Nucleosomes Impede Cas9 Access to DNA in vivo and in vitro

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Competing interests: MAH, LAG, and JSW have filed a patent application (US Appln. No. 62/024,373) related to CRISPRi screening technology.
Abstract

The prokaryotic CRISPR (Clustered Regularly Interspaced Palindromic Repeats)-associated protein, Cas9, has been widely adopted as a tool for editing, imaging, and regulating eukaryotic genomes. However, our understanding of how to select single-guide RNAs (sgRNAs) that mediate efficient Cas9 activity is incomplete, as we lack insight into how chromatin impacts Cas9 targeting. To address this gap, we analyzed large-scale genetic screens performed in human cell lines using either nuclease-active or nuclease-dead Cas9 (dCas9). We observed that highly active sgRNAs for Cas9 and dCas9 were found almost exclusively in regions of low nucleosome occupancy. In vitro experiments demonstrated that nucleosomes in fact directly impede Cas9 binding and cleavage, while chromatin remodeling can restore Cas9 access. Our results reveal a critical role of eukaryotic chromatin in dictating the targeting specificity of this transplanted bacterial enzyme, and provide rules for selecting Cas9 target sites distinct from and complementary to those based on sequence properties.
Introduction

CRISPR (Clustered Regularly Interspaced Palindromic Repeats) prokaryotic adaptive immune systems have yielded transformative tools for manipulating eukaryotic genomes. Most notably, the CRISPR associated Cas9 protein from *Streptococcus pyogenes*, together with a single chimeric guide RNA (sgRNA), provides a programmable endonuclease that has revolutionized our ability to edit genomes [1]. Cas9 has been further modified to a nuclease-dead form (dCas9) to provide a programmable DNA binding protein that can be fused to effector domains, making it possible to turn on or off targeted genes, mark specific genomic loci with fluorescent proteins, or alter epigenetic marks [1-8]. A central challenge in implementing these tools is identifying effective and specific sgRNAs. While much of the effort to define relevant rules has focused on the sequence of the target site and sgRNA [9-12], these only partially predict Cas9 activity and suggest that additional determinants likely exist.

Chromatin structure may represent a key parameter governing Cas9 efficacy in eukaryotic cells. CRISPR evolved in archea and bacteria [13] and is likely not optimized to explore and modify large, chromatin-bound eukaryotic genomes, a hypothesis supported by several studies that pointed to a correlation between rates of DNA binding and cleavage in regions of open chromatin as measured by DNase I hypersensitivity [9, 14, 15]. Additionally, recent single-molecule imaging studies have shown that dCas9 explores euchromatin more frequently than it does heterochromatin [16].

We hypothesized that nucleosomes, the basic unit of chromatin structure, are an important impediment to Cas9 recognition. Here, we addressed this hypothesis *in vivo* by leveraging large datasets collected from over 30 sgRNA tiling and genome-scale genetic screens [17]. We found that regions of high nucleosome occupancy *in vivo*, as determined by MNase-seq (micrococcal nuclease sequencing) [18, 19], were strongly depleted of highly active sgRNAs for CRISPR interference (CRISPRi) [3, 17] and nuclease-active Cas9. We complemented these results with *in vitro* experiments demonstrating that formation of a nucleosome provides a direct and profound block to dCas9 binding and Cas9 cleavage. Despite this strong barrier to Cas9 activity, we found that addition of a chromatin remodeling enzyme to chromatinized DNA *in vitro* can restore Cas9 access to nucleosomal DNA, highlighting one route by which CRISPR may still be able to modify chromatin *in vivo*. Our results reveal a fundamental aspect of the mechanism by which this transplanted bacterial enzyme interacts with eukaryotic chromatin, and provide a new dimension for selecting highly active sgRNAs.
Results

CRISPRi activity is periodic and out-of-phase with nucleosome positioning

In order to study features of chromatin that affect CRISPRi activity, we integrated data from whole-genome screens testing for a wide range of phenotypes performed with a previously described CRISPRi library targeting each gene with ~10 sgRNAs [17]. We selected 30 screens performed in the cell line K562 expressing dCas9-KRAB (M.L., B.A., B.B., C.Y.P., M.K., Y.C., J.F., and J.N., personal communication) and set a threshold for high-confidence hit genes, which allowed us to assess the relative strength of phenotypes for the sgRNAs targeting these genes. Specifically, we analyzed 18,380 sgRNAs targeting 1,539 genes, and generated “activity scores” by normalizing sgRNA phenotypes to the average of the 3 sgRNAs with the strongest phenotypes for each gene (Figure 1A and Supplementary File 1). To assess how CRISPRi activity varies with respect to the transcription start site (TSS), we plotted the average sgRNA activity score as a function of distance to the FANTOM-annotated TSS [20] (Figure 1B). This analysis revealed a robust, periodic pattern of activity, with peaks at ~190bp intervals relative to the TSS. This periodicity was highly reminiscent of patterns previously described for nucleosomes [21]. Indeed, analysis of K562 MNase-seq data from the ENCODE consortium [19, 22] revealed that the average nucleosome signal was strongly anti-correlated with CRISPRi activity (Figure 1B), suggesting that high nucleosome occupancy leads to low CRISPRi activity.

