Massively parallel genomic perturbations with multi-target CRISPR interrogates Cas9 activity and DNA repair at endogenous sites

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Here we present an approach that combines a clustered regularly interspaced short palindromic repeats (CRISPR) system that simultaneously targets hundreds of epigenetically diverse endogenous genomic sites with high-throughput sequencing to measure Cas9 dynamics and cellular responses at scale. This massive multiplexing of CRISPR is enabled by means of multi-target guide RNAs (mgRNAs), degenerate guide RNAs that direct Cas9 to a pre-determined number of well-mapped sites. mgRNAs uncovered generalizable insights into Cas9 binding and cleavage, revealing rapid post-cleavage Cas9 departure and repair factor loading at protospacer adjacent motif-proximal genomic DNA. Moreover, by bypassing confounding effects from guide RNA sequence, mgRNAs unveiled that Cas9 binding is enhanced at chromatin-accessible regions, and cleavage by bound Cas9 is more efficient near transcribed regions. Combined with light-mediated activation and deactivation of Cas9 activity, mgRNAs further enabled high-throughput study of the cellular response to double-strand breaks with high temporal resolution, revealing the presence, extent (under 2 kb) and kinetics (~1 h) of reversible DNA damage-induced chromatin decompaction. Altogether, this work establishes mgRNAs as a generalizable platform for multiplexing CRISPR and advances our understanding of intracellular Cas9 activity and the DNA damage response at endogenous loci.

Clustered regularly interspaced short palindromic repeats (CRISPR)–Cas nucleases such as Streptococcus pyogenes Cas9 have revolutionized biomedical science through genome manipulation1. For genome editing, Cas9 binds to DNA complementary to its guide RNA (gRNA), induces a double-strand break (DSB), then initiates DNA damage responses (DDRs) that repair and potentially modify the DNA sequence2. Although several studies have shed light on different stages of this process3–9, many aspects of intracellular Cas9 behaviour and ensuing DDR remain incompletely characterized. For instance, how Cas9 departs from genomic DNA after cleavage is unclear10–12, and how genomic context combines with mismatch levels to dictate Cas9 binding and cleavage requires more characterization. The cellular response to Cas9-induced DNA damage also warrants further study10,13, in particular, how damage response factors and chromatin interact with genomic DNA cleaved by Cas9 (refs. 5,14,15).

Better understanding of these CRISPR-associated processes would further mature CRISPR technologies and inspire future tools and applications14,16–18. However, current approaches have been limited to few target sites, in vitro measurements, reporter systems or expressed libraries of gRNAs. Limited target positions preclude exploring heterogeneity at different genomic locations to extract generalizable conclusions19,20, while in vitro measurements fail to capture the complex chromatin context and are not always generalizable to inside cells1,12,21. Reporter systems may not reflect endogenous phenotypes22, and expressed gRNA libraries introduce variability between individual gRNAs, thus obscuring readouts on relative Cas9 activity at different target sites23.

In this Technical Report, we present an approach whereby a single, multi-target gRNA (mgRNA) directs Cas9 to simultaneously target over a hundred endogenous positions genome-wide that are well mapped by high-throughput short-read sequencing. This technique enabled interrogation of Cas9 activity and the ensuing DDNs at endogenous sites at scale. Using mgRNAs, we made discoveries on the dynamics of Cas9 binding and post-cleavage release, the effects of chromatin context on Cas9 activity, and chromatin dynamics during the cycle of DNA damage to repair (Fig. 1a). Our findings establish multi-target CRISPR as a generalizable platform for advancing our understanding of CRISPR-based genome manipulation and cellular DNA damage and repair.

Results
Design, discovery and characterization of mgRNAs. To discover Cas9 gRNA sequences with multiple target positions in the genome, we searched for 20 bp sequences adjacent to a Cas9 protospacer adjacent motif (PAM) in the human genome with up to three mismatches from a 280 bp short interspersed nuclear element (SINE)24. Over 40,000 20 bp sequences were found, each targeting between 2 and over 1,000 putative on-target sites (Fig. 1b). The target sites are located throughout the genome, exhibit balanced representation at different genomic locations to represent multiple epigenetic states21 (Fig. 1c–e).

We then evaluated whether the targeted regions can be uniquely distinguished with high-throughput short-read sequencing. We generated simulated Illumina-style paired-end (PE) 2 × 36 bp reads at all target sites for each gRNA with 5–300 target sites, then determined the number of genome-wide alignments for each read using bowtie2 (ref. 24). For the majority of gRNAs, only a small minority of reads had ambiguous alignments, that is, more than one
Fig. 1 | Initial characterization of mgRNAs. a, Schematic of mgRNA discovery and applications. b, Plot of the thousands of unique target sequences identified in silico, sorted along the x-axis by the number of putative on-target sites in the hg38 human genome (y-axis). A maximum of 1,000 on-target sites were evaluated for each gRNA (even if the total was higher) to maintain efficient computation. c, d, RefSeq (c) and ChromHMM (d) classification of target sequences. In d, only labels with ≥ 1% representation are directly displayed; otherwise they are grouped under ‘Other’. e, For the unique target sequences shown in b, we plotted the mean distance between adjacent putative on-target sites on the same chromosome. The order of sequences along the x-axis is the same as in b, f, Using bowtie2, we determined the proportion of ambiguous reads from simulated PE 2 × 36 bp ChIP-seq reads. g, Same as f, but interpreted as two SE 36 bp ChIP-seq reads. h, i, Same as f and g, respectively, but using 75 bp ChIP-seq reads, instead of 36 bp. j, k, Nucleotide composition at each position in a window of 40 bp (j) or 1 kb (k) around all Cas9 cut sites for a select mgRNA. The x-axis represents the base-pair distance from the Cas9 cut site at x = 0 (fourth nucleotide from PAM). Protospacer, PAM, the ~280 bp SINE and its 3′ A-rich region are annotated. The gRNA sequence is written in white and black font and the PAM in red. l, Mutation rate of ten-target mgRNA. HeLa cells with Dox-inducible Cas9 were transduced with a ten-target mgRNA and grown in presence of Dox. Cells were collected at different timepoints (0, 2, 6 and 10 days). The genomic DNA was extracted, PCR amplified and sequenced at the Cas9 target sites (for details, see Methods). Data points are the average of two biological replicates. m, Day-10 samples from l were analysed to determine the mutational signatures resulting from Cas9 cleavage at each target (8/10) that showed detectable indels. Mutations were classified into five categories: long deletions (longer than 5 bp), short deletions (5 bp or less), SNVs (that is, base change), 1 bp insertions, and insertions longer than 1 bp. Only the mutated protospacers were considered for the analysis. Data points are the average of two biological replicates. n, Reproducibility analysis of m. Each point represents the percentage of a particular mutation type for a given target (akin to the bars in m, but with replicates analysed separately). Data points were fitted to a linear function (dotted line) and their Pearson correlation coefficient was computed, yielding r = 0.99.
**Fig. 2 | Validation of mgRNAs using ChIP-seq.**

**a**, **b**, Average profiles of Cas9 (**a**) and MRE11 (**b**) enrichment in a 2,000 bp window centred at the cut site, across all on-target sites. **c**, Distribution of PE Cas9 and MRE11 ChIP-seq read fragments that span the cut, at two particular cut sites. Coordinates in hg38 are listed in the title of each panel. Numerical value of enrichment is reported as reads per million (RPM). **d**, Proportion of ambiguous reads from measured MRE11 and Cas9 ChIP-seq reads with the three target sequences ‘GG’, ‘CT’ and ‘TA’. Reads at all discovered macs2 peaks were used for analysis, which include both on-target and potential off-target sites. **e**, Histogram of distances between all adjacent macs2 detected Cas9 binding sites (blue; median distance of 265 kb represented by blue dashed line), and between all adjacent on-target sites (green; median distance of 12.2 Mb represented by green dashed line). **f**, Correlation between two biological replicates of MRE11 ChIP-seq. Both axes have units of RPM enrichment in a 2.5 kb window centred at each cut site. **g**, All possible correlations between MRE11, Cas9, yH2AX and 53BP1 ChIP-seq enrichment around all on-target sites (black) and all binding sites (on-target and off-target; grey), between HEK293T cells and iPSCs. **h**, Average profiles of MRE11 ChIP-seq enrichment in a 2,000 bp window centred at each cut site. **i**, Average profiles of 53BP1 (**h**) and yH2AX (**i**) enrichment measured in a specific window centred at all on-target sites. A 2,500 bp window was used for MRE11; 1,500 bp for Cas9; and 20 kb for yH2AX and 53BP1. **j, k**, Average profiles of 53BP1 (**h**) and yH2AX (**i**) enrichment in a 2 Mb window centred at the cut sites. **l**, Plots of relationships between MRE11 and yH2AX (**j**), MRE11 and 53BP1 (**k**), and MRE11 and Cas9 (**l**) at each target site of the three different target sequences (‘CT’, red; ‘TA’, blue; ‘GG’, green). **m, n**, Average profile of MRE11 (**m**) and Cas9 (**n**) enrichment in a 2,000 bp window centred at all cut sites in WTC-11 iPSCs. **o, p**, Plot of MRE11 (**o**) and Cas9 (**p**) ChIP-seq enrichment around all on-target sites (black) and all binding sites (on-target and off-target; grey), between HEK293T cells and iPSCs. Pearson correlation coefficient displayed on graph. **q, r**, Plot of MRE11 (**m**) and Cas9 (**n**) ChIP-seq enrichment around all target sites, between iPSC biological replicates. Source numerical data are available in source data.
Distal
Other fragments
Fragments that abut DSB site
Fragments that span DSB site

a) Conceptual schematic of ChIP-seq analysis for PE reads, distinguishing between read fragments that abut or span the DSB site. Reads abut the DSB site if they fall within 5 bp from the cut site. Distal reads abut the DSB site if they fall within 5 bp from the cut site.

