

Competitive precision genome editing (CGE) assay for functional analysis at single base resolution

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Abstract

The two major limitations of applying CRISPR/Cas9-technology for analysis of the effect of genotype on phenotype are the difficulty of cutting DNA exactly at the intended site, and the decreased cell proliferation and other phenotypic effects caused by the DNA cuts themselves. Here we report a novel competitive genome editing assay that allows analysis of the functional consequence of precise mutations. The method is based on precision genome editing, where a target sequence close to a feature of interest is cut, and the DNA is then repaired using a template that either reconstitutes the original feature, or introduces an altered sequence. Introducing sequence labels to both types of repair templates generates a large number of replicate cultures, increasing statistical power. In addition, the labels identify edited cells, allowing direct comparison between cells that carry wild-type and mutant features. Here, we apply the assay for multiplexed analysis of the role of E-box sequences on MYC binding and cellular fitness.

Introduction

Adaptation of bacterial clustered regularly interspaced short palindromic repeats (CRISPR)–CRISPR-associated protein 9 (Cas9) system for editing of mammalian genomes has provided a robust and efficient tool for sequence manipulations. CRISPR/Cas9-based methods have been extensively used in mutating specific genomic loci in single-gene studies¹ as well as genome-wide screens^{2,3,4}. However, resolution of the CRISPR/Cas9-editing is limited by the suitable protospacer adjacent motif (PAM) sequences found in the close proximity of the region of interest. Homology-directed recombination (HDR)-mediated precision editing can be used to introduce genetic alterations exactly at the intended loci, but this method suffers from strong DNA damage response, low efficiency, and the incompatibility with pooled CRISPR-screening approaches. Because of the low efficiency of precision genome editing, pooled screens commonly use either Cas9 nuclease alone that generates a series of insertion and deletion alleles, or nuclease dead Cas9 fused to a transcriptional repressor (CRISPRi) or activator (CRISPRa) elements^{5,6,7}. These methods do not have single base or single allele resolution, and their precision is limited because they use an indirect measure, inferring the perturbation from the presence of a guide-sequence integrated into the cells at a (pseudo)random genomic position.

Furthermore, interpreting the functional consequence of targeted Cas9-induced mutations is confounded by the DNA damage introduced by Cas9, and from off-target effects of the Cas9 nuclease⁸. In particular, double-strand breaks (DSB) at on- or off-target loci cause DNA damage and genomic instability resulting in paused cell cycle or apoptosis^{9,10,11}. These problems are particularly acute in analysis of small intergenic features, such as transcription factor binding sites. This is because sgRNAs cannot be selected from a large number of possible sequences predicted to have the same effect, and the flanking sequences of most TF binding sites are generally less complex than sequences found in coding regions.

Results and discussion

Here we describe a novel competitive precision genome editing (CGE) approach utilizing CRISPR/Cas9 genome editing at precise loci to accurately analyse the effect of mutations on cellular properties and molecular functions, such as fitness, transcription factor (TF) binding, or mRNA expression. Importantly, the CGE method utilizes CRISPR/Cas9 technology combined with one or several HDR templates with sequence tags enabling lineage-tracing of the targeted cell populations. One of the HDR templates (control) reconstitutes the wild-type sequence of the region of interest by harboring the original genomic sequence, whereas the other template will replace it with desired mutated sequence, such as non-functional TF binding site (**Fig. 1a**). Importantly, each individual HDR template molecule has variable sequence tag(s) flanking the sequence of interest that can be detected from the Illumina sequencing reads of the target locus (**Fig. 1a**). This enables precise counting of the editing events facilitated by each repair template and direct comparison of the effect of the mutation to original genomic sequence. In addition, inclusion of a large set of different sequence tags allows excluding the possibility that the tags themselves, and not the intended mutations, cause the observed phenotype.

In the CGE experiment, DNA samples from cells edited with either mutant or control sequence are collected at two or more time points (early and late) and the cell lineages with particular editing event can be followed before and after subjecting the cells to selection pressure, such as competitive growth in culture, after which cellular fitness, TF binding to the target locus, and the expression levels of mRNA can be analyzed (**Fig. 1b**). Since the sequence tags are present in both repair templates, this experimental design allows precise comparison of the mutated vs. control sequence by excluding the non-edited wild-type sequences from the analysis. Sequencing reads will then be assigned to the distinct editing events based on their sequence tags, and the ratio between mutated and control sequences for

each tag can be determined at both time points resulting in dozens of replicate measurements for each editing event (**Fig. 1a**). Thus, statistical power to detect differences between the time points is very high. The experiment is a single-well assay in which both the repair templates are transfected to cells within one culture well and the genomic perturbation is compared directly to control in the same cell population. This eliminates the experimental bias and variation coming from transfection/transduction and Cas9 introduced DSBs, and well-to-well variation caused by differences in culture and experimental conditions between wells.

