Mutations and genetic alterations are often sequentially acquired in various biological and pathological processes, such as development, evolution, and cancer. Certain phenotypes only manifest with precise temporal sequences of genetic events. While multiple approaches have been developed to model the effects of mutations in tumorigenesis, few recapitulate the stepwise nature of cancer evolution. Here we describe a flexible sequential mutagenesis system, Cpf1-Flip, with inducible inversion of a single crRNA array (FlipArray), and demonstrate its application in stepwise mutagenesis in murine and human cells. As a proof-of-concept, we further utilize Cpf1-Flip in a pooled-library approach to model the acquisition of diverse resistance mutations to cancer immunotherapy. Cpf1-Flip offers a simple, versatile, and controlled approach for precise mutagenesis of multiple loci in a sequential manner.
In a large variety of biological and pathological processes, genetic mutations or alterations are often acquired in a sequential manner. In evolution and speciation, the genomes of organisms acquire mutations constantly and are subjected to natural selection. In genetically complex disorders such as cancer, multi-step mutagenesis is often a major obstacle for effective treatments. Cancers evolve through an ongoing process of mutation–selection balance, where initial mutations are selected for, or against, in vivo, followed by subsequent acquisition of additional mutations as the tumor grows. Since the initial set of oncogenic “driver” mutations is generally what starts and sustains tumor growth, targeted molecular therapies are often chosen to specifically attack such oncogenic dependencies. However, the selection pressures of treatment will favor secondary mutations that confer drug resistance, leading to relapse. Thus, the process of cancer evolution by sequential mutagenesis stymies these therapies via continuous diversification and adaptation to the tumor microenvironment, eventually exhausting available treatment options. Even with the advent of cancer immunotherapy, where checkpoint blockade is increasingly being utilized in the clinic, the acquisition of secondary mutations that abolish T cell receptor (TCR)–antigen–major histocompatibility complex recognition can still lead to immune escape and ultimately negate the effect of immunotherapy. Thus, the ability to perform sequential and precise mutagenesis is critical for studying biological processes with multi-stage genetic events such as development and evolution, as well as the pathogenesis of complex diseases such as cancer.

From a genetic engineering perspective, stepwise mutagenesis or perturbation is a powerful technique for precise genetic manipulation of cells and live organisms. Multiple methods have been employed to achieve this end. In the pre-recombinant DNA era, stepwise perturbation was often done by multiple rounds of random mutagenesis using chemical or physical carcinogens followed by artificial selection. The subsequent discovery and application of recombinase systems such as Cre-loxP, Flp-FRT, and σC31-att enabled inducible genetic events. In these systems, the DNA recombinase (i.e., Cre) specifically recognizes its target DNA sequence motif (i.e., loxP) and catalyzes recombination between two such target sites. Depending on the configuration of the target sites, targeted recombinases can be utilized for DNA excision, translocation, and/or inversion. However, the floxed genomic loci underlying Cre-based systems must be pre-engineered on a gene-by-gene basis. This process of generating new floxed alleles for each unique application is time and labor intensive, further limiting the feasibility of multiplexed Cre recombination.

More recently, precisely targeted and customizable mutagenesis was simplified by the discovery of RNA-guided endonucleases (RGNs) Cas9 and Cpf1 from clustered regularly interspaced short palindromic repeats (CRISPR) systems. RGNs can induce double-strand DNA breaks, subsequently generating insertions and deletions at the target site. This process is precisely targeted based on the sequences of CRISPR RNAs (crRNAs), which complex with RGNs to enable and guide their nuclease functions. Unlike with Cre recombination, CRISPR crRNAs can be easily transferred to target cells through transfection or viral vectors, thus obviating the need to pre-engineer the host genome for each targeted gene.

In contrast to Cas9, the most widely utilized RGN to date, Cpf1 is a single component RGN that does not depend on trans-activating RNA and can autonomously process crRNA arrays. These features have made Cpf1 particularly attractive for multiplexed mutagenesis. In addition to several studies in mammalian systems, Cpf1-mediated mutagenesis and transcriptional repression have now been successfully applied in plants. Furthermore, chemical modifications on Cpf1 mRNA and crRNAs have been identified that can improve cutting efficiency. It was recently demonstrated that Cpf1 can also process crRNAs from mRNAs expressed by a Pol II promoter, further enabling flexible transcriptional control.