b) Number of pE reads (RPm) that either span or abut each target site for MRE11 (c) and Cas9 (e) mgRNA on-target site, RPM on the PAM-proximal versus PAM-distal side for MRE11 (orange) versus Cas9 (green). ChIP-seq 3 h after Cas9 delivery. The numbers indicate linear regression slopes. MRE11 and Cas9 slopes from d-f are inversely correlated. For each target site, we plotted the MRE11 PAM-proximal bias versus Cas9 PAM-distal bias. Correlation was determined using Pearson correlation with its p value.

For deletions >5 bp or <5 bp, on the x-axis).

For deletions >5 bp or <5 bp, on the x-axis).

For cells exposed to 10 days of Cas9 with
j) Violin plot of the number of reads, for each Cas9 target site, categorized by the four read types (dist +4, dist −4, prox +4 and prox −4) described above. Comparison between dist +4 and sum of prox +4 and prox −4 using two-sided unadjusted Student's t-test. **P < 0.0001. Values are: 3 × 10^-10, 3 × 10^-15 and 3 × 10^-16 for CT, TA and GG, respectively. p) Two-sided unadjusted Student's t-test of significance for the number of PAM-proximal reads (prox +4 vs. prox −4) between different gRNA sequences. NS, not significant, ***P < 0.0001. Values from left to right are: 0.35, 2.6 × 10^-22 and 6.6 × 10^-19. q) Schematic of the ‘CT’, ‘TA’ and ‘GG’ sequences, which differ only in the most PAM-proximal two nucleotides: prox 4 prox −4. A fourth, hypothetical species is included for completeness: prox +4 dist −4 prox +4 nucleotide; prox −4: immediately PAM-proximal lacking +4 nucleotide. A fourth, hypothetical species is included for completeness: dist −4: immediately PAM-distal lacking +4 nucleotide. The +4 nucleotide is highlighted in cyan. m-o, Violin plot of the number of reads, for each Cas9 target site, categorized by the four read types (dist +4, dist −4, prox +4 and prox −4) described above. Comparison between dist +4 and sum of prox +4 and prox −4 using two-sided unadjusted Student's t-test. **P < 0.0001. Values are: 3 × 10^-10, 3 × 10^-15 and 3 × 10^-16 for CT, TA and GG, respectively. p) Two-sided unadjusted Student's t-test of significance for the number of PAM-proximal reads (prox +4 vs. prox −4) between different gRNA sequences. NS, not significant, ***P < 0.0001. Values from left to right are: 0.35, 2.6 × 10^-22 and 6.6 × 10^-19. q) Schematic of the ‘CT’, ‘TA’ and ‘GG’ sequences, which differ only in the most PAM-proximal two nucleotides (highlighted in purple). The ‘GG’ target sequence has an extra PAM. ‘NGG’ PAM(s) are labelled in red. The blunt-end cleavage possibility is displayed with red triangles annotating the cleavage position. Source numerical data are available in source data.
alignment with the same ‘best’ bowtie2 alignment score (Fig. 1f). As expected, treating the PE 2 × 36 bp reads as single-end (SE) reads increased the percentage of reads with ambiguous alignments (Fig. 1g), whereas increasing the number of sequenced base pairs to 75 at each end (that is, PE 2 × 75 bp) reduced this percentage (Fig. 1h, i). We then aligned the sequence around each expected on-target site for a gRNA with under 1% ambiguous alignments. The nucleotide composition at each position in a 40 bp window confirmed the expected Cas9 protospacer (Fig. 1j). Expanding to a 1 kb window confirmed features of the Alu SINE, such as its 280 bp approximate length and A-rich 3′ end (Fig. 1k). The sequences beyond 150–200 bp from the cut sites were evenly distributed between the four nucleotides and probably correspond to regions that can be uniquely mapped by sequencing. PE sequencing reads can therefore be uniquely mapped given the sequence diversity even within the short repetitive element and the high probability of at least one DNA end being positioned outside the element. We replicated the same analysis using a mouse and a zebrafish genome and different SINEs (Extended Data Fig. 1a–k). Together, our computational pipeline robustly identified diverse candidate gRNAs across different species.

Experimental validation of gRNAs. We validated the activity of gRNAs by measuring genome editing outcomes (insertions and deletions, or indels) at gRNA targeted sites. Cas9/gRNA with ten predicted target sites active in HeLa cells over 10 days revealed robust indel generation at eight out of the ten sites (Fig. 1l and Extended Data Fig. 1l). The mutation identity was predominantly one-nucleotide insertions, consistent with the repair profiles of Cas9-generated DSBs (Fig. 1m). Similar results were obtained with a different gRNA (Extended Data Fig. 1m,n), and mutation distributions showed high reproducibility between biological replicates (Fig. 1n and Extended Data Fig. 1o). Together, these results demonstrate efficient intracellular activity with gRNAs. To interrogate Cas9 binding and recruitment of DNA repair factors in a high-throughput manner, we tested three gRNAs (‘CT’, ‘GG’ and ‘TA’) with 145, 126 and 117 on-target sites, respectively. We electroporated Cas9 protein pre-assembled with gRNA into HEK293T cells, followed 3 h later by chromatin immunoprecipitation with sequencing (ChIP–seq) for Cas9 and an early DDR protein, MRE11 (refs. 4,5,13,28). ChIP–seq profiles averaged across all on-target sites revealed high enrichment with shapes consistent with previous literature (Fig. 2a–c). Cas9 on- and off-target sites were called using MACS2 software (Extended Data Fig. 1p–r), and showed less than 0.3% of sequencing reads with ambiguous alignments (Fig. 2d), verifying that ChIP–seq accurately quantified enrichment at sites targeted by these gRNAs. Median distances between adjacent Cas9 binding sites and adjacent on-target sites were both large, at 270 kb and 13 Mb, respectively (Fig. 2e). MRE11 enrichment was highly correlated between biological replicates (Fig. 2f) and with other DNA repair markers such as 53BP1 and phosphorylated H2AX (γH2AX) (Fig. 2g–k). In contrast, correlations between Cas9 and DNA repair factors were weaker and dependent on gRNA sequence (Fig. 2l). MRE11 and Cas9 ChIP–seq after gRNA delivery was also performed in pluripotent stem cells (iPSCs) (Fig. 2m,n). ChIP–seq enrichments at target sites were only moderately correlated between iPSCs and HEK293T cells (Fig. 2o,p) despite high correlation between biological replicates (Fig. 2q,r). Altogether, these results demonstrate multiplexed Cas9 activity and robust ChIP–seq readout for Cas9 and DNA repair factors at endogenous sites targeted by gRNAs.