To further explore this inverse relationship between CRISPRi activity and nucleosome organization, we exploited the previous observation that nucleosome phasing is more pronounced in well-expressed genes [19]. We grouped genes by their expression in K562 [22, 23] and analyzed CRISPRi activity and nucleosome occupancy within each group. In support of a connection, we found that peak to trough amplitudes of both features were larger for highly expressed genes (Figures 1C,D).

We also analyzed the MNase signal at each sgRNA target site to determine whether nucleosome occupancy could directly explain variation in CRISPRi activity between individual sgRNAs (Figure 1E). Consistent with the hypothesis that nucleosomes exclude dCas9, almost all of the highly active sgRNAs targeted sites corresponded with low MNase-seq signal. However, not all sgRNAs targeting sites with low nucleosome occupancy were highly active, matching previous findings that sgRNA and target DNA sequence features also influence efficiency of the CRISPR system [9-12]. To exclude the possibility that differences in sequence features alone between nucleosome-bound and -free regions could explain the periodicity of CRISPRi activity (rather than the presence of nucleosomes per se), we performed linear regression using MNase signal, sgRNA length [12, 17], and a validated sgRNA sequence...
scoring algorithm [10]. We found that each parameter individually correlated with sgRNA activity (p < 10\(^{-58}\) for each; Figure 1F). Importantly, correction for sequence and length features had minimal impact on the ability of the MNase signal to predict CRISPRi activity (p < 10\(^{-85}\) after correction). Indeed, a linear fit of all three features provided a still stronger correlation (p < 10\(^{-221}\); Figure 1F, far right column), suggesting that incorporating nucleosome occupancy in future sgRNA design algorithms could significantly improve predictive value. We have recently developed a comprehensive algorithm for predicting highly active CRISPRi sgRNAs that, by accounting for nucleosome positioning, higher order sequence features, and non-linear relationships in these parameters, shows even greater correlation with this dataset that was already enriched for active sgRNAs using our original CRISPRi library design principles [17] (cross-validation R\(^2\)=0.32; Horlbeck et al., manuscript in preparation [24]).

**Nuclease-active Cas9 activity anti-correlates with nucleosome occupancy**

In order to generate a dataset for evaluating the effect of nucleosome positioning on nuclease-active Cas9 in K562 cells, we took advantage of our previously described library densely tiling sgRNAs in 10kb windows around the TSS of 49 genes known to modulate susceptibility to the toxin ricin [17, 25] and tested Cas9-expressing cells for resistance or sensitivity to ricin (Supplementary File 2). We observed phenotypes consistent with the expected knockdown phenotype primarily in coding sequences (CDS) but also in some promoter regions, consistent with recent results showing that modifications introduced by Cas9 can disrupt cis-regulatory regions [26]. Analysis of CDS-targeting sgRNAs revealed that strong phenotypes were found predominantly in regions of low MNase-seq signal (Figure 2A), although this relationship was less pronounced than for CRISPRi. This may be due to decreased phasing of nucleosomes within the gene body [21]. When we analyzed all sgRNAs in the library, thus incorporating information from cis-regulatory regions where nucleosomes are well phased, we found the effect of nucleosome position to be even stronger (Figure 2-figure supplement 1). Although nucleosome occupancy was predictive of CRISPR activity independent of sgRNA sequence features, a much stronger correlation was obtained when all features were considered (Figure 2B). Therefore, nucleosome organization likely represents an important feature for CRISPR sgRNA design and should be considered a key contributing factor in interpreting future tiling mutagenesis experiments of coding and non-coding regions.
Nucleosomes are sufficient to fully block cleavage by Cas9 *in vitro*

Our *in vivo* experiments reveal a strong anti-correlation between Cas9 mediated downstream phenotypic outputs and nucleosome positioning, but do not directly report on the ability of Cas9 to access nucleosomal DNA. Factors other than Cas9 access could contribute to the observed correlation in our data. For example, (d)Cas9 may bind or cut equally well in nucleosome-bound and un-bound regions, but may exert the observed modulation of gene expression through interference with transcriptional pausing, splicing, regulatory looping, or binding of important regulatory factors, processes which also correlate with nucleosome organization [27-33].