To preserve potentially functional flanks of the sequence of interest, it is important to introduce silent or near-silent mutations. For coding regions, this can be accomplished by introducing synonymous mutations of codons and avoiding splice-junctions. Since less is known about functional elements within non-coding regions, we decided to use a diverse library that largely conserves the wild-type sequence, introducing only one or few point mutations per cell within a region wider than a typical TF binding site (~ 10 bp). In our case, each of the ten positions within the sequence tag was mutated with probability of 24%, thus keeping most of the flanking sequence intact (**Fig. 1a**) but introducing typically 2-3 mutations per repair oligo (in ~53% of the sequences; **Supplementary Fig. 1a**). This generates 30 distinct sequences whose sequence differs from the native sequence by exactly one nucleotide (**Supplementary Fig. 1b**), 405 distinct sequences with 2-nt difference to the native sequence, and 3240 distinct sequences with three mutations. In the oligo synthesis for HDR templates, the probability for any individual sequence tag with one mutation is higher than for tag with two or three mutations, which is reflected in the data with single-mutation tags having higher read counts than double and triple mutants (**Fig. 1c**), consistent with the fact that single-mutation sequence tags are present in the original mixture of synthesized oligos in more copies than double- and triple-mutants. After assigning the read counts for each editing event with mutated or native sequence at the two experimental time-points, the

ratio of mutated / native sequences can be plotted for each sequence tag separately, enabling robust and accurate measurement for the effect of the mutation on cellular fitness for each cell lineage separately (**Fig. 1d**).

To evaluate our novel CGE approach in functional studies, a 6-nucleotide binding motif (E-box) for transcription factor MYC was mutated at the promoters of MYC-target genes to study their effect on cell proliferation and fitness. If a particular E-box is essential for cell growth, the alleles containing tags and the wild-type sequence should be enriched in the cell population compared to the E-box deleted alleles after one week of culture, which can be analyzed from genomic DNA collected at the beginning and at the end of the experiment (**Fig. 1b, d**). MYC is an oncogenic transcription factor that regulates expression of genes involved in major cellular pathways promoting cell growth and proliferation, such as ribosome biogenesis and nucleotide biosynthesis¹². However, the functional consequence for cell proliferation resulting from MYC binding to a promoter of a particular gene has not been previously shown. For the purpose of this study, putative MYC target genes were selected for editing on the basis of functional genomics studies in human colon cancer cell lines and previously published data sets in HAP1 haploid cell line using the following criteria: (1) Gene must be essential in HAP1 cells, i.e. found in both publications (refs ^{13,14}), and (2) display robust MYC binding at the gene promoter within open chromatin on the basis of ATAC-seq signal, and clear change in expression upon MYC silencing in colon cancer cells (ref. ^{15,16}; **Supplementary Fig. 2**). In addition, (3) the gene should preferably only contain one E-box within the ChIP-nexus peak (ref. ¹⁵; **Supplementary Fig. 2**).

The experiments were carried out in HAP1 haploid cell line that is derived from chronic myelogenous leukemia cell line KBM-7. HAP1 cells grow adherently and no longer express hematopoietic markers. Most of these cells in early passage cultures are haploid for all chromosomes, and thus particularly useful for mutational screens since only one editing

event is sufficient for a full knock-out. Importantly, targeted mutation of the E-box sequence to a non-functional TATTTA at the promoters of four MYC target genes – ribosomal protein L23 (*RPL23*), hexokinase 2 (*HK2*), phosphoribosyl pyrophosphate amidotransferase (*PPAT*), and midasin AAA ATPase 1 (*MDN1*) – resulted in reduced cell growth as measured from the read counts for lineage-tracing sequence tags with two mutations at day 8 as compared to day 2 (**Fig. 2a**). However, there were E-boxes at promoters of MYC target genes that can be mutated to non-functional sequence without affecting cell proliferation, such as serine hydroxymethyltransferase 2 (*SHMT2*) and phosphoribosylaminoimidazole carboxylase and phosphoribosylaminoimidazolesuccinocarboxamide synthase (*PAICS*; **Fig. 2a**), demonstrating the strength of this approach in dissecting the contribution of each individual transcription factor binding site to cell proliferation. Furthermore, the CGE assay can robustly measure the effect of each E-box on cellular fitness also for genes that harbor several of them within their regulatory region, as demonstrated for the *MDN1* gene. Out of the two E-boxes within the *MDN1* promoter, mutation of the E-box closer to the TSS (TSS +32) had an effect on cell proliferation, whereas the mutation of the E-box farther away (TSS -151) had no effect, despite MYC binding detected at both of these sites using ChIP-nexus (ref. ¹⁵; **Supplementary Fig. 3**).

After demonstrating the power of the precision genome editing assay in analyzing the functional consequence of mutating TF binding sites within the gene regulatory regions, we set out to validate the approach by introducing mutations to the coding regions of genes. To this end, we mutated previously described phosphorylation sites of the cyclin-dependent kinase 1 (*CDK1*) and the growth factor receptor-binding protein 2 (*GRB2*) genes. In coding region, sequence tags were generated by randomizing the degenerate positions of the adjacent codons in the repair template. Phosphorylation sites were abolished by alanine (A) or phenylalanine (F) substitutions of the phosphorylated serine (S), threonine (T) or tyrosine (Y)

residues. To mimic phosphorylation, the same amino-acids were also mutated to the acidic residues glutamate (E) or aspartate (D), which in many proteins can lead to the same effect as phosphorylation of the serine, threonine and tyrosine residues¹⁷.