Sequential mutagenesis using Cas9 was demonstrated in ex vivo organoid cultures. However, this approach required sequentially introducing each sgRNA in culture, one at a time, limiting its broader applicability. In particular, the sequential introduction of different sgRNAs would be impractical for library-scale screening or any in vivo experimental designs. To our knowledge, conditional sequential mutagenesis using RGNs has not yet been demonstrated. Here we describe a flexible sequential mutagenesis system through inducible inversion of a single crRNA array (Cpf1-Flip) and demonstrate its simplicity in stepwise multiplexed gene editing in mammalian cells for modeling sequential genetic events, such as in cancer. We further apply Cpf1-Flip to model the acquisition of resistance mutations to immunotherapy in a pooled mutagenesis setting, demonstrating the feasibility of Cpf1-Flip for conducting sequential genetic studies. This system can be utilized for multi-step mutagenesis of any genes in the genome for interrogating complex genetic events with temporal control.

Results

Construction of a Cpf1 sequential mutagenesis system. When loxP sites are arranged such that they point towards each other, Cre recombination leads to inversion of the intervening sequence. However, this process leads to the complete regeneration of the loxP sites, thereby allowing Cre to continually catalyze DNA inversion. As continuous Cre-mediated inversion would be counterproductive in many applications, mutant loxP sites have been characterized that enable unidirectional Cre inversion. When the mutant loxP sites lox66 and lox71 are recombined, they generate a wild-type loxP site and a double-mutant lox72. Cre has a substantially lower affinity for lox72, thus leading to mostly irreversible inversion of the floxed DNA segment.

We designed a U6 expression cassette containing two inverted BsmI restriction sites, flanked by a lox66 sequence and an inverted lox71 sequence (Fig. 1a). In the same lentiviral vector, an EFS promoter drives the expression of Lachnospiraceae bacterium Cpf1 (LbCpf1, or Cpf1 for short) and a puromycin resistance gene (EFS-Cpf1-Puro). After BsmI restriction digestion, the vector linearizes and allows for insertion of a crRNA array. To enable stepwise mutagenesis, we designed crRNA arrays in which the first crRNA is encoded on the sense strand, while the second crRNA is inverted. We hereafter refer to this construct as a crRNA FlipArray. Six consecutive thymidines (6xT) are present in cis at the 3’ end of each crRNA, terminating U6 transcription. Each crRNA is preceded by the LbCpf1 direct repeat (DR) sequence, which guides Cpf1 to process the crRNA array.

Cre-mediated recombination of the lox66 and lox71 mutant loxP sites leads to inversion of the FlipArray, generating a wild-type loxP and a double-mutant loxP, lox72. As the affinity of Cre recombinase for lox72 is substantially lower than for wild-type loxP, inversion of the FlipArray is mostly irreversible. After inversion, the two crRNAs trade places and the second crRNA becomes expressed. Thus, in the absence of Cre, Cpf1 generates indels at the target site of the first crRNA; after Cre recombination, Cpf1 is directed to the target site of the second crRNA. We hereafter term this approach Cpf1-Flip. In short, the Cpf1–Flip system leverages CRISPR-Cpf1 mutagenesis and melds it with the inversion capabilities of Cre/lox66/lox71 to enable programmable two-step mutagenesis.
Sequential mutagenesis in murine and human cells. To demonstrate sequential editing of cancer genes, we first applied Cpf1-Flip to generate Nf1 and Pten mutations in a mammalian lung cancer cell line (KPD)\textsuperscript{26}. We cloned in a FlipArray containing a spacer targeting Nf1 (crNf1) and an inverted spacer targeting Pten (crPten) (crNf1-crPten FlipArray, or NPF). We infected the cells with lentivirus containing EFS-Cpf1-puro; U6-FlipArray. After 7 days, cells were then infected with lentivirus containing EFS-Cre to induce inversion of the FlipArray. Prior to Cre recombination, only crNf1 is expressed; following Cre recombination, crPten becomes expressed.