To determine whether (d)Cas9 activity is indeed directly affected by the presence of nucleosomes, we turned to a purified *in vitro* reconstituted system. Mouse histones were recombinantly expressed, purified, and assembled into a nucleosome using 147bp of the Widom 601 positioning sequence (Figures 3A,B) [34]. Conveniently, the 601 sequence contains numerous NGG protospacer adjacent motifs (PAMs) required for Cas9 recognition, spanning the full length of the DNA at different helical positions, allowing us to test the effects of position and solvent accessibility within the nucleosome (Figure 3C). We first tested the ability of Cas9 to cleave nucleosomal DNA. We pre-loaded purified Cas9-HaloTag with *in vitro* transcribed sgRNA, then introduced either naked 601 DNA or that same DNA assembled into a nucleosome (Figures 3A,D and Figure 3-figure supplement 1A ). Fluorescent labeling of the DNA allowed us to visualize the cleavage products on a denaturing Urea-PAGE gel. In agreement with our *in vivo* data, the nucleosome protected its DNA from cleavage by Cas9, and complete protection from cleavage was observed to be independent of target position within the nucleosome (Figures 3E,F and Figure 3-figure supplement 1B,C). While our manuscript was under review, a study by Hinz and colleagues reported the similar *in vitro* finding that Cas9 nuclease activity is inhibited within the nucleosome but not at adjacent linker sequences[35]. Previous single-molecule and biochemical studies have established that nucleosomal DNA undergoes transient unwrapping or breathing at the entry and exit sites, creating a gradient of accessibility along the nucleosome [36-41]. This property is often credited for the observed position dependent binding patterns of many transcription factors and DNA binding proteins. Interestingly, our data suggests that the cleavage activity of Cas9 *in vitro* is not detectably influenced by this effect, although less stable nucleosomes than the Widom 601 nucleosome or target sites closer to the nucleosome edge may exhibit more breathing and thus more accessibility than those tested here.
Cas9 is unable to bind nucleosomal DNA in vitro

Cleavage of DNA by Cas9 has been described as a stepwise process [15, 16, 42] in which Cas9 must first scan for PAMs, unzip the DNA duplex, and fully pair the guide RNA and target DNA prior to cleavage. While our in vitro data show that full pairing and cleavage is prevented by the presence of a nucleosome, we wondered if binding without cleavage, especially at the more accessible ends of the nucleosome, might still occur. Additionally, the first step in binding, PAM recognition, may be governed by helical location within the nucleosome as well as its proximity to the more dynamic ends. Histones make contacts with the DNA at every helical turn, thus a PAM may fall on the outside of the nucleosome, exposing it to solvent, or on the inside at the DNA-histone interface. To test the influence of target location within the nucleosome on Cas9 binding, we used an electrophoretic mobility shift assay to monitor binding of dCas9 to Cy3 end-labeled 601 DNA (Figures 4A,B), either free or assembled into a nucleosome containing Alexa Fluor 647-labeled histone H2B (Figure 4-figure supplement 1A). Consistent with our cleavage results, binding by dCas9 was abolished by the presence of the nucleosome, regardless of the targeted dCas9 binding site (Figure 4C-E and Figure 4-figure supplement 2A,B).

To better understand how a nucleosome might impede Cas9 binding, we aligned the available crystal structures of DNA-bound Cas9 and the structure of the 601 nucleosome ([43], PDB ID 3LZ0; [44], PDB ID 4UN3). We superimposed the target DNA in the Cas9 crystal structure with the DNA in the nucleosome structure at a site where the two DNA paths gave the best fit. The resulting combined structure reveals that the Cas9 protein poses significant steric clashes with the histones (Figure 4F). Given the extent of overlapping densities in the two structures, it seems unlikely that the histones and Cas9 could co-occupy the same piece of DNA. Additionally, it may be important to note that unlike other DNA binders such as transcription factors, binding by Cas9 constitutes melting of target DNA, which may pose an additional barrier to binding on a nucleosome. This hypothesis leaves two possible outcomes of targeting Cas9 to nucleosomal DNA: either Cas9 is capable of displacing histones in order to engage nucleosomal DNA, or it is excluded altogether. Our data support the latter conclusion.