Previous mutagenesis screen by Blomen et al. (ref. ¹³) suggests that BCR/ABL-GRB2-RAS/MAPK signal transduction is essential for KBM-7 but not for HAP1 cells. To test this, we mutated two key residues, Y160 and Y209 in the adaptor protein GRB2 that links tyrosine kinase signaling to the RAS-mitogen-activated protein kinase (MAPK) pathway. GRB2 phosphorylation at Y209 impairs interaction between GRB2 and Son-of-sevenless (Sos), which promotes GTP binding to RAS and activation MAPK signaling¹⁸. Phosphorylation of Y160 acts in opposing fashion by favoring the monomeric state of GRB2 that is capable of interacting with Sos¹⁹. In agreement with this, we observed that KBM-7 but not the HAP1 cells were sensitive to GRB2 mutation (**Supplementary Fig. 4**), indicating that the precision genome editing assay can be used to identify functionally important phosphorylation events in cells.

We next used the CGE assay to evaluate the fitness effect of CDK1 regulatory phosphorylation site mutations in human cells. CDK1 activation and onset of mitosis requires phosphorylation of T161 in the activation segment, and dephosphorylation of T14 and Y15²⁰. The non-phosphorylatable double-mutant T14A/Y15F cells were almost completely lost after one week of precision editing (**Fig. 2b**). These findings are consistent with earlier work reporting that the T14A/Y15F double-mutant can be activated prematurely during the cell cycle²¹, and overexpression of this mutant in cells results in cell death due to mitotic catastrophe²². The effect of the phosphorylation site mutation in the CDK1 activating segment, T161A, was less prominent. Loss of phosphorylation resulted in markedly decreased cell proliferation, whereas T161E phosphomimetic mutation allowed cells to proliferate normally (**Fig. 2b**). This is consistent with the lack of requirement of regulation of

the CDK activating kinase in human cells²³. We also tested recently reported prime editing method²⁴ for mutating a phosphorylation site and for introducing the sequence tag within the *CDK1* coding region. Using this approach, we observed reduced fitness of HAP1 cells as a result of Y15F mutation (**Fig. 2b**), demonstrating that also prime editing can be utilized for generating the targeted mutations and sequence tags for our precision genome editing assay.

Since the competitive precision genome editing assay showed clear effects on cell proliferation resulting from a mutation of a single TF binding motif or protein phosphorylation site, we set to analyze the direct effects of E-box mutation on MYC binding and histone 3 lysine 27 acetylation (H3K27ac). For this, we performed chromatin-immunoprecipitation using anti-MYC and anti-H3K27ac antibodies from the HAP1 cells after precision editing. To quantify the editing events, each targeted locus was amplified using PCR and the amplicons were Illumina sequenced. Importantly, we detected fewer antibody-enriched sequences with TATTTA mutated sequence compared to CACGTG original sequence, demonstrating less MYC binding to the mutated sequences at RPL23, MDN1, and SHMT2 E-boxes, as opposed to the input sample with equal ratios of TATTTA and CACGTG (**Fig. 2c**). We also observed decrease in H3K27 acetylation at RPL23 and MDN1 E-boxes (**Fig. 2c**), suggesting lower expression of these target genes. However, there was no changes in the level of H3K27 acetylation at SHMT2 locus, consistent with the observation that mutation of this E-box had no effect on cell proliferation (**Fig. 2a, c**).

Here we show a novel method for precise analysis of the effect of mutations on cellular phenotype by utilizing CRISPR/Cas9 precision editing combined with lineage-tracing sequence tags in HDR templates. Previously, next-generation sequencing based methods, such as GUIDE-seq, have been developed for assessing the off-target DNA cleavage sites²⁵ and random sequence labels have been used for increasing precision and accuracy of CRISPR-screens²⁶. In a recent saturation mutagenesis screen, a single-nucleotide

variants (SNV) targeting BRCA1 gene were transfected to target cells along with Cas9 and sgRNA and targeted gDNA and RNA sequencing was performed to quantify SNV abundances²⁷. However, our approach enables for the first time a precise assessment of the effect of particular intended mutations. Our approach of using parallel editing of the target loci with two HDR templates in a single cell culture has two key advantages over previously described genome-editing assays. First, silent mutations that generate sequence tags to HDR templates provide means to discard all confounding information from the next-generation sequencing output of the assay. Second, direct comparison of the mutated sequence to the reconstituted native sequence mitigates all the detrimental off-target effects, as well as enables lineage tracing of edited clones thus providing statistical power to the analysis. When measuring allele-specific phenotypes, the method also allows the use of diploid cells for analysis of phenotypes such as TF binding or RNA expression. Measuring more complex phenotypes in diploid cells is also possible, but requires either prior deletion of one allele from the targeted locus, or dilution of the two repair templates by a template that inactivates the wild-type allele in such a way that most cells carry either two inactive alleles, or one inactive allele, and one targeted allele. This will be easier when targeting coding regions, as failure of targeted repair commonly leads to inactivation of the target gene due to generation of frameshift or deletion alleles by non-homologous end-joining.