RNA from the double-infected KPD cells at various timepoints. After CDNA synthesis, we utilized inversion-specific primers to detect inverted crRNA FlipArray transcripts (Fig. 2d). The induction of inverted FlipArray transcripts steadily increased through the course of the experiment, illuminating the kinetics of Cre-mediated inversion of the FlipArray and its subsequent transcription. The low-levels of inverted FlipArray transcripts at baseline could be due to spontaneous inversion, or an artifact of the primer design.

We sequenced the target sites of crNf1 and crPten to determine whether the NPF construct had indeed created mutations in a controlled stepwise manner. Uninfected controls did not have any significant variants at crNf1 (Supplementary Data 1) or crPten target sites (Supplementary Data 2) (Fig. 2c, e, f). 7 days following the initial lentiviral infection with EFS-Cpf1-puro; U6-NPF, indels were found at the crNf1 target site, but not the crPten site (Fig. 2g, h, k). Since the second crRNA is not transcribed prior to Cre recombination, this result affirms that inversion of NPF has not yet occurred at this time point. After another 10 days following infection with EFS-Cre lentivirus (17 days following the initial infection with EFS-Cpf1-puro; U6-NPF), indels were found at both crNf1 and crPten target sites at high frequencies (Supplementary Data 3; Fig. 2i-k).

To further demonstrate the utility of Cpf1-Flip in diverse biological systems, we designed a FlipArray targeting two human...
Fig. 2 Inducible sequential mutagenesis in murine cells through Cpf1-Flip. a Schematic for PCR-based detection of Cre-mediated inversion of the crRNA FlipArray (targeting Nf1 and Pten). b PCR-based detection of non-inverted and inverted FlipArrays at D0 (n = 3) and D10 (n = 3) following Cre, along with input control. c Quantification of gel intensities in b, normalized to input and expressed as a percentage of total FlipArray abundance. d Detection and quantification of Cre-mediated inversion of the crRNA FlipArray at the RNA transcript level using RT-PCR (n = 2 infection replicates). The expression of the inverted FlipArray was assessed at multiple timepoints following EFS-Cre infection using sequence-specific primers for the inverted FlipArray transcript as normalized to the Cpf1 mRNA level. e, f Representative Illumina targeted amplicon sequencing of the crNf1 target site (g) and crPten target site (h) 7 days after infection with lentivirus containing EFS-Cpf1-puro; U6-NPF-FlipArray. Where relevant, the top 5 most frequent variants are shown, with the associated variant frequencies in the boxes to the right. g, h Representative Illumina targeted amplicon sequencing of the crNf1 target site (i) and crPten target site (j) 17 days after infection with lentivirus containing EFS-Cpf1-puro; U6-NPF-FlipArray and 10 days following EFS-Cre infection. Where relevant, the top five most frequent variants are shown, with the associated variant frequencies in the boxes to the right. k Dot plot detailing the total variant frequencies at the crNf1 and crPten target sites in uninfected cells (red), 7 days after FlipArray transduction (−Cre) (green), and 17 days after FlipArray transduction (+Cre) (blue). n = 2 cell replicates for uninfected group, n = 3 for other conditions. All error bars are mean ± s.e.m.
genes, DNMT1 and VEGFA. The crRNA in the first position targets DNMT1 (crDNMT1) while the second, inverted crRNA targets VEGFA (crVEGFA) (crDNMT1-crVEGFA FlipArray, or DVF) (Fig. 3a). Cre activation induces recombination of the lox66/lox71 sites, such that crVEGFA becomes expressed. We transduced human HEK293T cells with EFS-Cpf1; U6-DVF lentivirus, followed by puromycin selection. To assess the functionality of the FlipArray, we then infected the cells with EFS-Cre lentivirus. Using primers specific to the non-inverted or inverted DVF FlipArray, we confirmed that Cre administration drives efficient inversion
consequence of random sampling, as only a subset of the D7 cells to be consistently lower at D21 than at D7. This is likely a role in maintaining DNA methylation. The cutting efficiency of the crRNA itself, as well as inefficiencies in FlipArray expression or subsequent crRNA array processing. Taken together, these results demonstrate that Cpf1-Flip is a flexible tool for sequential mutagenesis based on the Cpf1:crRNA complex, temporally controlled by Cre recombinase.