The chromatin remodeling enzyme yChd1 can restore access to nucleosomal DNA in vitro

The nucleosome landscape in eukaryotic chromatin is strongly fixed in certain regions, including those proximal to the TSS of actively transcribed genes, but is highly dynamic in others. Nucleosome positioning is regulated by both intrinsic DNA sequence preferences as well
as extrinsic factors such as chromatin remodeling enzymes [21, 45, 46]. In order to model how these dynamics affect Cas9 accessibility to DNA in vitro, we turned to a chromatinized plasmid system. We dialyzed plasmid DNA containing a single 601 nucleosome positioning sequence with a sub-saturating quantity of purified histone octamers, and confirmed the stability of the resulting chromatin assemblies by MNase digestion (Figures 5A,B). We tested whether the 601 sequence was occupied by a nucleosome using restriction enzyme accessibility mapping, and found that sites with the 601 sequence were protected from digestion while sites immediately adjacent were not (Figure 5C). To recapitulate the effects of chromatin remodeling in vitro, we used a purified, truncated form of the Snf2-like chromatin remodeling enzyme Chd1 from Saccharomyces cerevisiae (yChd1), which had previously been shown to mediate nucleosome sliding in an ATP-dependent manner without additional co-factors [47, 48]. Consistent with this sliding (rather than nucleosome ejecting) activity, when we digested the chromatinized plasmid assembly with the frequent cutter HaeIII, we observed a shift towards lower molecular weight bands upon addition of yChd1, indicative of diminished protection of HaeIII sites, but overall the digest retained higher bands relative to the naked plasmid control (Figure 5D).

We next sought to test whether yChd1 affected the accessibility of the DNA at the 601 positioning sequence. As with the restriction enzyme digestion of the chromatinized plasmid in the absence of yChd1, we found that sites along the nucleosome positioning sequence were protected from Cas9 cleavage, consistent with our mononucleosome results. When we assayed restriction enzyme or Cas9 accessibility in the presence of yChd1, however, we found that the cleavage efficiency was restored to 50-80% of the corresponding naked plasmid control (Figures 5E-G). Notably, the percent protection at the EcoRI site adjacent to the positioned nucleosome did not decrease upon addition of yChd1, demonstrating that the decrease in protection along the 601 sequence was mediated by the nucleosome displacement activity of yChd1 rather than by a non-specific effect on cleavage efficiency. While our data with the chromatinized plasmid system confirm our findings that a well-positioned nucleosome provides a profound block to Cas9 cleavage, our further finding that chromatin remodeling restores access to nucleosomal DNA provides one potential mechanism by which Cas9 may efficiently modify broad portions of eukaryotic genomes. This plasmid model could be further exploited to assay the activity of Cas9 at nucleosome-free and boundary sites, and thus derive biophysical parameters governing Cas9-chromatin interactions.
Despite its swift success as a repurposed tool for gene editing, imaging, and transcription modulation, the ability of this prokaryotic CRISPR/Cas9 system to effectively navigate eukaryotic chromatin has remained poorly understood. Here, we show that the nucleosome, the basic unit of chromatin, poses a strong barrier to Cas9, both in vitro and in vivo. By masking ~147bp of DNA, the nucleosome effectively reduces the size of the eukaryotic genome available to Cas9. Previous studies using ChIP-seq to assay Cas9 binding have shown that off-target binding at PAM plus seed sequences more frequently occurs in regions of open chromatin [15, 49]. Our data expand upon these findings to show that the discrete pattern of nucleosome organization is able to modulate the efficiency of Cas9 binding and cleavage at on-target sites. The practical implications of these observations are underscored by our finding that accounting for nucleosome occupancy offers a significant improvement in predictive power for sgRNA design.

While our data show that nucleosomes strongly protect their DNA from Cas9 binding and cleavage in vitro, their organization in cells is not static. Transient displacement of nucleosomes occurs during replication, remodeling, and transcription. By adding the chromatin remodeling enzyme yChd1 to nucleosomes in vitro, we demonstrate that this displacement can in fact restore Cas9 access to DNA. However, despite brief exposure of nucleosomal DNA during remodeling and various other cellular processes, we still observe a clear anti-correlation between Cas9 activity and nucleosome occupancy in vivo, suggesting that the barrier to Cas9 target recognition exists even in a cellular environment. Indeed, the balance between nucleosome disruption, turnover, and repositioning in the cell leads to the average level of occupancy and positioning at each site observed by MNase-seq [19, 21]. Thus, our data suggest that it is likely the overall effect of this average nucleosome positioning that leads to the observed anti-correlation with Cas9/sgRNA activity. It is important to note, however, that there is likely a fundamental difference between applications that use dCas9 versus nuclease-active Cas9. Knock-down of transcription by CRISPRi likely requires persistent binding by dCas9 in order to continually block transcription, and would be largely ineffective during S-phase when transcription is globally shut down. In contrast, to make a genomic edit, Cas9 must succeed in cleaving DNA only once, and could potentially take advantage of nucleosome turnover during replication. The role these differences play in the dependence of Cas9 on nucleosome position is still not clear. We expect, however, that nucleosome position and occupancy will be of particular concern to applications that use the nuclease-dead Cas9 and require sustained binding. Future investigations into the role of cell cycle and nucleosome disruption may provide
an additional piece to our understanding of the mechanism of Cas9 in eukaryotic cells.
Furthermore, nucleosome organization represents only one aspect of eukaryotic chromatin, and
thus, our results contribute a first step in understanding and exploiting how chromatin affects
Cas9 activity in order to enable more sophisticated and precise rules for targeting Cas9.
Methods