Cas9 binding and cleavage mechanics at endogenous loci. Characterizing how Cas9 interacts with genomic DNA is important to better understand Cas9 genome editing10,28. For example, how Cas9 departs from genomic DNA after cleavage is unclear; RNA polymerase4 and histone chaperone FACT5,29 have both been proposed to evict Cas9, but direct evidence inside cells is lacking. To dissect these dynamics in a highly multiplexed fashion while controlling for the target sequence, we exposed HEK293T cells to ‘GG’, ‘CT’ or ‘TA’ gRNAs for 3 h, and categorized the resulting Cas9 and MRE11 ChIP–seq reads as either spanning or abutting the cut site, corresponding to protein-associated DNA fragments that are either intact or cleaved by Cas9, respectively (Fig. 3a). MRE11 ChIP–seq reads predominantly abutted the cut sites (Fig. 3b), consistent with MRE11 loading on cleaved DNA10,28, whereas Cas9 ChIP–seq reads predominantly spanned the cut sites (Fig. 3c), consistent with Cas9 residing on the target before cleavage and departing quickly thereafter. Of the reads that abut each cut site, MRE11 exhibited enrichment bias for the PAM-proximal side of the cut for most target sites, while Cas9 showed bias for the PAM-distal side (Fig. 3d–f and Extended Data Fig. 1p–r). The extent of PAM-proximal/PAM-distal bias was inversely correlated between MRE11 and Cas9 though not all target sites exhibited this bias (Fig. 3g–i and Extended Data Fig. 1s). These results suggest stable Cas9 binding before cleavage, possibly to check for sequence complementarity, followed by cleavage and rapid release of DNA preferentially from the PAM-proximal side, facilitating MRE11 loading. Consistent with this model, sequencing of indel products showed preferential short deletions at the PAM-proximal (MRE11-resident) side (Fig. 3 and Extended Data Fig. 1t). Preferential Cas9 dissociation from the PAM-proximal side was observed previously, but only for a single target sequence and in vitro40. Our results validate this observation in cells and further suggest that Cas9 binding to a cleaved DNA terminus can obfuscate it from MRE11 and the cellular DDR.
To further characterize post-cleavage Cas9 mechanics, we modelled Cas9 ChIP–seq read species derived from DNA fragments bound to Cas9 after either staggered or blunt cleavage. From staggered cleavage, DNA end repair during ChIP–seq library preparation fills in the 3’ end, resulting in presence of the fourth nucleotide (from PAM) at both sides of the cut (Fig. 3k). We refer to these ChIP–seq reads on PAM-proximal and PAM-distal sides as ‘prox + 4’ and ‘dist + 4’, respectively. In contrast, from blunt-end cleavage, only the PAM-distal read contains the fourth nucleotide, that is, ‘dist + 4’, whereas the PAM-proximal read does not, resulting in a ‘prox – 4’ read species (Fig. 3l). ‘dist + 4’ was significantly more enriched than the sum of ‘prox + 4’ and ‘prox – 4’ (P < 1 × 10^{-15}, Student’s t-test), recapitulating clear PAM-distal binding bias (Fig. 3m–o). These results suggest that the 16–17 bp of gRNA to genomic DNA base-pairing interactions at the PAM-distal side of the cut are stronger than the 3–4 bp of base pairing and PAM–Cas9 interactions at the PAM-proximal side. Interestingly, Cas9 with the ‘GG’ gRNA exhibited significantly stronger association
with the PAM-proximal side compared with the other two gRNAs \((P < 1 \times 10^{-18}, \text{Student's } t\)-test\) (Fig. 3p), which we speculate may be due to the additional 'NGG' PAM sequence in the first three nucleotides of the protospacer (Fig. 3q).

**Linking Cas9 binding and DNA repair to local epigenetic states.** Genome editing efficiencies are difficult to predict but are probably influenced by both sequence and epigenetic factors. Epigenetic influences have been challenging to decipher owing to confounding effects of gRNA sequence; mgRNAs are uniquely suited for this task because a common gRNA sequence targets different epigenetic contexts. To characterize Cas9 binding alone, we measured occupancy of (cleavage-deficient) dCas9 using ChIP–seq after mgRNA/dCas9 delivery. For the 'GG' mgRNA, we detected 5,236 dCas9 binding sites (Fig. 4a), a number of off-target sites comparable to single-targeting gRNAs. To evaluate Cas9-mediated DNA damage, we measured occupancy of MRE11 after delivery of (cleavage-competent) Cas9. MRE11 was only enriched at sites with two or fewer mismatches whereas some sites exhibited clear dCas9 binding for up to over eight mismatches, and both

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**Graphical Abstract**

- **a** Graph showing RPM versus base pairs from cut site.
- **b** Graph showing RPM enrichment versus base pairs (thousands) from cut site.
- **c** Graph showing RPM enrichment versus distance from cut site (kb).
- **d** Graph comparing MRE11 ChIP–seq and ATAC–seq for different chromosomes.
- **e** Graph showing RPM versus base pairs from cut site for MRE11 ChIP–seq FWHM.
- **f** Graph showing RPM versus base pairs from cut site for ATAC–seq FWHM.
- **g** Graph showing correlation coefficient versus distance from cut site (kb).
- **h** Graph showing spanning enrichment versus distance from cut site (kb).

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**Figure Legends**

- **Fig. 1** Linking Cas9 binding and DNA repair to local epigenetic states.
- **Fig. 2** Genome editing efficiencies are difficult to predict but are probably influenced by both sequence and epigenetic factors.
- **Fig. 3** Epigenetic influences have been challenging to decipher owing to confounding effects of gRNA sequence; mgRNAs are uniquely suited for this task because a common gRNA sequence targets different epigenetic contexts.
- **Fig. 4** To characterize Cas9 binding alone, we measured occupancy of (cleavage-deficient) dCas9 using ChIP–seq after mgRNA/dCas9 delivery. For the 'GG' mgRNA, we detected 5,236 dCas9 binding sites (Fig. 4a), a number of off-target sites comparable to single-targeting gRNAs. To evaluate Cas9-mediated DNA damage, we measured occupancy of MRE11 after delivery of (cleavage-competent) Cas9. MRE11 was only enriched at sites with two or fewer mismatches whereas some sites exhibited clear dCas9 binding for up to over eight mismatches, and both

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**Table**

| Base pairs from cut site | RPM |
|--------------------------|-----|
| -1,500 -1,000 -500 0 500 1,000 |    |

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**Diagrams**

- **Diagram a** Shows RPM versus base pairs from cut site.
- **Diagram b** Shows RPM enrichment versus base pairs (thousands) from cut site.
- **Diagram c** Shows RPM enrichment versus distance from cut site (kb).
- **Diagram d** Compares MRE11 ChIP–seq and ATAC–seq for different chromosomes.
- **Diagram e** Shows RPM versus base pairs from cut site for MRE11 ChIP–seq FWHM.
- **Diagram f** Shows RPM versus base pairs from cut site for ATAC–seq FWHM.
- **Diagram g** Shows correlation coefficient versus distance from cut site (kb).
- **Diagram h** Shows spanning enrichment versus distance from cut site (kb).
enrichments were higher if the mismatch resided solely in the PAM-distal region (≥12th position, counting from PAM) (Fig. 4b,c and Extended Data Fig. 2a), consistent with known properties of Cas9 binding and cleavage\(^{5,11}\). Interestingly, there was high heterogeneity in both dCas9 and MRE11 enrichment even between identical on-target sequences (Fig. 4b,c), probably stemming from epigenetic factors.

To infer the epigenetic state, we obtained ten publicly available genome-wide epigenetic maps from the same cell line\(^{36}\) and determined their enrichments in specified windows centred around each Cas9 target site (Fig. 4d). dCas9 enrichment was most strongly correlated with markers of DNA accessibility, as measured by assay for transposase-accessible chromatin using sequencing (ATAC-seq) and DNase I-hypersensitive site sequencing (DNase-seq), consistent with previous reports\(^{34,37,38}\). In contrast, MRE11 recruitment was correlated with additional chromatin features besides accessibility (Fig. 4e), suggesting that additional epigenetic factors are at play beyond Cas9 binding. To characterize the MRE11 damage response independent of Cas9 binding, we normalized MRE11 signal by dCas9 signal, which yielded the strongest correlation with gene bodies (H3K27me3 and RNA sequencing (RNA-seq)), promoters (H3K4me3 and RNA polymerase II) and enhancers (H3K27ac) (Fig. 4f). This suggests either higher Cas9 cleavage efficiencies or more efficient MRE11 recruitment at these regions, which we can distinguish by directly measuring DSBR levels genome-wide using breaks labelling in situ and sequencing (BLISS)\(^{39}\). BLISS enrichment was highly correlated with MRE11 (\(r = 0.7\)) (Extended Data Fig. 2bc), and the pattern of epigenetic correlation for dCas9-normalized BLISS enrichment (unrepaired DSBRs given the same amount of Cas9 binding) mirrored dCas9-normalized MRE11 enrichment (Fig. 4f). These results suggest that identical Cas9 on-target sites bound by the Cas9–gRNA complex are cleaved at different rates. In particular, regions near gene bodies, promoters and enhancers exhibit intrinsically higher cleavage activity by a bound Cas9. Together, improved Cas9 binding at accessible regions, followed by increased Cas9-mediated DNA damage near enhancers, promoters and gene bodies, provides an explanation for previous studies using sgRNAs that report higher genome editing efficiencies at these exact regions (Fig. 4g)\(^{11,34,35}\).