In summary, we report here an advanced method for measuring the phenotypic effects of precise targeted mutations. The method allows controlling for the effect of DNA damage, the major confounder in CRISPR-based methods. We also demonstrate the power of the technology by robustly detecting small fitness effects of individual transcription factor binding motifs and single amino-acid substitutions. The method is widely applicable and extends the utility of CRISPR/Cas9-mediated genome editing to address important biological questions that have been difficult to address using existing technologies.

Methods

Genome editing constructs

Precision editing of each genomic locus was performed by introducing a CRISPR/Cas9-mediated DSB and HDR templates harboring either the mutated or original genomic feature along with a sequence tag. Guide sequences were designed using CRISPOR²⁸ tool, giving preference to the protospacers with closest distance to the genomic feature to be edited, and the crRNAs were obtained from Integrated DNA Technologies (**Supplementary Tables 1, 2**). Single-stranded 100-nucleotide (nt) DNA molecules were used as HDR templates. For editing E-box sequences, two HDR templates were designed for each targeted locus, one with CACGTG sequence for reconstituting the original E-box, and another with mutated sequence that replaces the E-box with non-functional TATTTA sequence. In each oligo, the original or mutated E-box was flanked by a 10-nt sequence tag and two 42-nt homology arms complementary to the target strand. Sequence tags were generated by mutating each of the ten nucleotides with probability of 24%, i.e. 8% probability for each of the three non-consensus bases (oligo synthesis using custom hand-mixed bases from Integrated DNA Technologies; **Supplementary Table 1**). As a negative control for E-box mutation experiments, the coding region of the *MYC* gene was targeted with two HDR templates, one reconstituting the original coding sequence, and another replacing nucleotides encoding Val-5 and Ser-6 with synonymous codons (GTTAGC > GTAAGT), and the sequence tag was created by randomizing the third degenerate position in the two codons flanking the targeted region on both sides. For targeting protein phosphorylation sites in the coding regions of the *CDK1* and *GRB2* genes, HDR oligos with 40-nt homology arms and sequence tags generated by randomizing the degenerate positions of the adjacent codons in the repair template were designed (**Supplementary Table 2**). Phosphorylation sites were abolished by alanine or phenylalanine substitutions of the

phosphorylated serine/threonine or tyrosine residue, as well as mutated to glutamate or aspartate that in some cases mimic the phosphorylated state¹⁷.

Prime editing guides (pegRNA) to target CDK1 were designed according to the recommendations from ref. ²⁴. Similar to HDR templates described above, the pegRNA pool introduces a mutation (Y15F) or reconstitutes the original sequence, and in both cases the third degenerate position in the codons flanking the targeted region was randomized to create the sequence tags for lineage-tracing (**Supplementary Table 2**).

PCR primers for amplifying genomic DNA at each targeted locus were designed so that neither forward nor reverse primers overlap with the genomic sequence used in the HDR templates. All custom oligoes used for targeting and amplifying the E-box targets and the phosphorylation sites are listed in **Supplementary Tables 1 and 2**, respectively.

Cell lines and transfections

HAP1 (#C631) and KMB-7 (#C628) cell lines were obtained from Horizon Discovery and maintained in low-density cultures in Iscove's Modified Dulbecco's Medium (IMDM) with 10% FBS, 2 nM L-glutamine, and 1% antibiotics according to the vendor's guidelines.

Precision editing experiments measuring cellular fitness were done by transfecting 200,000-400,000 early-passage HAP1 or KBM-7 cells with ribonucleoprotein (RNP) complex and the two HDR templates. For sgRNA molecules, equimolar ratios of target-specific crRNAs and ATTO550-tracrRNA (Integrated DNA Technologies) were annealed. RNP complexes used for the transfections were constituted from S.p. HiFi Cas9-protein (Integrated DNA Technologies; 1000 ng / 200,000 cells) and target-specific sgRNA (250 ng / 200,000 cells) and transfected to cells using CRISPRMAX (Life Technologies) as per manufacturer's recommendation along with HDR template (1:1 mixture of the original and mutant HDR templates) with final concentration of 3 nM. Half of the cell population was

harvested for genomic DNA isolation 48 h after transfection (day 2), and the rest of the cells were plated for culture and harvested on day 8. For ChIP assays measuring the effect of E-box mutation on MYC occupancy and H3K27 acetylation, 15 million cells were transfected for each condition on two 15-cm dishes, scaling up the transfection reagents according to the cell numbers, and the cells were harvested, and chromatin cross-linked 48 h after transfection.

For prime editing experiments, Prime editor 2 was expressed from pCMV-PE2 and pegRNAs from pU6-pegRNA-GG-acceptor plasmids²⁴ (Addgene #132775 and #132777, respectively). Plasmid transfection was performed using FuGENE HD (Promega) according to manufacturer's instructions. Rest of the experiment was performed in the same way as the homology-directed repair editing experiment.