Analysis of CRISPRi sgRNA activity

The K562 dCas9-KRAB-BFP cell line was obtained from [17] and had been constructed from K562 cells obtained from ATCC. The resulting cell line tested negative for mycoplasma (MycoAlert Kit, Lonza, Basel, Switzerland) in regular screenings, and cytogenetic profiling by array comparative genomic hybridization (not shown) closely matched previous characterizations of the K562 cell line [50]. Data from 30 published [17] and unpublished screens (M.L., B.A., B.B., C.Y.P., M.K., Y.C., J.F., and J.N., personal communication), conducted using the CRISPRi sgRNA library described in Gilbert 2014 [17] in K562 cells constitutively expressing dCas9-KRAB-BFP, were processed through a standardized pipeline adapted from Bassik et al. 2013 [25], and Kampmann et al. 2013 [51]. Briefly, sgRNA phenotypes were calculated as the log2 enrichment of sequencing read counts between two conditions (e.g. initial and final timepoints for growth screens, untreated and treated for drug/toxin screens) and normalized to cell doubling differences where appropriate. Most screens were conducted in duplicate, and sgRNA phenotypes from the duplicates were averaged. To determine hit genes, each gene was given an effect size (average of strongest 3 sgRNA phenotypes by absolute value) and a confidence value (Mann-Whitney p-value of all ~10 sgRNAs compared to negative controls), and hits were selected using a score that integrates effect size and statistical confidence (| effect Z score * log10 p-value | = 20 in any screen). For genes with multiple TSS, each TSS was analyzed separately and the gene was assigned the highest score. Finally, sgRNA phenotypes were extracted for hit genes from the screen in which the gene scored as a hit and normalized to the average of the strongest 3 phenotypes to generate the “sgRNA activity score.”

sgRNA positions were defined as the genomic coordinate of the 3’ G of the NGG PAM (all genomic coordinates referenced in this text are from hg19). TSS positions were determined from the FANTOM5 project annotation (Riken) (http://fantom.gsc.riken.jp/5/datafiles/phase1.3/extra/TSS_classifier/TSS_human.bed.gz; accessed March 2, 2015), using the downstream genomic coordinate of the corresponding “p1@gene” BED file entry. All local averages were calculated in 50bp windows centered around the indicated point. As sgRNA lengths including the PAM were ~24bp and position was calculated relative to PAM, a window size of 50bp captures all sgRNAs that directly neighbor the center point at either the 3’ or 5’ end. In order to quantify amplitude, the local averages were first smoothed using a low-pass Butterworth filter (N = 4, Wn = 0.03; SciPy signal processing module) and then peaks and troughs were calculated by determining the local maxima and
minima, respectively. As described for Figure 1D, Peak 0 was defined as the local maximum
closest to the TSS, peak 1 was defined as the next maximum downstream of peak 0, and
troughs were defined as the minima immediately downstream of the respective peaks.

**MNase-seq data analysis**

K562 MNase-seq data was obtained from the ENCODE consortium as processed
continuous signal data (BigWig file format; accession number ENCFF000VNN, Michael Snyder
lab, Stanford University). sgRNA target site signal was calculated as the average signal at all
positions between the 5’ end of the sgRNA and the 3’ end of the PAM.

**RNA-seq data analysis**

K562 RNA-seq data was obtained from the ENCODE consortium as transcript
quantifications (accession number ENCFF485YKK, Thomas Gingeras lab, Cold Spring Harbor
Laboratories). Genes were assigned expression levels in units of FPKM according to their
highest-expressed transcript.

**Linear regression**

Sequence score was calculated by passing the specified 30bp target site for each sgRNA
to the on_target_score_calculator.py script provided by Doench et al., 2015 (accessed October
9, 2015) [10]. This sequence score, MNase signal as calculated above, and sgRNA length
(base pairs in protospacer and PAM) were each compared to sgRNA activity scores by Pearson
correlation. Linear fits of the specified parameters were computed using multidimensional linear
regression (Sci-kit learn linear_model package), and correction of the activity scores for
sequence and length features was performed by subtracting the predicted scores based on the
combined fit.