The biophysical mechanism for improved Cas9 cleavage near transcribed regions requires further investigation. One possible explanation is DNA supercoiling; transcribed regions are known to be negatively supercoiled\(^{40,41}\), and single-molecule biophysical studies showed that Cas9 cleaves more efficiently on DNA negatively supercoiled at physiologically relevant levels\(^{42}\). Other potential mechanisms include DNA-binding proteins such as RNA polymerase\(^{43}\) and the histone chaperone FACT\(^{44}\) influencing Cas9 residence on gDNA.

**Prediction of genome editing processes using machine learning.** To further explore the determinants of Cas9 binding and DNA damage induction, we trained random forest machine learning models to predict both dCas9 and MRE11 enrichment at all binding locations. From solely mismatch information, dCas9 and MRE11 enrichment at 3 h could be adequately predicted for an independent test dataset with \(r = 0.78\) and 0.64, respectively (Fig. 4h,i). Using solely epigenetic information led to comparable levels of performance with \(r = 0.75\) for dCas9 and 0.59 for MRE11 (Fig. 4j,k). However, using both mismatch and epigenetic information greatly improved prediction, resulting in \(r = 0.86\) for dCas9 and 0.83 for MRE11 (Fig. 4l,m). Comparable levels of predictive power were also achieved for the 30 min timeframe (Extended Data Fig. 2d–i). These results highlight the importance of local epigenetic state in modulating Cas9 activity and provide further evidence.

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**Fig. 5 | Chromatin accessibility change after DNA damage.** a. Averaged background-subtracted ATAC-seq profiles across all on-target sites for cells without Cas9 (‘neg’, dark blue); or 3 h after Cas9/‘GG’–gRNA (‘Cas9’, red), dCas9/‘GG’ (‘dCas9’, medium blue) or D10A Cas9 nickase/‘GG’ delivery (‘nCas9’, light blue). ‘Background-subtracted’ enrichment was obtained by subtracting the number of reads at each position for Cas9-negative cells from Cas9-exposed signals at four representative cut sites. The genomic coordinate (hg38) of the cut sites are displayed on top of each panel. b. Measurement of FWHM for MRE11 ChIP–seq enrichment (523 bp) versus MRE11 enrichment (722 bp) averaged across all on-target cut sites. c. Top: average background-subtracted ATAC–seq enrichment (in RPM) at each target site for samples from a. Comparison using two-sided unadjusted Student’s t-test. ****\(P < 0.0001\). **b** Subtraction of the ‘Cas9’ sample by ‘neg’ sample from excess ATAC–seq enrichment due to Cas9-mediated DNA damage. \(n = 2\) biologically independent experiments. d. Histogram of ATAC-seq DNA lengths for cells without Cas9 (‘neg’, blue), or 3 h after Cas9/‘GG’–gRNA delivery (‘Cas9’, red). j. Subtraction of the ‘Cas9’ sample by ‘neg’ sample from j. Fitting (black dotted curve) was performed using an exponential decay model. k, i. Same as a and b, for cells without Cas9 (‘neg’, dark blue) or 3 h after Cas9/‘GG’–gRNA delivery (‘Cas9’, red), but only measuring the subset of ATAC–seq reads that span the cut site. Comparison using two-sided unadjusted Student’s t-test. ****\(P < 0.0001\). P value is 1.44 \times 10^{-6}. Source numerical data are available in source data.
that combining epigenetic with mismatch information improves the prediction of genome editing activity21,43.

**Increase in chromatin accessibility at Cas9-induced DSBs.** It has been proposed that local chromatin decompaction occurs after DNA damage to facilitate repair, but direct evidence has not been observed at single Cas9 DSBs44,45. To measure chromatin accessibility changes after DNA damage, we performed ATAC–seq46 with and without exposure to Cas9/mgRNA. Averaged background-subtracted ATAC–seq enrichment centred at Cas9 target sites exhibited locally increased chromatin accessibility at 10 min compared to 0 min at DSB sites (Figure 1). This increase in accessibility was specific to target sites and not seen at random non-target sites (Figure S5).

**Mouse ChIP and ATAC-seq.** To confirm the accessibility changes measured by ATAC–seq, we performed BLISS MRE11 ChIP and ATAC–seq with and without exposure to Cas9/mgRNA. MRE11 recruitment was increased at 10 min compared to 0 min at target sites (Figure 2). This increase in MRE11 recruitment was specific to target sites and not seen at random non-target sites (Figure S5).

**Other repair factors.** To further validate the accessibility changes measured by ATAC–seq, we performed BLISS MRE11 ChIP and ATAC–seq with and without exposure to Cas9/mgRNA. MRE11 recruitment was increased at 10 min compared to 0 min at target sites (Figure 2). This increase in MRE11 recruitment was specific to target sites and not seen at random non-target sites (Figure S5).

**Figure 1:** Diagram showing the steps of the repair process. 1. Cas9 binds target DNA, 2. Cas9 cleaves DNA, 3. MRE11 binds PAM-proximal end 10 min after DSB, 4. Chromatin opens around DSB 30 min after DSB, 5. DSB is repaired, 6. Chromatin is restored 1 h after repair completion, 7. Other repair factors.
We further uncovered a subpopulation of ATAC–seq reads spans 1442sibility that can persist after DNA ligation, which potentially nucleosome-depleted, kilobase-scale region of increased acces-
sability after ligation. In conclusion, Cas9 cleavage induces a localized,
gests that chromatin recompaction does not occur immediately
respond to post-cleavage DNA that has undergone ligation and sug-
ping the target sites that significantly increased after Cas9 delivery
2 kb accessible region from Fig. 5c lost up to ten nucleosomes47,48.
with distances between adjacent Tn5 transposition events that are
assumed to be a Poisson point process (Fig. 5j). Assuming nucleo-
distribution of the excess reads lacked the nucleosomal footprint-
Cells exposed to Cas9 had excess ATAC–seq reads; the length
corresponded to nucleosome occupancy footprinting46 (Fig. 5i).
Next, we inferred the lengths of all PE ATAC–seq reads within
1.5 kb from expected target sites. For cells without Cas9, the distribu-
tion of sequencing read lengths showed a local maximum that corresponded to nucleosome occupancy footprinting46 (Fig. 5i).
Cells exposed to Cas9 had excess ATAC–seq reads; the length distribution of the excess reads lacked the nucleosomal footprinting
signature and was well fit by an exponential decay, consistent with distances between adjacent Tn5 transposition events that are
assumed to be a Poisson point process (Fig. 5i). Assuming nucleo-
some spacing length of around 200 bp, this implies that the up to
2 kb accessible region from Fig. 5c lost up to ten nucleosomes47,48.
We further uncovered a subpopulation of ATAC–seq reads span-
ing the target sites that significantly increased after Cas9 delivery
\( P = 1.44 \times 10^{-18}, \) Student’s t-test (Fig. 5k,l), which must corre-
spond to post-cleavage DNA that has undergone ligation and sug-
gets that chromatin recombination does not occur immediately
after ligation. In conclusion, Cas9 cleavage induces a localized,
nucleosome-depleted, kilobase-scale region of increased acces-
sibility that can persist after DNA ligation, which potentially facilitates the binding of DNA damage-associated proteins such as repair factors, cohesin and transcription factors to promote successful repair49–51.

Chromatin accessibility dynamics in DSB repair. The temporal sequence of events after Cas9 cleavage has not been well charac-
terized but can be explored using the very fast light-activatable CRISPR (vfCRISPR) based on a photocaged gRNA (cgRNA)51. We delivered Cas9 with the multi-target ‘GG’ cgRNA to HEK293T cells, waited 12 h for stable Cas9 binding, then light-activated Cas9 and performed time-resolved BLISS, MRE11 ChIP–seq and ATAC–seq. DSBs and MRE11 damage responses were undetectable before light activation, confirming that Cas9 is inactive without light exposure (Fig. 6a,b). As early as 10 min after activation, BLISS exhibited the strongest relative enrichment increase followed by MRE11 ChIP–seq signal (Fig. 6a,b), consistent with initial DSB induction followed by repair protein recruitment. ATAC–seq enrichment increased by 30 min after Cas9 activation but not 10 min (Fig. 6c,d and Extended Data Fig. 2j), suggesting that DSB-induced increase in accessibility occurs downstream of initial repair protein recruitment.