Genomic DNA isolation and target-specific sequencing

Genomic DNA was isolated using AllPrep DNA/RNA Mini kit and Blood & Cell Culture DNA Maxi kit (Qiagen) from day 2 and day 8 time points, respectively, and treated with RNase A (0.2 µg/ul; Thermo Fisher Scientific) for 2 h at 37 C. To eliminate the carry-over of single-stranded DNA from the HDR templates, gDNA samples were treated with exonuclease I and VII (New England Biolabs) in 10 mM Tris-HCl, 50 mM KCl, 1.5 mM MgCl₂ for 30 min at 37 C followed by enzyme inactivation for 10 min at 95 C and DNA extraction using phenol:chloroform:isoamyl alcohol (Sigma). All gDNA from day 2 and 10 µg gDNA corresponding to 3 million haploid cells from day 8 was amplified in PCR reactions with maximum of 2.5 µg of DNA per reaction using NEBNext High Fidelity master mix (New England Biolabs) and target-specific primers with Illumina adaptor flanks (**Supplementary Tables 1, 2**) for 20 cycles, followed by DNA purification using 1.5x Ampure XP beads (Beckman Coulter). For amplifying the genomic regions with E-box

targets, biotinylated primers were used for PCR1, and the 30% volume of the purified PCR products was used for streptavidin capture with M-280 Dynabeads (Thermo Fisher Scientific) according to manufacturer's protocol. Prime-editing samples are not affected by the presence of HDR template, and thus they were prepared without the exonuclease I/VII treatment and affinity purification of biotinylated PCR products. Second PCR amplification with 8 cycles was used for generating sequencing-ready libraries using NEBNext High Fidelity master mix and Illumina Universal and Index primers (E7335S, New England Biolabs) in four and twelve parallel reactions on M-280 beads for day 2 and day 8 samples, respectively. PCR products were purified using 0.9x Ampure XP beads and sequenced for 150 cycles on NovaSeq 6000, HiSeq 4000, and NextSeq 500 platforms (Illumina).

Analysis of precision editing data

Sequencing reads were assigned to each genomic target by matching the first 20 nucleotides of each read to the sequence of PCR products amplified using target-specific primers. Then, the reads originating from cells that had undergone successful precision editing were identified based on the presence of a sequence tag, and the reads originating from non-edited wild-type cells were discarded. Reads matching to each individual sequence tag flanking the original and mutated sequence features were counted in different timepoint samples, and a pseudocount of +1 was added to each read count to avoid the zeros in subsequent calculations. For experiments targeting E-boxes, all cell lineages harboring sequence tags with read count > 50 in both day 2 samples were included in the analyses. In **Fig. 1d**, the results from all sequence tags with exactly one flanking mutation are shown to demonstrate the power of the assay in tracing the growth of individual cell lineages over time. However, to further increase the robustness of the analysis, only the sequence tags with exactly two flanking mutations were included in the analyses for **Fig. 2a, c** and

Supplementary Fig. 3. For experiments targeting protein phosphorylation sites, all the sequence tags with read count > 20 for GRB2 experiments and > 5 for CDK1 experiments were included in the analyses due to lower sequencing depth in these experiments.

To analyze the effect of each mutation on cellular fitness, the ratio of cells harboring mutated and original sequence features was compared at each time point. If mutating a particular E-box or phosphorylation site hampers cell growth or proliferation, the cells harboring the mutated allele will be underrepresented in the final pool of cells after one-week culture compared to the cells harboring the original sequence feature, and vice versa. Thus, the read count ratios were calculated for each sequence tag for mutated vs. original sequence. To eliminate the potential effect of near-silent flanking mutations on cellular fitness, the sequence tags with similar flanking mutations were compared in the analysis. Finally, the ratio for day 8 vs. day 2 was calculated for each sequence tag. In **Fig. 2**, the results are presented as \log_2 (fold change) as follows: $\log_2[\text{day 8 read count (mutated / original)} / \text{day 8 read count (mutated / original)}]$ for each cell lineage. Wilcoxon signed rank test was used for testing whether the median of \log_2 (fold change) values is unequal to zero.

Chromatin-immunoprecipitation (ChIP) with target-specific sequencing and ChIP-seq

Wild-type HAP1 cells and genome-edited HAP1 cells 48 h after RNP transfection were crosslinked with 1% formaldehyde and chromatin samples were prepared as described previously²⁹. Chromatin was sonicated to an average fragment size of 500 bp using micro-tip sonicator (Misonix Inc) and used for immunoprecipitation (IP) with antibody-coupled Dynal-beads (Thermo Fisher Scientific) for MYC, H3K27ac, and normal rabbit IgG (Millipore #06-340, Abcam #ab4729, and SantaCruz #sc-2027, respectively). Chromatin from ten million wild-type cells and 20 million transfected cells was used for each IP. After overnight incubation, washed with LiCl buffer and reverse cross-linking was performed as described in

ref. ²⁹, followed by DNA purification using phenol:chloroform:isoamyl alcohol and ethanol precipitation.