**Ricin tiling screen**

Screens for ricin susceptibility were performed essentially as previously described [17].
Briefly, K562 cells constitutively expressing Cas9-BFP from an SFFV (spleen focus-forming
virus) promoter were transduced with our previously described pooled sgRNA tiling library
packaged into lentivirus for a multiplicity of infection below 1. Duplicate screens were infected
and subsequently treated independently. Infected cells were allowed to recover for 2 days, then
selected with 0.75μg/mL puromycin (Tocris) for 2 days, and finally allowed to recover from
puromycin treatment for 2 days. Cells were then cultured for 19 days and were either treated
with three pulses of 0.5ng/mL ricin administered for 24 hours and followed by re-suspension in fresh media, or passaged untreated. Genomic DNA was harvested from the endpoint untreated and treated samples and processed for high-throughput Illumina sequencing as previously described [17]. Screens were conducted at a minimum library coverage of 1,000 cells per sgRNA, and sequenced to a median depth of ~500 reads per sgRNA. Phenotypes were calculated as log, enrichments of read counts between untreated and treated conditions, normalized to cell doubling differences, and averaged between duplicates. Phenotype-signed Z scores were calculated by dividing all scores by the standard deviation of negative control phenotypes and then multiplying phenotypes by -1 for sgRNAs targeting genes shown to produce sensitizing phenotypes upon knockdown [25] such that positive values represent “expected” phenotypes.

**Protein Purification**

Mouse histones H2B(T115C), and H4 were recombinantly expressed in BL21(DE3)pLysS cells from expression plasmids gifted by Dr. Karolin Luger. Expression and purification of mH4 was conducted as previously described by the Luger lab [52, 53], while mH2B(T115C) was expressed and purified as described by the Cairns lab [54] with the following exception: after purification, histones were dialyzed against multiple changes of double distilled water and 1mM β-mercaptoethanol (BME) before lyophilizing for storage. Purified recombinant mouse (*Mus Musculus*) histones H2A and H3 were gifts from Dr. Karolin Luger.

The labeling mutant, mH2B(T115C) was fluorescently labeled with ~5-fold molar excess Alexa Fluor 647 C2 maleimide dye (Thermo Fisher Scientific, Waltham, MA, USA) as follows. 2 mg of lyophilized mH2B(T115C) was dissolved at 2 mg/mL in labeling buffer (7 M GuHCl, 50 mM Tris-HCl pH 7.5, 1 mM TCEP) and nutated at room temperature for 30 minutes. 1 mg of the dye was then dissolved in 50 µl of anhydrous DMF under Argon gas, and approximately half of the dye solution was slowly mixed with the dissolved mH2B(T115C) in the dark at room temperature to begin the labeling reaction. After nutating the reaction for 1 hour, the rest of the dissolved dye was slowly added and the reaction was moved to 4ºC overnight in the dark. In the morning, the reaction was quenched by adding over 100-fold molar excess of BME.

Histone octamers were refolded and purified as previously described [52]. Specifically, 110 nmoles each mH3 and mH4 were refolded with 130 nmoles each mH2A and mH2B(T115C). The resulting octamer was concentrated using a 10,000 MWCO Spin-X UF concentrator (Corning, Tewksbury, MA, USA), then purified on a Superdex 200 HR (10/30) column (GE Life Sciences, Pittsburgh, PA, USA) using an Akta Explorer FPLC (GE Life Sciences, Pittsburgh, PA, USA).
Sciences, Pittsburgh, PA, USA) at 0.2 mL/min. Selected fractions were concentrated, flash frozen, and stored at -80°C until use.