After repair of DNA damage, the duration of accessibility increase remains unknown. However, without an effective method for CRISPR deactivation, intracellular Cas9 will repeatedly cleave repaired loci and preclude measurements of chromatin restora-
tion52. We therefore employed a light-deactivatable Cas9 based on a photocleavable gRNA (pcRNA) to synchronize Cas9 deactivation, facilitating chromatin profiling through repair completion53. We delivered Cas9 with multi-target ‘GG’ pcRNA to HEK293T cells, deactivated Cas9 after 2 h and performed time-resolved MRE11 ChIP–seq and ATAC–seq. After Cas9 deactivation, MRE11 enrichment rapidly declined across all target sites with 75% reduction.
in enrichment within the first 15 min (Fig. 6e), which probably corresponds to completion of DNA repair\(^\text{13}\). In contrast, the level of chromatin accessibility increase persisted for the first 15 min before declining (Fig. 6f–h), consistent with our previous results in Fig. 5k,l and suggesting that accessibility reversal is delayed compared with MRE11 departure. There was no detectable correlation between MRE11 departure and the tested epigenetic markers (Fig. 6i). Inhibition of DNA–PKcs using KU-60648 prevented MRE11 departure\(^\text{2}\), suggesting that the repair events are dependent on non-homologous end-joining (Fig. 6j–k)\(^\text{4}\).

Our findings on Cas9 activity and DDR are summarized in Fig. 6l. After binding and cleavage of target DNA, Cas9 quickly releases the DNA preferentially from the PAM-proximal side, enabling binding of MRE11 to this DNA end within 10 min. Within 30 min of DSB, chromatin undergoes decompaction whereby nucleosomes ~1 kb from the cut site are evicted, potentially facilitating recruitment of additional DNA repair factors. Once the lesion has been repaired, the nucleosomes are repositioned around the cut site, restoring the chromatin accessibility landscape to pre-cleavage levels.

**Discussion**

We report the discovery and applications of multiplexed CRISPR using mgRNAs. We identified tens of thousands of mgRNAs that each target 2 to over 1,000 positions across multiple genomes, providing an extensive resource for rapid adoption. We then combined mgRNAs with high-throughput sequencing readouts to provide the most comprehensive study thus far of Cas9 genome editing and ensuing DDRs at endogenous loci (Supplementary Table 1). The large number and diversity of target sites enables generalizable observations such as the destabilizing impact of even one PAM-distal mismatch on Cas9 binding and better cleavage by bound Cas9 near transcribed regions. Aggregating data across multiple target sites boosts readout signal, allowing us to use ATAC–seq reads across all target sites to measure local nucleosome depletion after Cas9 DNA damage. Furthermore, compatibility with very fast CRISPR activation and deactivation\(^\text{15,16}\) allowed quantification of the dynamics of chromatin accessibility change during and after DNA repair with high temporal resolution. Cas9 with mgRNAs also exhibits advantages over ‘multi-target’ meganucleases\(^\text{17}\) including programmable target positioning, precise time control using CRISPR activation and deactivation, facile delivery without need to generate a stable cell line, and relevance to CRISPR genome editing. Finally, the ability to read mutational outcomes of mgRNA paves the way towards its use as a genetic barcoding tool. Supporting this claim, the indels at eight target sites generated by the ten target mgRNA in HeLa cells demonstrated high barcoding diversity as measured by Shannon entropy\(^\text{18}\) (Fig. 7a).

Our study is not without limitations. First, the mgRNA model system may not translate to native DSBS or single-targeting Cas9. However, this is unlikely given that most of our findings are corroborated with existing literature. Second, our assumption that every mgRNA target site is independent could be challenged if Cas9 binding/cleavage events physically influence measurements at adjacent target sites. However, the median distance between adjacent binding sites (265 kb) and adjacent on-target sites (13.2 Mb) (Fig. 2c) is orders of magnitude greater than the ~2 kb window used for the bulk of analysis, so the effect of nearby off-target Cas9 activity is probably minor. Third, bulk sequencing cannot deconvolute heterogeneity between individual cells, which may be overcome by combining mgRNAs with single-cell imaging\(^\text{19}\) or sequencing readouts\(^\text{20}\). Finally, mgRNA can generate high numbers of simultaneous DSBS in each cell, averaging under 50 per cell for a 126-targeting mgRNA based on the number of 53BP1 foci in immunofluorescence microscopy (Fig. 7b,c). A single DSB delayed cell division, consistent with a previous report\(^\text{21}\), and 50 DSBS blocked cell division (Fig. 7d). We believe the high DSB count is unlikely to influence our results because all experiments were conducted within 3 h of Cas9 delivery during which no altered cellular phenotypes were observed (Fig. 7e and Extended Data Fig. 2j), and relative Cas9 kinetics between different target sites are probably unaffected by the high mutation load.

In conclusion, we developed mgRNAs as an approach to multiplex CRISPR–Cas9 at endogenous sites. Using mgRNAs, we revealed insights on Cas9 target recognition and cleavage activity, and determined the dynamics of chromatin accessibility during repair of Cas9-induced DSBS. We envision that mgRNAs will be a powerful tool to further advance our understanding of CRISPR technologies and DNA repair processes.

**Online content**

Any methods, additional references, Nature Research reporting summaries, source data, extended data, supplementary information, acknowledgements, peer review information; details of author contributions and competing interests; and statements of data and code availability are available at https://doi.org/10.1038/s41556-022-00975-z.

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Methods

**SpCas9 purification.** SpCas9 purification was done using BLS1-CodonPlus (DE3)-RIL competent cells (Agilent Technologies 230245) that were transformed with CaSp9 plasmid (Addgene, #67881). Bacteria were grown in 1 L of LB medium, induced with isopropyl-β-D-thiogalactoside overnight and then lysed. The supernatant was clarified and then purified using Ni-NTA beads. A detailed description can be found in ref. 56.

**Cell culture.** HEK293T cells (ATCC CRL-3216) and HeLa (ATCC CCL-2) cells were cultured at 37 °C under 5% CO2, in Dulbecco's Modified Eagle's medium (DMEM, Corning) supplemented with 10% FBS (Clontech), 100 units/mL penicillin and 100 μg mL−1 streptomycin (DMEM complete). Cells were tested every month for mycoplasma.

A human iPSC, WTC11 cell line was used for all iPS experiments in this study. We followed the guidelines of Johns Hopkins Medical Institute for the use of this human iPSC line. Briefly, frozen WTC11 cells were first thawed at 37 °C water bath and washed in Essential 8 Medium (E8; Thermo Fisher Scientific, #A1517001) by centrifugation. After resuspension, WTC cells were plated onto a 6 cm cell culture dish pre-coated with human embryonic cell-qualified Matrigel (1:100 dilution, Corning, #354277). Plate coating should be performed for at least 2 h. Subsequently, 10 μM ROCK inhibitor (Y27632; STEMCELL, #72308) was supplemented into the E8 medium to promote cell growth and survival. For subculture, WTC11 cells were dissociated from the plate using accutase (Sigma, #A6964) and passed every 2 days. WTC11 cells were maintained in an incubator at 37 °C with 5% CO2.

**Electroporation of Cas9 ribonucleoprotein.** A Cas9 mRNA ribonucleoprotein was assembled and electroporated into HEK293T or WTC-11 iPS cells using 4D-Nucleofector Kits (Lonza, SF Cell Line kit for HEK293 and P3 Primary Cell kit for WTC11) following the manufacturer's instruction. Oligos used for trans-activating CRISPR RNA (tracrRNA) and CRISPR RNAs (crRNAs) are presented in Supplementary Table 3. A detailed protocol can be found in ref. 56.

**Chromatin immunoprecipitation sequencing.** The ChIP protocol was adapted from previous literature39. Oligonucleotide sequences for library preparation are in Supplementary Table 3. A detailed protocol can be found in ref. 56. Briefly, protein A beads were washed twice using BSA buffer and incubated with the antibody for 1–3 h with rotation. Bead–antibody mixtures were washed twice with BSA buffer right before ChIP. Cells were collected and fixed with formaldehyde (1% final) for 1–3 h with rotation. Bead–antibody mixtures were washed twice with BSA buffer right before ChIP. Cells were collected and fixed with formaldehyde (1% final) at room temperature. The reaction was quenched using glycine (130 mM final). Cells were then lysed sequentially using three different buffers, sonicated and spun down. The supernatant was collected, and the bead–antibody mixture was added. The ChIP reaction incubated overnight. Bead mixtures were then washed on a magnet seven times, resuspended in reverse crosslink buffer and incubated at 65 °C for 6+ hours. After proteinase K and RNase A treatments, the DNA was column purified. To prepare ChIP-seq libraries, we performed end repair/adapter ligation and PCR using PE_i5 and PE_i7XX primer pairs. Final DNA was purified using AMPure beads, quantified via Qubit, pooled and sequenced on a NextSeq 500 (Illumina).