Modeling acquired resistance to immunotherapy with Cpf1-Flip. We next sought to apply Cpf1-Flip to model acquired resistance to immunotherapy in breast cancer cells (E0771 cell line). We designed a small pool of FlipArrays in which the first crRNA targets Nf1 while the inverted second crRNA targets a panel of immunomodulatory factors (Cd274, Ido1, B2m, Fasl, Jak2, and Lgals9 referred to as TSG-Immune FlipArray library). These factors are thought to influence anti-tumor immunity and have been implicated in acquired resistance to checkpoint inhibitors. After pooled lentiviral transduction of E0771 cells with the TSG-Immune FlipArray library, we infected the cells with EFS-Cre lentivirus to induce FlipArray inversion (Fig. 4a). Upon Cre-mediated inversion, the second crRNA is expressed and triggers the knockout of various immunomodulatory factors, thus mimicking the sequential evolution of cancers in the face of immunotherapeutic pressures.

Targeted amplicon sequencing confirmed efficient mutagenesis of Nf1 (Fig. 4b), followed by mutagenesis of the immunomodulatory factors upon Cre-mediated FlipArray inversion (Fig. 4c) (Supplementary Data 7, 8). Given the pooled nature of these experiments, lower population-level cutting efficiencies are anticipated at the second loci, as only a sixth of the total cell population, on average, is infected with a given FlipArray. The lack of consistent mutagenesis at the crB2m and crCd274 target sites may be intrinsic to the crRNA sequences themselves, a result of inefficient Cre infection/recombination and FlipArray processing, or simply a consequence of biased representation within the cell pool. Of note, we observed high cutting efficiencies at the Jak2 locus despite the pooled nature of the experiment. Since these cells were processed completely in parallel as a minipool, the observation that crJak2 and crLgals9 showed consistent mutagenesis points to intrinsic differences in crRNA targeting efficiencies as the key factor underlying the lack of consistent cutting by crB2m and crCd274. Collectively, these data demonstrate the potential of Cpf1-Flip to facilitate sequential genetic screens—for instance, to model the acquisition of resistance mutations to cancer immunotherapy.

Discussion
We introduced Cpf1-Flip, an inducible sequential mutagenesis system using invertible crRNA FlipArrays. As a proof-of-concept, we demonstrated sequential mutagenesis in both mouse and

(Fig. 3b). In this system, inversion efficiency was 85.42 ± 2.90% by 2 weeks following EFS-Cre (Fig. 3c).

Next, to determine whether the Cpf1-Flip system had enabled sequential mutagenesis at the crDNMT1 and crVEGFA target sites, we performed deep sequencing. As anticipated, uninfected controls did not have significant mutations at either site (Supplementary Data 4, 5; Fig. 3d, e, j, k). Seven days after transduction with EFS-Cpf1; U6-DVF lentivirus, significant indels were found at the crDNMT1 target site but not at the crVEGFA target locus (Fig. 3f, g, j, k). The cells were then infected with EFS-Cre to cause FlipArray inversion, leading to expression of crVEGFA. 21 days after the initial transduction (14 days after EFS-Cre administration), significant indels were observed at both crDNMT1 and crVEGFA target sites (Supplementary Data 6; Fig. 3h–i). In these data, the DNMT1 cutting efficiency appeared to be consistently lower at D21 than at D7. This is likely a consequence of random sampling, as only a subset of the D7 cells were subsequently taken forward for Cre infection. In addition, it is possible that DNMT1 loss affects cell viability, given its crucial role in maintaining DNA methylation. The cutting efficiency at crVEGFA was notably lower compared to crDNMT1. This contrast may be due to lower efficiency of the crRNA itself, as well as inefficiencies in FlipArray expression or subsequent crRNA array processing. Taken together, these results demonstrate that Cpf1-Flip is a flexible tool for sequential mutagenesis based on the Cpf1:crRNA complex, temporally controlled by Cre recombinase.
human cells, while additionally performing pooled sequential mutagenesis in a cancer cell line. These data revealed that the cutting efficiency of the second target loci can be low with certain crRNAs despite successful FlipArray inversion. The most likely explanation for the discordance between FlipArray inversion and subsequent mutagenesis of the second target locus is the differing efficiencies of the crRNAs themselves. This is corroborated by the variance observed across independent crRNAs in the pooled TSG-Immune library (Fig. 4), where consistent cutting efficiencies were observed at the Jak2 and Lgals9 target sites, but not at B2m or Cd27 sites. Moreover, cells with different crRNAs in a pool can undergo random drift or selection, further diverting their relative fractions and thereby indel frequencies. Nevertheless, the FlipArray library can be readout by barcoded PCR of the specific crRNA cassette followed by high-throughput sequencing. Thus, as with all CRISPR screens, pooled screen footprints can be potentially used for sequential and reversible gene activation. Through the use of tethered Cpf1 variants, FlipArrays could also allow for even more complex multi-step gene editing programs. Recombinases and recognition sites in the crRNA array would empower stepwise double knockouts (2 fl–Rev.Complement(crRNA 2)–DR)-3

