAC
ASCL1.10	ACTCGCCCTCCCTGGCCGGA
ASCL1.11	CTGCTGCTTCTGCTTTTTTT
RAB5A.1	TCCTCCTCCGCCGCCGCCGCGC
RAB5A.2	TTCCGCGGGCGGGGCGAGGCAGG
RAB5A.4	GTACAGTAAAGAGCGAAGGGAAA
RAB5A.5	GGCTGGGGGGTCTCTGGGCTCCT
RAB5A.6	GTTCTGTGAAGGAGCCGGCGGCTG
RAB5A.7	GGGACTGACTGAGGGAGCGACGG
RAB11A.1	AAGAGGTAGTCGTACTCGTCGTC
RAB11A.2	AAGGTGAGGCCATGGGCTCTCGC
RAB11A.4	GTCACTAAGTAATTGAACAATA
RAB11A.5	CCCTTTGAGCCTCCTTTAGCGAC
RAB11A.6	GCGACTAAAGCTTGAAGCCCCAC
RAB11A.7	GCTCCTCGGCCGCGCAATGGGCA

qPCR Primer

Gene	sequence
ADRB2_forward	TTGCTGGCACCCAATAGAAGC
ADRB2_reverse	CAGACGCTCGAACTTGGCA
ASCL1_forward	CCCAAGCAAGTCAAGCGACA
ASCL1_reverse	AAGCCGCTGAAGTTGAGCC
CCR4_forward	TCTCGCCAAGACACTGAACAG
CCR4_reverse	GGCCCTGCATTCTCAAGAAG
CHRM3_forward	GGCCTGTGCCGATCTGATTAT
CHRM3_reverse	CGGCCTCGTGATGGAAAAG
CHRM4_forward	CAGCTCGGGCAATCAGTCC
CHRM4_reverse	GCCTATGATGAGATCAGCACAC
EEA1_forward	AGCAACTCCTATAAACACAGTGG
EEA1_reverse	AGCAAGATTAGACTCTCCTCCAT
FZD1_forward	AGCCATCCAGTTGCACGAG
FZD1_reverse	GAGTCGGGCCACTTGAAGTT
HBB_forward	AGGAGAAGTCTGCCGTTACTG
HBB_reverse	CCGAGCACTTTCTTGCCATGA
HTR6_forward	GCAACACGTCCAACCTTCTTCC
HTR6_reverse	TGCAGCACATCACGTCGAA
IL1B_forward	AAACAGATGAAGTGCTCCTTCC
IL1B_reverse	AAGATGAAGGGAAAGAAGGTGC
IL1R2_forward	ATGTTGCGCTTGTACGTGTTG
IL1R2_reverse	CCCGCTTGTAATGCCTCCC
PIK3C3_forward	CCTGGAAGACCCAATGTTGAAG
PIK3C3_reverse	CGGGACCATACACATCCCAT
PLCB1_forward	GGAAGCGGCAAAAAGAAGCTC
PLCB1_reverse	CGTCGTCGTCACCTTCCGT
PRKC1_forward	GACAACGAACAGCTCTTCACC
PRKC1_reverse	CCAGGACGTTCTGGTACACA
RAB5A_forward	AGACCCAACGGGCCAAATAC
RAB5A_reverse	GCCCCAATGGTACTCTCTTGAA
RAB7A_forward	GTGTTGCTGAAGGTTATCATCCT
RAB7A_reverse	GCTCCTATTGTGGCTTTGTACTG
RAB9A_forward	AGGGACAACGGCGACTATC
RAB9A_reverse	TCTGACCTATCCTCGGTAGCA
RAB11A_forward	CAACAAGAAGCATCCAGGTTGA
RAB11A_reverse	GCACCTACAGCTCCACGATAAT
ZFP42_forward	AGAAACGGGCAAAGACAAGAC
ZFP42_reverse	GCTGACAGGTTCTATTTCCGC

NGS primer

CD47_forward	ACACTCTTTCCTACACGACGCTCTTCCGATCT NNAAGATCTTACAGTACAGACTTC
CD47_reverse	GTGACTGGAGTTCAGACGTGTGCTCTTCCGATC NNCAGGACAAATAAAAAAAGAAGC

Indel analysis primer

ASCL1_forward	CGAGGAACCCGAAGAGAATAACAGTGAG
ASCL1_reverse	CCGCTGGCGCCTTCTTGTTTCTA
CD47_forward	TATCAGTTCAGCAAGTTCTATTTAGCAGTGTGTT
CD47_reverse	AGTACATTACCCCAGGACCAATCAGCCAAA
DNMT1_forward	CTGGGACTCAGGCGGGTCAC
DNMT1_reverse	CCTCACACAACAGCTTCATGTCAGC
EMX1_forward	TGGATGCCCGTGTCATTAAGAGAGAGACTTT
EMX1_reverse	CCCTTCTGTGAATGTTAGACCCATGGGAGC
FANCF_forward	GTATTAGGGCTTTTAAGTTGCCAGAGTCAAGGA
FANCF_reverse	GTCTGTTAGCAGACCCAGATAGACAGGAGAC
GRIN2B_forward	GAATGCAGGGCTTGTGTACTTATAGCCCC
GRIN2B_reverse	ACAAATGCATGGTTTAGTCCTCAGCACAAAC
IL1B_forward	CCATGAGATTGGCTAGGGTAACAGCA
IL1B_reverse	AGAGACAGAGAGACTCCCTTAGCACC
PIK3C3_forward	GTGGGCGCCTTGTGCACATGC
PIK3C3_reverse	CACCTCCCCTGCTAATACACCATGTGCTC
RAB5A_forward	CGCCCCGCGAACAAACCTAGGC
RAB5A_reverse	CCAGGACGGAGACCAGGCGGAACC
VEGFA_forward	AAACTCCCCCACCCCCTTTCC
VEGFA_reverse	ATTCCAGCACCGAGCGCCC