All immunoprecipitated DNA isolated from transfected cells was amplified for 30 cycles in two reactions using similar PCR strategy and conditions as described above for genomic DNA samples. In addition, 10 µg of input DNA from each transfected condition was similarly amplified in four parallel reactions. PCR products were purified using 1.5x Ampure XP beads and 20% of purified DNA was used as a template for the second PCR amplification step for 8 cycles with Illumina primers as above. Final libraries were purified using 0.9x Ampure XP beads and sequenced for 150 cycles on NovaSeq 6000 (Illumina). Data was analyzed essentially as described for fitness experiments: after excluding the reads originating from wild-type cells, a pseudocount +1 was added to the reads originating from the edited cells with distinct sequence tags, the read count ratios between mutated and original sequences were calculated for each condition, and $\log_2(\text{fold change})$ between each IP and respective input sample was calculated. Only cell lineages harboring sequence tags with read counts > 50 in the input sample were included in the analyses. Wilcoxon signed rank test was used for testing whether the median of $\log_2(\text{fold change})$ values is unequal to zero.

Wild-type HAP1 samples were used for standard ChIP-seq library preparation with NEBNext Ultra II DNA Library Prep kit (New England Biolabs), followed by sequencing on NovaSeq 6000. The reads were aligned to human genome (hg19) using bowtie2³⁰ and peaks were called using MACS2³¹ against input with default narrow peak parameters. The bedgraph files were used for genome browser snapshots. For colon cancer cell lines GP5d, LoVo, and COLO320DM, previously published ChIP-nexus data sets from ref. ¹⁵ (EGAD00001004099) were used. In the genome browser snapshots, the traces from BAM coverage files are shown.

Chromatin accessibility and gene expression analysis

ATAC-seq for chromatin accessibility was performed from 50,000 HAP1 cells as previously described³². Briefly, the cells were washed with ice-cold PBS, lysed in 50 µl of lysis buffer for 10 min in ice, and treated with Tn5 transposase in 2x tagmentation buffer (Illumina) for 30 min at 37 C. DNA was purified using MinElute PCR Purification kit (Qiagen) and prepared for sequencing using Nextera library preparation kit (Illumina) by five cycles of PCR amplification. The sample was sequenced on NovaSeq 6000 for 2 x 50 cycles, and the paired-end data was analyzed using an in-house pipeline as described in ref. ¹⁶. For GP5d cells, the ATAC-seq data from ref. ¹⁶ (GSE180158) was used. In the genome browser snapshots, the traces from BAM coverage files are shown.

For gene expression analysis, previously published RNA-seq data from ref. ¹⁵ (EGAD00001004098) was used. The data sets for MYC silencing using siRNA (siMYC) and respective control samples transfected with non-targeting siRNAs (siNon-target) for GP5d and LoVo cells were re-analyzed by aligning the reads from fastq files to GRCh37 human genome using tophat2³³ and by analyzing the differentially expressed genes between siMYC and siNon-target samples using cuffdiff³⁴ using default parameters and the option for first-strand library type. Log₂(fold change) values for the selected genes are shown in

Supplementary Fig. 2.

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Data availability: All next-generation sequencing data generated in this study is available under ENA accession XXX. Previously published data sets for colon cancer cells were used as follows: RNA-seq from EGAD00001004098, ATAC-seq from GSE180158, and ChIP-nexus from EGAD00001004099.