**Genome-wide DSB detection with BLISS.** The BLISS protocol was adapted from previous literature40. All oligonucleotide sequences are provided in Supplementary Table 4. A detailed protocol can be found in ref. 56. In short, BLISS adapters were annealed and phosphorylated. RA3 oligonucleotides (Integrated DNA Technologies, 400,000 cells were seeded into a 24-well plate for each reaction, washed once with PBS, fixed with 4% paraformaldehyde for 10 min, then washed three times with PBS. Cells were then subjected to a first round of lysis, followed by a PBS wash, a second round of lysis and two PBS washes. Cells were then washed twice with CutSmart Buffer (NEB), and subjected to DNA end-blunting reaction. Cells were then added with CutSmart Buffer followed by adenylation of DNA ends. Cells were washed twice with CutSmart Buffer and with T4 Ligase Buffer, followed by in situ adapter ligation. Samples were then washed four times with high-salt buffer to remove unligated adapters. DNA was extracted by adding extraction buffer and proteinase K, incubating at 55 °C overnight and column purifying DNA the day after. DNA was then sonicated, in vitro transcribed and purified. RA3 oligonucleotide adapter was ligated to the purified RNA, and the product was purified. Samples were reverse transcribed and PCR amplified, and the final DNA was purified using AMPure beads. Samples were pooled, quantified with QuBit, Bioanalyzer and qPCR, then sequenced on a NextSeq 500 using high-output paired sequencing, with 64 bp for read 1 and 36 bp for read 2. Only the subset of reads with the perfectly matching 13 bp constant adapter region (CCGCATACGCGC) in read 1 was used for subsequent analysis.

**Measurements of mutations at mRNA targets.** A PiggyBac system was used to transpose HeLa cells with a vector carrying Cas9 under the control of a Tet-Off inducible promoter. After transforming the ligated product into the gel purification plasmid and following the manufacturer’s instructions. The following day, individual colonies were selected and grown in selection media; plasmids were purified the next day using QIAprep Kit (Qiagen, #27016). Correct insertion of the mRNA was verified via Sanger sequencing. For lentivirus production, Lenti-X-293T cells (Addgene, #16380) were grown in 60 cm dishes up to ~70% confluency. Then, 5.25 μg of transfer plasmid was mixed with 0.75 μg of PMD2.G (Addgene, #12259) and 1 μg of psPAX2 (Addgene, #12260), and with 21 μl of TransIT-Lenti (Mirus, #6603). The mixture was incubated for ~15 min and added dropwise to the cells. The viral supernatant was collected at 36h, 48h and 60h, and filtered and concentrated using Lenti-X Concentrator (takarabio, #631232), according to the manufacturer’s instructions. Doxycycline (Dox)-inducible Cas9 monoclonal cells were grown to ~60 % confluency in six-well plates. Cells were exposed to virus carrying mRNA (~0.3 – 0.5 multiplicity of infection) and 8 μg mL−1 polybrene for 24h. Two days after infection, cells were exposed to 100 μg mL−1 hygromycin and kept under such selection conditions for all subsequent experiments. Half of the cells confirmed successful plasmid integration at the estimated multiplicity of infection. An initial set of stably transduced cells were collected before Dox addition as timepoint zero. Cells were then grown in 24-well plates under exposure to 2 μg mL−1 of Dox. At different timepoints after induction, a number of cells were collected during passaging and their gDNA was extracted. For the ten-target mRNA, a no-Dox control experiment was performed in parallel. gDNA was extracted from using Qiagen DNeasy kit (Qiagen, #69506), eluted in 60 μl of elution buffer and quantified using QuBit (Thermo). One nanogram of gDNA was amplified via three PCRs: two nested PCRs to amplify the target region and a third, indexing PCR to add the NGS adapters and indices. PCR-1 was run to 20 cycles using the primers presented in Supplementary Table 6. One microliter of 1:10 dilution of unpurified PCR-1 product was used for PCR-2, which was run to 20 cycles with the primers presented in Supplementary Table 7. The PCR-2 product was purified using 1x volume of AMPure XP beads (Beckman Coulter) and eluted in 15 μl of IDTE buffer (IDT DNA). One microliter of this product was PCR-3 was run to 7 cycles with the primers presented in Supplementary Table 8. The final product was purified using 0.8x volume of AMPure XP beads, eluted in 15 μl of IDTE and quantified using QuBit. Products from different samples were pooled and sequenced using a MiSeq (Illumina). For classification of pooling primers from different targets that yielded a balanced representation of all the sequenced targets among the NGS reads. For the ten-target mRNA, we pooled all the PCR-1 primers and all the PCR-2 primers in equimolar amounts to a final concentration of 5 μM per oligo. For the 20-target mRNA, we made three sets of primers per PCR: set 1 with targets 2–6, set 2 with targets 8–11 and set 3 with targets 1, 7 and 12. Targets were then de-multiplexed during the data analysis (see below).

**Determining mutation levels and mutation outcomes of mRNAs.** To determine the mutation levels of the different mRNA targets, we first de-multiplexed these targets (which were amplified in a multiplexed fashion) by aligning the first 50bp of each PE read to the genome. A given read was considered to contain an mRNA target if the PE alignment fell within a window of 1,000 bp from the expected genomic location of the target. A mutation was called if the intact theoretical protospacer sequence was not found in the read.

For classification of the mRNA target mutations, we defined for each target site two key sequences that were, respectively, 20 bp upstream and downstream of the expected genomic location of the cut site. For each read aligning to a target site, these two key sequences were identified and the distance between them was computed. Reads with distances shorter than the expected value were classified as deletions, while reads with distances longer than expected were classified as insertions. Reads with the expected distance between the key sequences but with mutations in the protospacer were classified as single-nucleotide variants (SNVs).

**ATAC-seq.** ATAC-seq was performed following the Omni-ATAC protocol41 using the amplification protocol and primers described in ref. 56. Primers are also presented in Supplementary Table 9. A detailed protocol can be found in ref. 56. Cells were washed with PBS, collected via scraping and counted. A total of 50,000 cells were used for ATAC. Collected cells were then pelleted, the supernatant was removed and the cells were resuspended in 50 μl of cold lysis buffer, gently mixed and kept on ice for 3 min. Then, 5.25 μl of transfer wash buffer was then added and gently mixed. Cells were then pelleted, resuspended in 50 μl of transposition reaction and incubated at 37 °C for 30 min. Transposed DNA was column purified and eluted in 21 μl of EB. Samples were pre-amplified, followed by qPCR to determine the number of cycles needed for final amplification (one-third of saturation). Final elution was purified using AMPure beads and eluted in 32 μl IDTE. Final libraries were quantified using 2% agarose gel, pooled, quantified with QuBit, Bioanalyzer and qPCR, then sequenced on a NovaSeq 500 (Illumina) using paired 2 × 50 bp reads.
CRISPR activation and deactivation. The special gRNA or pcrRNAs were used in the place of normal cRNAs when complexed with tracrRNA. For activation, Cas9/gRNA was first electroporated into cells, plated onto 12-well plates, then incubated for 12h to allow stable Cas9 binding but not cleavage. Next, cells were exposed to 1 min of 365nm light exposure from a handheld blacklight (https://www.amazon.com/AXMAN-Ultraviolet-365nm-Detector-Flashlight/dp/B06XW7S1CS/). Either one, three or six flashlights were used at once. When multiple flashlights are used, they are conveniently held together using a 3D-printed flashlight holder (https://github.com/rogerous/chipseq_pcrRNA/blob/master/jaxman_1LED_flashlight_holder_design/files/8zeFEGPVoSaat). Samples were collected without light exposure, or 10 and 30 min after light exposure. For deactivation, Cas9/pcRNA was first electroporated into cells, plated onto 12-well plates, incubated for 2h, then exposed to light of the same dose. Samples were collected during the time of light exposure, or at 1h, 2h and 4h after light exposure.

Immunofluorescence microscopy of 33BP1 foci after multi-target Cas9 activation. The number of endogenous 53BP1 foci in cells was evaluated through Immunofluorescence microscopy of 53BP1 foci after multi-target Cas9 activation. The number of endogenous 53BP1 foci in cells was evaluated through Immunofluorescence microscopy of 53BP1 foci after multi-target Cas9 activation. The number of endogenous 53BP1 foci in cells was evaluated through Immunofluorescence microscopy of 53BP1 foci after multi-target Cas9 activation.

To determine the number alignments for each target, we outputted each gRNA + PAM into a FASTA file and ran bowtie2 with `-k 1000` mode, which searches up to 1,000 alignments for each line in the FASTA, that is, each target sequence. To determine the number alignments for each target, we outputted each gRNA + PAM into a FASTA file and ran bowtie2 with `-k 1000` mode, which searches up to 1,000 alignments for each line in the FASTA, that is, each target sequence.

Discovery and characterization of mrRNA sequences. Starting from a 280 bp SINE sequence, for all 20 bp substrings in both the forward and reverse complement direction, we obtained all 20 bp sequences with up to three mismatches. The template restricted to the nine most PAM-proximal nucleotides. GC content was restricted to 40–70%. This resulted in 75,626 unique target sequences.