A new purification scheme was conceived to achieve exceptionally high purity (d)Cas9 (Figure 4 – figure supplement 3). Nuclease active *S. pyogenes* Cas9-HaloTag was recombinantly expressed and purified from BL21(DE3)pLysS-Rosetta cells (Novagen/EMD Millipore, Darmstadt, Germany) using the expression plasmid pET302-6His-wtCas9-Halo-NLS, while the nuclease dead *S. pyogenes* dCas9(D10A,H840A)-HaloTag was recombinantly expressed and purified from BL21(DE3)pLysS cells using the expression plasmid pET302-6His-dCas9-Halo [16]. Bacterial cultures were grown in Terrific Broth II (MP Biomedicals, Santa Ana, CA, USA) at 37°C until an OD$_{600}$ reached 0.4. Cultures were then transferred to an ice bath for ~15 minutes until an OD$_{600}$ reached 0.5, at which point expression was induced with 0.2mM IPTG, and the cultures were moved to an 18°C shaker for 16 hours. Cells were harvested at 3000xg for 20 minutes, then resuspended in lysis buffer (500 mM NaCl, 50 mM Hepes pH 7.6, 5% glycerol, 1% Triton X-100, 10 mM imidazole, 1 mM benzamidine, 2.3 µg/mL aprotinin, 0.5 mM PMSF, and 1 tablet per 50 mL of Protease Inhibitor Cocktail (Roche, Basel, Switzerland). Cells were lysed using sonication on ice at 50% duty cycle, power 8, 30 seconds on, 1 minute off (Misonix/ Qsonica, LLC, Newtown, CT, USA). The lysed cells were then ultracentrifuged at 4º, 40K rpms, for 40 minutes to remove cell debris. The supernatant was then allowed to bind to Ni-NTA agarose resin (Qiagen, Hilden, Germany) by nutating at 4°C for 30 minutes. The resin containing bound (d)Cas9 was poured into a mini column (Bio-Rad, Hercules, CA, U.S.A.) and washed with 10 column volumes (CV) of lysis buffer, and 5 CVs of 20 mM Imidazole buffer (same as lysis buffer but with 20 mM imidazole). Elution of the (d)Cas9 was achieved using 250 mM Imidazole buffer (same as lysis buffer but with 250 mM imidazole), and fractions were checked on an SDS-PAGE gel. Chosen fractions were pooled and diluted to a starting NaCl concentration of 200 mM using Buffer A (0 M NaCl, 50 mM Hepes pH 7.6, 5% glycerol, 1 mM DTT, 0.5 mM PMSF). A 5 mL HiTrap Q-HP (GE Life Sciences, Pittsburgh, PA, USA) and a 5 mL HiTrap SP-HP (GE Life Sciences, Pittsburgh, PA, USA) column were attached in tandem (Q first in line) and equilibrated on an Akta FPLC at 10% Buffer B (same as Buffer A except with 2 M NaCl). The pooled (d)Cas9 was filtered, then loaded onto the tandem columns at 2 mL/min. The columns were washed with 10% Buffer B until A$_{280}$ and A$_{260}$ returned to baseline, at which point the Q column was removed and (d)Cas9 was eluted from the SP column using a gradient from 10% to 50% Buffer B over 10 CVs. Fractions were chosen using SDS-PAGE, pooled, and dialyzed into storage buffer (200 mM NaCl, 50 mM Hepes pH 7.6, 5% glycerol, 1 mM DTT).
Aliquots were flash frozen in liquid nitrogen and stored at -80°C.

**Target DNA purification**

To make the fluorescent DNA used in this study, the 601 DNA sequence was amplified by PCR from plasmid pBSIISK+601 (parent 601 sequence plasmid gifted by K. Luger) with the primers listed below. Two different PCR products were produced; one labeling the Watson strand of the 601 sequence with a 5’ Cy3 dye (IDT, Coralville, IA, USA), and the other labeling the Crick strand. Large scale (~2 mL) PCR reactions using in-house produced Pfu DNA polymerase were ethanol precipitated before loading onto a 20cm x 20cm 12% native TBE-PAGE gel for DNA purification. The PCR product was cut out of the gel, and the gel slice was crushed and soaked in 0.3 M Sodium Acetate pH 5.2 with multiple buffer changes. The pooled extract was ethanol precipitated, resuspended in 10 mM Tris-HCl pH 7.5, 10 mM NaCl, and stored at -20º until use.

**Watson Primer pair:**

LW021: 5’- /5Cy3/ATCGGATGTATATATCTGACACGTGC -3’
LW022: 5’- ATCTGAGAATCCGGTGCCG -3’

**Crick Primer pair:**

LW025: 5’- /5Cy3/ATCTGAGAATCCGGTGCCGAG -3’
LW030: 5’- ATCGGATGTATATATCTGACACGTGC -3’

**sgRNA Production for in vitro experiments**

sgRNA was produced by T7 transcription of a short DNA oligo template. To create this template, two oligos, one containing the T7 promoter and DNA target sequence, and the other encoding the invariable scaffolding of the sgRNA, were annealed and filled in using a single PCR cycle. The DNA template was ethanol precipitated, then transcribed using the T7 Quick High Yield RNA Synthesis Kit (New England Biolabs, Ipswich, MA, USA) according to vendor instructions. The resulting sgRNAs were purified via 6% Urea-PAGE gel. The correct band was cut out and crushed and soaked in 0.3M NaOAc. After ethanol precipitation, and resuspension in double distilled water, aliquots were stored at -80ºC until use. SgRNAs targeting the 601 sequence included 20bp of complementarity 5’ to the targeted PAM, with the exception of PAM 4, which has room for only 19bp of complementarity. SgRNAs were used against PAMs 1, 3-15, and 17-
20. PAM 2 did not have a long enough target sequence available within the 601. The non-sense guide used in this study contains the target sequence ‘ACATGTTGATTTTCTGAAA’.