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Figures and figure legends

Figure 1

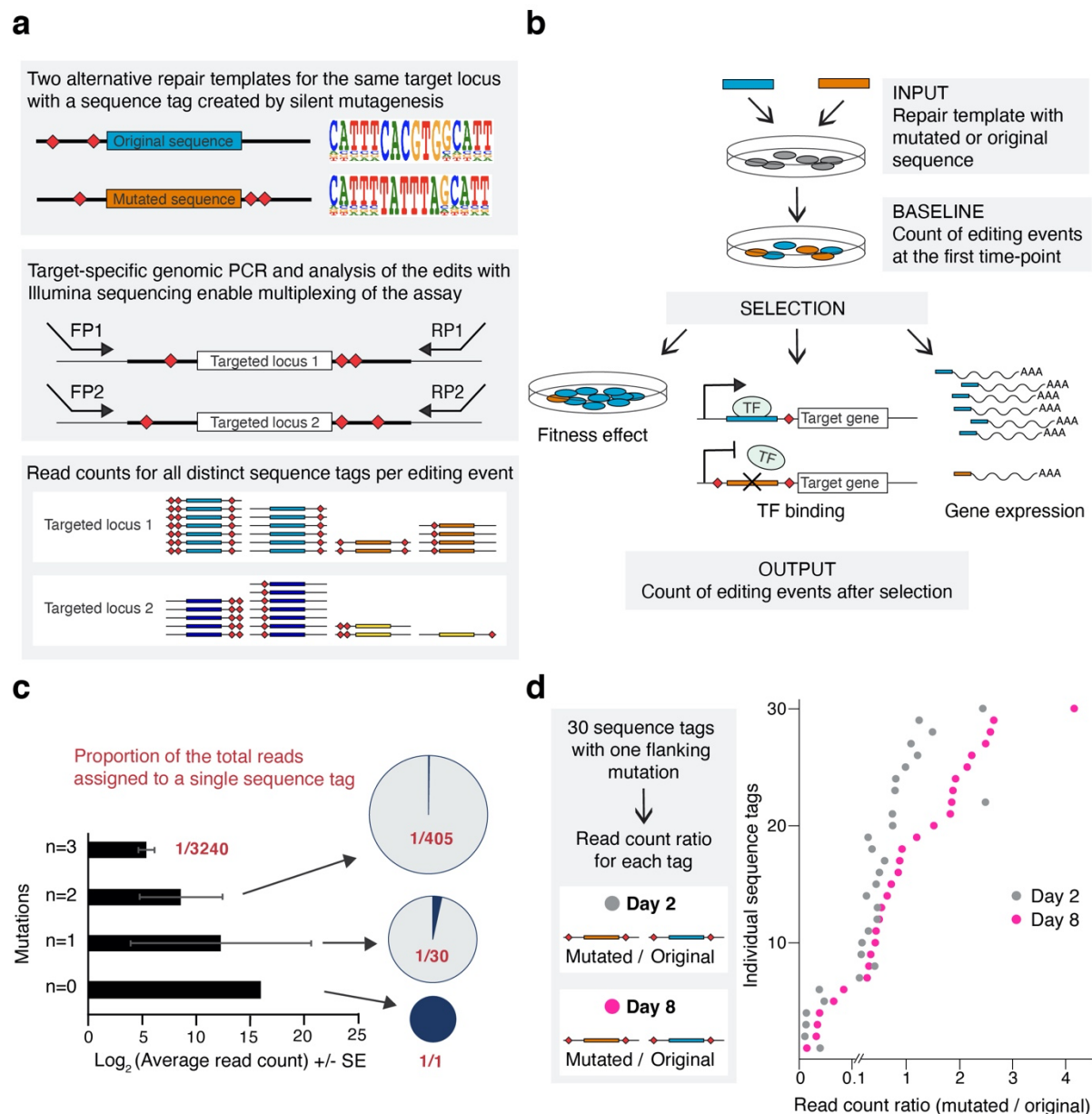


Figure 1. Strategy for lineage-tracing of cells with distinct genome editing events utilizing sequence tags with silent mutations.

a, Conceptual outline of the precision editing strategy utilizing two HDR templates, one harboring original genomic sequence and another with the desired mutation. Five nucleotides on both side flanking the region of interest are mutated in the HDR oligos with the probability of 24%, a strategy that maintains most of the flanking sequence intact. Editing events at

targeted locus can be identified using Illumina sequencing and cell lineages traced according to their mutation signatures.

b, Schematic presentation of the experimental workflow utilizing a 1:1 mixture of two HDR templates for the same target. The abundance of each HDR template in the cell population can be analyzed at the baseline (first time point), and after selection pressure using different output assays, such as cellular fitness (after one week of culture), TF binding (after immunoprecipitation with specific antibody), and mRNA expression.

c, Number of distinct sequences with zero, one, two, and three mutations in the ten flanking nucleotides when mutated with the probability of 24% and their average read counts from the Illumina sequencing reads at the targeted SHMT2 locus at baseline (from ChIP input sample). Pie charts illustrate the proportion of the reads assigned to each distinct sequence tag with different number of mutations.

d, The effect of E-box mutation at the *RPL23* gene promoter on fitness of HAP1 cells. Precision editing results are shown separately for each cell lineage harboring a sequence tag with exactly one flanking mutation. Read count ratios for mutated vs. original sequence are shown in two time points, day 8 (pink) and day 2 (grey).