Methods

FlipArray design and construction. The empty EFS-Cpf1-Puro; U6-FlipArray vector was constructed by modification of the pY109 lentiviral vector (Addgene plasmid #84740)33. After BsmB digestion (FastDigest Esp3I, ThermoScientific), the expression cassette, oligo cloning was then used to invert FlipArrays baseline would crRNAs within an invertible FlipArray at baseline would crRNA spacer: TAAGCATAATGATGATGCCATTTTTTTAAAAAAAAGCAGCTA-

The following crRNA spacer sequences were also used, with analogous oligo designs for cloning into the Cpf1-Flip vector: crDNMT1: CTGATGTCCTACGTCGTGTA crVEGFA: CTAGGAAATTTGAGGCGG crFadu: GTGGCGGCCCTTCAGGCCCA crIdo1: CTACAGGGACTGCCAGAGT crJak2: AACTATCTGGGAAAGAGTAA crLgals9: TGCAGTTACCAACCCGGTGA crPten: TGGACCCAGAGAAAGATTACG crZd74: TAAAGCGTACTACCGAGG

Lenti-Cre vector design and construction. The Lenti-Cre vector was designed to express the Cre recombinase under a constitutive EFS promoter. The plasmid was generated by PCR amplification of Cre and EFS fragments followed by Gibson assembly into a previous lentiviral vector backbone (lentiGuidePuro)34.

Cell culture and genomic DNA extraction. KPD cells were cultured in DMEM supplemented with 10% FBS and 1% penicillin/streptomycin. Experiments were conducted with at least two independent cellular replicates. For genomic DNA extraction, ~200,000 cells were added to each tube, followed by rigorous vortexing for 15 s and centrifuging at 12,000 × g for 10 min. The supernatant containing DNA was then purified using a Qiagen RNeasy Kit following the RNA cleanup protocol. cDNA was generated by reverse transcription with random hexamers. PCR reactions specific for non-inverted and inverted FlipArrays were performed and analyzed simultaneously for each sample. Quantification was done on 2% E-gel using low-range quantitative ladder (ThermoFisher), and was normalized to the first PCR product. Full gel images are shown in Supplementary Fig. 2a–c.

Quantification of inverted FlipArray transcripts. KPD cells were cultured in DMEM supplemented with 10% FBS and 1% penicillin/streptomycin. For RNA extraction, ~200,000 cells were isolated and spun down at 500 × g for 5 min. After a PBS wash, cells were resuspended in 450 μl TRIzol, 100 μl of chloroform was then added to each tube, followed by rigorous vortexing for 15 s and centrifuging at 12,000 × g for 10 min. The supernatant containing RNA was then purified using a Qiagen RNeasy Kit following the RNA cleanup protocol, cDNA was generated by reverse transcription with random hexamers. PCR detection of inverted crRNA FlipArray transcripts was done using the following primers: Inv_FlipArray_F: TCTTTGTGGAAAGGACGAAACCCG Inv_FlipArray_R: AAGCAGCTATTTCCCATGATTCCTTCATATTT RdR: ACGTAGCAGGGAGAGAAGATGTA PCR conditions: 98 °C 2 min, 32 cycles of (98 °C 1 s, 62 °C 5 s, 72 °C 15 s), 72 °C 2 min, 4 °C hold. Following Qiagen PCR purification, 2 μl of the first PCR were used for the second inversion-specific or non-inverted-specific PCR. The following primers were used for detection of non-inverted or inverted FlipArrays: NPF_F: TCTTGTTGGAAGGGAGGAAACCCG NPF_R: TGGCATACGATCTAGCTGCTTGGTAAAAATATTGCGA NPF_R_inv: TAAGCATAATGATGATGCCATTTTTTTAAAAAAAAGCAGCTAG Inv_FlipArray_R: AAGCAGCTATTTCCCATGATTCCTTCATATTT RdR: ACGTAGCAGGGAGAGAAGATGTA PCR conditions: 98 °C 2 min, 32 cycles of (98 °C 1 s, 62 °C 5 s, 72 °C 15 s), 72 °C 2 min, 4 °C hold. PCR reactions specific to non-inverted and inverted FlipArrays were performed and analyzed simultaneously for each sample. Quantification was done on 2% E-gel using low-range quantitative ladder (ThermoFisher), and was normalized to the first PCR product. Full gel images are shown in Supplementary Fig. 2a–c.
Detection of Cpf1 mutagenesis.