To determine whether each alignment was within a ReSeq gene annotation and the ChromHMM epigenetic states. We then analyzed the UCSC Genome Browser (https://genome.ucsc.edu). The UCSC Genome Browser was used to pre-determine sample size. No data were excluded from the analyses.

Ambiguous read proportions from simulated ChIP–seq reads. For gRNA with 100–300 on-target sites in the genome, we simulated 100 PE 200–600-bp-long (uniform distribution) sequencing reads. The reads were randomly chosen to either span the cut site, reside PAM-distal or reside PAM-proximal to the cut. For PAM-distal or PAM-proximal reads, the distance from the edge of the DNA to the cut site was drawn from an exponential distribution. Both 2 x 36 PE reads and 2 x 75 PE reads were simulated. The PE reads were outputted to FASTA files (read 1 and read 2), and bowtie2 was used to determine up to ten alignments for each simulated read pair:

```
bowtie2 -f -p 9 -local -k 10 -X 1000 -no-mixed -no-discordant -x [path to genome] -1 [path to read1] -2 [path to read2] -S [path to output SAM]
```

The code subsequently determines whether the original position of the read pairs matches the best alignment based on bowtie2, and whether this best alignment has the uniquely best alignment score. The proportion of reads that satisfy these requirements represent the proportion of uniquely best alignments. The proportion of ambiguous alignments is 1 minus this value.

Ambiguous read proportions from real ChIP–seq reads. We used all dCas9 binding positions for analysis. For each binding position, we generated PE ChIP–seq reads found within a specified window width centred at the Cas9 binding site into FASTA read 1 and read 2 file formats. Then the section 'Ambiguous read proportions from simulated ChIP–seq reads' was followed, starting with use of bowtie2. Window widths of 1500 bp were used for Cas9 ChIP–seq, and 2,500 bp for MRE11.

Calculating enrichment for MRE11, Cas9, yH2AX and 53BP1 ChIP–seq. For calculation, we determined the reads per million (RPM) in specific window widths centred at all cut sites. We used a window of 200 kb for both 53BP1 and yH2AX, 2,500 bp for MRE11 and 1,500 bp for Cas9. For MRE11 and Cas9, additional code analyses the exact read positions and determines if a PE sequencing read fragment spans the cut site (‘span’), or if a sequenced DNA fragment begins within 5 kb from the cut site (‘abut’). To determine ‘dist’ = 4, ‘dist’ = 4’, ‘prox’ + 4’, or ‘prox – 4’, we analysed the DNA fragment position according to the rules specified for these read species.

Enrichment profiles for MRE11 and Cas9 ChIP–seq (also spanning ATAC–seq) at base-pair resolution. At each genomic position in a window centred at each cut site, each PE read within this window is retrieved. The number of PE reads that map to each base pair is tallied. The middle region of PE read fragment that is not likely to be sequenced is also included in this tally. We used a window of 2,500 bp for MRE11, 1,500 bp for Cas9 and 3 kb for ATAC–seq.

Enrichment profiles for yH2AX, 53BP1 and ATAC–seq at window widths. To determine profiles of yH2AX and 53BP1, we calculated the number of sequencing reads (RPM) in each 10 kb window from the cut site, extending to 2 mb both upstream and downstream of cut sites. For ATAC–seq, we calculated RPM in a 4 kb sliding window incremented every 1 bp, extending to 1.5 kb both upstream and downstream of cut sites.

To determine wider levels of potential ATAC-seq enrichment, we used the same function to calculate RPM in each 1 kb window from the cut site, extending to 50 kb both upstream and downstream of cut sites.

Genome-wide Cas9 binding from dCas9 ChIP–seq. We used macs2 to find all dCas9 binding peaks, using a no-Cas9 sample for negative control, via the command:

```
macs2 callpeak -t [path/to/sample] -c [path/to/peak/track] -outdir [path/to/output] -name [name/of/output] -f BAMPE -g hs
```

Next, for each macs2 discovered peak with fold enrichment ≥4, a custom algorithm attempts to identify the target sequence position for Cas9 binding or cleavage that best explains the peak. This may be problematic for target sites with multiple mismatches. We used the following assumption to simplify the problem:

1. There is only one correct Cas9 binding/cleavage sequence within the 400 bp window of the macs2 predicted peak centre, and
2. The correct Cas9 binding/cleavage sequence is one with the fewest mismatches.

Enrichment measurements of epigenetic markers. Datasets used are indicated in 'Data availability'. For enrichment, we use a 50 kb radius for RNA-seq, H3K4me1, H3K4me3, H3K27ac and H3K36me3, a 50 bp radius for DNase I and ATAC–seq, and a 10 bp radius for micrococcal nuclease digestion with deep sequencing (MNase–seq). The number of reads that are found in each specified window width is outputted, normalized by the total RPM.

Machine learning model. We used the random forest regressor from scikit-learn. For mismatch information, features were obtained from one-hot encoding of mismatch state at each position along the protosparse. For epigenetic information, the RPM enrichment was directly used as features. The predicted output is the level of dCas9 binding or MRE11 enrichment, also measured as RPM. The machine learning model was trained using five-fold cross-validation on a training dataset composed of a random 70% of the total dataset. The remaining 30% was used for evaluation and featured in these figures comparing predicted versus actual values.

ATAC–seq read length distributions. For each PE ATAC–seq read fragment in a 3 kb window centred at all Cas9 on-target sites, its length was recorded. The distribution of DNA length across all target sites, along with exponential decay curve fitting, was computed in Microsoft Excel.

Statistics and reproducibility. ChIP–seq, ATAC–seq, amplicon sequencing and BLISS experiments were performed in biological replicates. No statistical method was used to pre-determine sample size. No data were excluded from the analyses.
The experiments were not randomized. The investigators were not blinded to allocation during experiments and outcome assessment.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability**

Deep-sequencing data generated for this study have been deposited in Sequence Read Archive under BioProject accession PRJNA733683. Sequencing data were analysed using the hg38 genome assembly ([https://www.ncbi.nlm.nih.gov/assembly/GCF_000001405.26](https://www.ncbi.nlm.nih.gov/assembly/GCF_000001405.26)). Previously published, publicly available epigenetic datasets used in this study are from HEK293 cell lines: ATAC–seq (SRR6418075), DNase I (ENCFF12OFX), H3K4me1 (ENCFF909FBEY), H3K4me3 (ENCFF122XLY), H3K9me3 (ENCFF141ZEQ), H3K27ac (ENCFF588KSR), H3K36me3 (ENCFF593SUW), MNase–seq (ERR2403161) and RNA-seq (SRR5627161). Datasets starting with ENCF can be found and downloaded from ENCODE ([https://www.encodeproject.org/](https://www.encodeproject.org/)). Dataset starting with SRR or ERR can be found and downloaded from NIH’s SRA ([https://www.ncbi.nlm.nih.gov/sra](https://www.ncbi.nlm.nih.gov/sra)). Source data are provided with this paper. All other data supporting the findings of this study are available from the corresponding author on reasonable request.

**Code availability**

Analysis code is available on GitHub ([https://github.com/rogerzou/multitargetCRISPR](https://github.com/rogerzou/multitargetCRISPR)). This software is open-source, modular and well documented. It enables in silico discovery and characterization of mgRNAs, alongside comprehensive analysis of ChIP–seq, BLISS and ATAC–seq datasets.

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**Author contributions**

R.S.Z. and T.H. conceived the project. R.S.Z. and A.M.-G designed and performed experiments and wrote the manuscript with contributions from all authors. R.S.Z. analysed data and prepared figures. Y.L. and H.B.L. assisted with ChIP–seq. L.S. and R.K.D. assisted with cell line generation and cloning. J.X.J.L. assisted with data analysis. R.K. supervised the design of mutation kinetics experiments. T.H. supervised the project.

**Competing interests**

Johns Hopkins University has submitted two patent applications, in which R.S.Z., Y.L. and T.H. are authors, on previously published methods for Cas9 activation and deactivation that were used in this study. The remaining authors declare no competing interests.

**Additional information**

Extended data is available for this paper at [https://doi.org/10.1038/s41556-022-00975-z](https://doi.org/10.1038/s41556-022-00975-z).