Figure 2

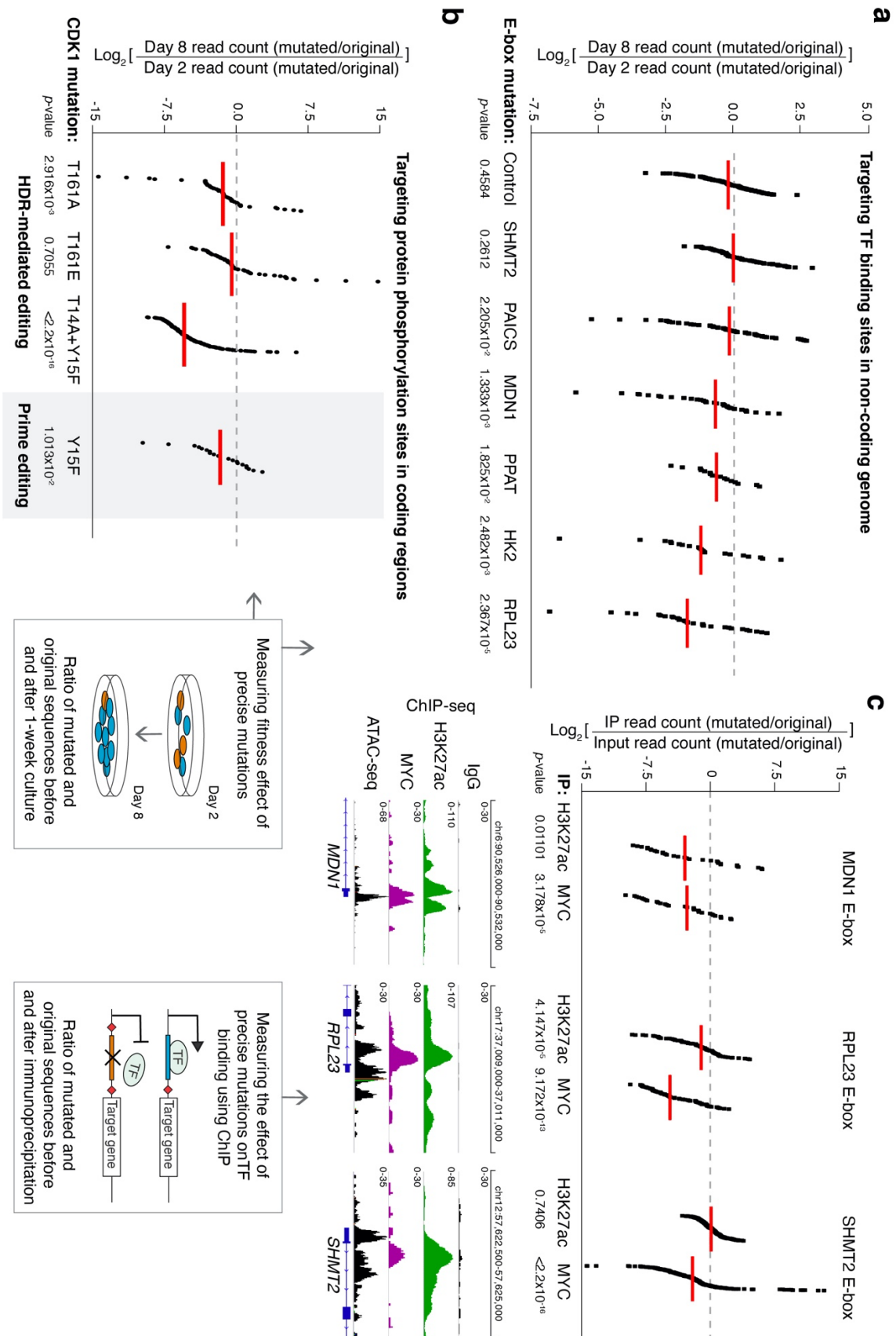


Figure 2. The effect of mutating TF binding sites and protein phosphorylation sites on cellular fitness determined by lineage-tracing of editing events.

a, The effect of mutating MYC binding motifs (E-box) at promoters of MYC target genes on fitness of HAP1 cells. E-box mutation at six MYC target gene promoters (see **Supplementary Fig. 2** for E-box locations) as well as negative control region (synonymous mutation in the MYC coding region) are shown. Precision editing results are shown for all cell lineages harboring sequence tags with exactly two flanking mutations with read count > 50 on day 2. Log₂ values for day 8/day 2 ratios are shown for each sequence tag pair after calculating the ratio of read counts for mutated vs. original sequence at both timepoints. *p*-values from Wilcoxon signed rank test are shown for each experiment.

b, The effects of mutating protein phosphorylation sites in the *CDK1* gene on fitness of HAP1 cells. Precision editing results are shown for all cell lineages harboring sequence tags with read count > 5 on day 2. Log₂ values for day 8/day 2 ratios are shown for each sequence tag pair after calculating the ratio of read counts for mutated vs. original sequence at both timepoints. Note that T161E is a phosphomimetic mutation whereas T161A abolishes the phosphorylation site. Phosphorylations at T14 and Y15 inhibit CDK1 function, but the presence of non-phosphorylatable T14A/Y15F double-mutant can result in premature mitosis and mitotic catastrophe²². The effect of Y15F on the fitness was also studied after introducing this mutation to the HAP1 cells using prime editing²⁴. *p*-values from Wilcoxon signed rank test are shown for each experiment.

c, The effect of E-box mutation on MYC occupancy and acetylation of H3K27 at promoters of MYC target genes *MDN1*, *RPL23*, and *SHMT2*. ChIP using MYC and H3K27ac antibodies followed by target-specific PCR and Illumina sequencing was performed 48 h after RNP transfection. Input from crosslinked and sonicated chromatin was used as a control. Precision editing results are shown for cell lineages harboring all sequence tags with exactly two flanking

mutations with read count > 50 in the input. Log₂ values for immunoprecipitated (IP) sample/input ratios are shown for each sequence tag pair after calculating the ratio of read counts for mutated vs. original sequence in both timepoints. *p*-values from Wilcoxon signed rank test are shown for each experiment. IGV snapshots show ChIP-seq and ATAC-seq tracks from wild type HAP1 cells for each of the targeted loci, demonstrating robust MYC binding to these sites in HAP1 cells.