The genomic regions flanking the crRNA target sites were amplified from genomic DNA using the following primers:

\[ \text{Lgals9}_R: \text{TAAGCCTGGACTAAGTAAGTGAATGCC} \]
\[ \text{Lgals9}_F: \text{TTTGGCATCTTCACCAAGGTAGATTGT} \]
\[ \text{Jak2}_R: \text{GTAGGTGTTCTCTTTCTGCTTCTCTGCCA} \]
\[ \text{Jak2}_F: \text{AGCTGATCGTTGGTTGCTGCATCAAGGT} \]
\[ \text{B2m}_R: \text{ATTGGGCACAGTGACAGACTTCAATTA} \]
\[ \text{B2m}_F: \text{TGTCAGGTGGAGTCTAGTGGTAGAAAA} \]
\[ \text{IDO1}_R: \text{CCCATGACTTTCCTAAGGAGTGTGAAA} \]
\[ \text{IDO1}_F: \text{TTTGGCATCTTCACCAAGGTAGATTGT} \]
\[ \text{VEGFA}_F: \text{CTCAGCTCCACAAACTTGGTGCC} \]
\[ \text{PTEN}_F: \text{ACTCACCAGTGTTTAACATGCAGGC} \]
\[ \text{Nf1}_R: \text{AACGTGCACCTCCCTTGTCA} \]
\[ \text{Nf1}_F: \text{TTTGGCATCTTCACCAAGGTAGATTGT} \]

The genomic DNA from ~1000 cells was used for the PCR with the NPF and DVF FlipArrays. For the TSG-Immune FlipArray library experiments, genomic DNA from ~6000 cells were used to account for the pooled nature of the PCR. The resultant PCR products were used for Nextera library preparation following manufacturer protocols. Reads were mapped to the mm10 or hg38 genome using BWA-MEM\(^1\), with settings -t 8 -w 200. After identification of indel variants using the pipeline\(^2\)indef function in VarScan v2.3.9, a 1% variant frequency threshold was used for the TSG-Immune experiments due to their pooled nature. All variant calls are detailed in Supplementary Data.

Sample size determination. No specific methods were used to predetermine sample size.

Blinding statement. Investigators were blinded for sequencing data analysis with generic sample IDs, but not blinded for PCR or RT-PCR.

Code availability. Custom scripts will be available to the academic community upon reasonable request.

Data availability. The FlipArray base vector has been submitted to Addgene (ID: 109349) and is available to the academic community. Cell lines, and additional data will be available to the academic community upon reasonable request. The genomic sequencing data sets generated during the current study are available in NCBI SRA under accession SRP136201.

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Author contributions
R.D.C. and S.C. conceived and designed the study. R.D.C. performed experiments and data analysis. H.K. assisted with experiments. R.D.C. and S.C. wrote the manuscript. S.C. supervised the work.

Additional information
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Competing interests: A provisional patent (Composition and Methods for Multiplexed Genome Editing and Screening, US provisional patent no. 62/660,467, filed April 20, 2018) has been filed by R.D.C. and S.C. related to sequential mutagenesis in this study. The remaining author declares no competing interests.

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