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Extended Data Fig. 1 | See next page for caption.
Extended Data Fig. 1 | Additional in silico and experimental characterization of multi-target gRNAs. a,b, Same as Fig. 1b and Fig. 1h, respectively, but for the Alu SINE in the mouse mm10 genome. c,d, Same as Fig. 1b and Fig. 1h, respectively, but for the B4 SINE in the mouse mm10 genome. e,f, Same as Fig. 1b and Fig. 1h, respectively, but for the DR-1 SINE in the zebrafish danRer11 genome. g,h, Same as Fig. 1b and Fig. 1h, respectively, but for the DR-2 SINE in the zebrafish danRer11 genome. i-k, Same as Fig. 1b and Fig. 1h, respectively, but for two different mgRNAs from the (i) Alu SINE in the human hg38 genome, (j) B4 SINE in the mouse mm10 genome, and (k) DR-1 SINE in the zebrafish danRer11 genome. l, No-dox control mutation curve. Dox-inducible Cas9 cells genomically integrated with a 10-target mgRNA (same cells as in Fig. 1l) were grown and passaged without dox exposure. Cells were harvested at different time points (0, 2, 6 and 10 days). m, Mutation curve of 20-target mgRNA in HeLa cells with dox-inducible Cas9. Figure details are the same as Fig. 1l. n, Mutation signatures for 10-day samples from m. Figure details are the same as in Fig. 1m. o, Same reproducibility analysis and figure details as in Fig. 1n but using the mgRNA from m. p-s, For all 4 target sites, the PAM-distal side is oriented on the left. (p-r) The first three examples exhibit Cas9 PAM-distal bias and MRE11 PAM-proximal bias. However, the last example (s) in chr2:140756521-140759021 illustrates an exception where both Cas9 and MRE11 exhibit PAM-distal bias. t, Same as Fig. 3j, but with the 20-target mgRNA 2. Source numerical data are available in source data.
Extended Data Fig. 2 | Further analysis and machine learning modeling at 30 min. a, Mean quantification of Fig. 4b,c for two biological replicates. b,c, Two replicates of correlation between MRE11 and BLISS enrichment in 2500 bp windows centered around all cut sites. d-i, Same as Fig. 4 h-m but for samples evaluated at 30 minutes after (d)Cas9 delivery. j, Violin plots of background-subtracted ATAC-seq reads per million (RP) enrichment at all on-target sites. ‘neg’ indicates Cas9-negative cells. ‘00m’, ‘10m’, and ‘30m’ indicate cells with vCRISPR Cas9/cgRNA without light exposure, 10 minutes after light exposure, and 30 minutes after light exposure, respectively. Comparison using two-sided unadjusted Student’s t-test. n.s. indicates no significance, **** indicates p < 0.0001. p-values from left to right are: 0.64, 0.25, and 6.85E-8. k, Quantification of Fig. 7e. n=3 biologically independent experiments, data presented as mean ± 1 SD. Source numerical data are available in source data.
Reporting Summary

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- A full description of the statistical parameters including central tendency (e.g., means) or other basic estimates (e.g., regression coefficient) AND variation (e.g., standard deviation) or associated estimates of uncertainty (e.g., confidence intervals)
- For null hypothesis testing, the test statistic (e.g., t, F, r) with confidence intervals, effect sizes, degrees of freedom and P value noted
  *Give P values as exact values whenever suitable.*
- For Bayesian analysis, information on the choice of priors and Markov chain Monte Carlo settings
- For hierarchical and complex designs, identification of the appropriate level for tests and full reporting of outcomes
- Estimates of effect sizes (e.g., Cohen’s d, Pearson’s r), indicating how they were calculated

*Our web collection on [statistics for biologists](https://www.nature.com/nbt/features/analysis/) contains articles on many of the points above.*

Software and code

Policy information about [availability of computer code](https://example.com)

**Data collection**

No software was used for data collection

**Data analysis**

bc2fastq2, bowtie2, macs2, samtools1.10, scikit-learn1.1, custom code available in [https://github.com/rogerzou/multitargetCRISPR](https://github.com/rogerzou/multitargetCRISPR)

For manuscripts utilizing custom algorithms or software that are central to the research but not yet described in published literature, software must be made available to editors and reviewers. We strongly encourage code deposition in a community repository (e.g. GitHub). See the Nature Portfolio [guidelines for submitting code & software](https://www.nature.com/natureportfolio/policies) for further information.

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All manuscripts must include a [data availability statement](https://example.com). This statement should provide the following information, where applicable:

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Deep-sequencing data that support the findings of this study have been deposited in Sequence Read Archive under BioProject accession PRJNA733683. Sequencing data was analyzed using the hg38 genome assembly ([https://www.ncbi.nlm.nih.gov/assembly/GCF_000001405.26](https://www.ncbi.nlm.nih.gov/assembly/GCF_000001405.26)). Previously published, publicly available epigenetic datasets used in this study are from HEK293 cell lines: ATAC-seq (SRR6418075), DNaseI (ENCCFF1200XFB), H3K4me1 (ENCCFF909ESY), H3K4me3 (ENCCFF921BHL), H3K9me3 (ENCCFF1412EQ), H3K27ac (ENCCFF5BBKSR), H3K36me3 (ENCCFF593SUW), MNase-seq (ERR2403161), and RNA-seq (SRR5627161). Datasets starting with ENCCF can be found and downloaded from ENCODE ([https://www.encodeproject.org](https://www.encodeproject.org/)). Dataset starting with SRR or ERR can be found and downloaded from NIH's SRA ([https://www.ncbi.nlm.nih.gov/sra](https://www.ncbi.nlm.nih.gov/sra)). Source data have been provided in Source Data. All other data supporting the findings of this study
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Life sciences study design

All studies must disclose on these points even when the disclosure is negative.

| Sample size | No statistical method was used to predetermine sample size. |
|-------------|------------------------------------------------------------|
| Data exclusions | No data was excluded from the analysis |
| Replication | Reproducibility was confirmed by computing Pearson correlation between replicates (when applied). This analysis is clearly stated and shown in the text, methods section, or supplementary materials. All experiments were performed in biological replicates. |
| Randomization | The experiments were not randomized. This is not required for the current study because there is no participants involved in the experiments. |
| Blinding | The Investigators were not blinded to allocation during experiments and outcome assessment. This is not relevant for this study because there is no participants involved in the experiments. |

Reporting for specific materials, systems and methods

We require information from authors about some types of materials, experimental systems and methods used in many studies. Here, indicate whether each material, system or method listed is relevant to your study. If you are not sure if a list item applies to your research, read the appropriate section before selecting a response.

### Materials & experimental systems

| n/a | Involved in the study |
|-----|-----------------------|
| ☑ Antibodies | |
| ☑ Eukaryotic cell lines | |
| ☑ Palaeontology and archaeology | |
| ☑ Animals and other organisms | |
| ☑ Human research participants | |
| ☑ Clinical data | |
| ☑ Dual use research of concern | |

### Methods

| n/a | Involved in the study |
|-----|-----------------------|
| ☑ ChiP-seq | |
| ☑ Flow cytometry | |
| ☑ MRI-based neuroimaging | |

#### Antibodies

Antibodies used: Cas9 – Diagenode C15310258; MRC11 – Novus N8100-142; yH2AX – Abcam ab81299; 53BP1 – Novus N8100-305; Alexa647 Secondary antibody – ThermoFisher, A-21235

Validation: For all the antibodies used, there was a statement in the manufacturer’s website indicating that they are suitable for immunoprecipitation assays. Also, we performed a no-treatment (negative) control for all the antibodies and cell lines tested.

#### Eukaryotic cell lines

Policy information about [cell lines](http://cell-lines.org)

- **Cell line source(s)**: HEK293T (ATCC® CRL-3216™) and Hela (ATCC® CCL-2™)
- **Authentication**: The cell lines were not authenticated
- **Mycoplasma contamination**: Cells were tested regularly for mycoplasma contamination and tested negative.
- **Commonly misidentified lines (See [ICGAR](http://icgar.org), register)**: None
## Data deposition

- Confirm that both raw and final processed data have been deposited in a public database such as [GEO](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA733683/).

### Data access links
- May remain private before publication.
  - [https://www.ncbi.nlm.nih.gov/bioproject/PRJNA733683/](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA733683/)

### Files in database submission
- fastq files of all ChIP-Seq and ATAC-Seq experiments, including biological replicates

### Genome browser session
- [UCSC](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA733683/)

## Methodology

| Replicates                  | Two biological replicates. |
|-----------------------------|---------------------------|
| Sequencing depth            | Paired-end, 2x36 bp, 15 million paired-end reads per sample |
| Antibodies                  | Cas9 – Diagenode C15310258; MRE11 – Novus NB100-142; γH2AX – Abcam ab81299; 53BP1 – Novus NB100-305 |
| Peak calling parameters     | macs2 callpeak -t [path/to/sample] -c [path/to/negctrl] --outdir [path/to/output] --name [name/of/output] -f BAMPE -g hs |
| Data quality                | Reads were filtered for mapping quality >= 25. Singleton reads, potential PCR duplicates and index reads were removed. |

### Software
- bowtie2, macs2, samtools
- custom code available at [https://github.com/rogerzou/multitargetCRISPR](https://github.com/rogerzou/multitargetCRISPR)