Template oligonucleotides:

5'- TTAATACGACTCACTATAGNNNNNNNNNNNNNNNNNGTTTTAGAGCTAGAAATAGC
-3'

5'- AAAAGCACGACTCGGTGCCACTTTTTCAAGTTGATAACGGACTAGCCTTATTTTAACTTG
CTATTTCTAGCTCTAAAAC -3'

Nucleosome assembly

Nucleosomes were assembled by salt gradient dialysis as previously described [52] using 50 µL home made dialysis buttons and ~1 µM DNA. The Watson and the Crick labeled 601 DNA were assembled into separate nucleosomes. The best DNA:octamer molar ratio for optimal nucleosome assembly was selected by titrating octamer. The most homogenous assembly as judged by 6% Native TBE-PAGE gel was chosen for future assays.

Cleavage and binding assays

(d)Cas9 cleavage and binding assays were both conducted in the same manner, exchanging dCas9 for wtCas9 during binding reactions. First, the complete ribonucleoprotein complex was formed by incubating 5-fold molar excess of the chosen sgRNA with (d)Cas9 at 37°C for 10 minutes. Next, the DNA substrate (either naked DNA or nucleosome) was added to 40 nM (5-fold less than Cas9), and the reactions were returned to 37°C for an hour. Reaction buffer contained 20 mM Hepes pH 7.5, 100 mM NaCl, 5 mM MgCl2, 1 mM EDTA, and importantly, 2.5 mg/mL insulin (Roche, Basel, Switzerland). Importantly, we found that including a nonspecific protein such as insulin prevents the nucleosomes from falling apart and getting lost to surfaces during binding and cleavage assays (Figure 4 – figure supplement 1B). Additionally, inclusion of insulin reduces nonspecific protein-protein interactions and aggregation. Binding reactions also contained 14.6% sucrose to allow direct loading onto a native gel without a loading buffer. At 1 hour, binding reactions were loaded onto a pre-run 6% native TBE-PAGE gel at 4°C, while cleavage reactions were stopped using a 5X stop buffer (250 mM EDTA, 2% SDS), and prepared to load onto a 10% Urea-PAGE gel by adding 2X loading buffer (95% formamide, 20% DMSO, 5 mM EDTA, 0.025% Orange G) and heating to
95°C for 5 minutes before snap cooling on ice. Binding and cleavage reactions were repeated at least twice each for most of the PAMs targeted. Gels were imaged using a PharosFX Plus (Bio-Rad, Hercules, CA, U.S.A.) and quantified using Image Lab (Bio-Rad, Hercules, CA, U.S.A.). For binding and cleavage reactions, gels were scanned in the Cy3 fluorescence channel, while Alexa Fluor 647 fluorescence was also imaged for binding reactions. For cleavage reactions, labeling of either the Watson or Crick strand was chosen such that the fluorophore was always attached to the strand complimentary to the sgRNA used.

Structure alignment and solvent accessibility

Molecular graphics and analyses were produced using the UCSF Chimera package (supported by NIGMS P41-GM103311) [55]. The crystal structures for the 601 nucleosome [43] (PDBID 3LZ0) and the Cas9-sgRNA-DNA ternary complex [44] (PDBID 4UN3) were superimposed by aligning the DNA path in both structures using MatchMaker and manual manipulation. The solvent accessible surface area of the DNA in the 601 nucleosome structure was computed using residue areaSAS.

Reconstitution of nucleosomes on a plasmid

Supercoiled plasmid was produced using a Qiagen Maxi prep kit (Hilden, Germany). Histone octamers were prepared from purified histones as described above. DNA and histone octamer were mixed at a weight ratio of 1:0.8 (DNA to octamer) at a concentration of 20µg of DNA in 1X TE with 2M NaCl. 50µL homemade dialysis buttons were used to dialyse the solution by a step-wise gradient in 500mL at room temperature. The gradient was as follows; 1.5 hr in 1M NaCl in TE, 2 hr in 0.8M NaCl in TE, 1.5 hr in 0.6M NaCl in TE, and 2 hr in 0.05M NaCl in TE. The resulting chromatin was stored at 4ºC until use.

Micrococcal nuclease digestion assays of chromatin assembly reactions

MNase (Sigma-Aldrich, St. Louis, USA) was resuspended at 200 units/mL in water and then diluted at 1:100, 1:500, or 1:1000 in a solution of 1X MNase reaction buffer (50mM Tris-HCl pH7.9 and 5mM CaCl$_2$ dihydrate), 0.1 mg/mL insulin, and 10% glycerol. Chromatinized plasmid assemblies in 1X MNase buffer were added to each MNase dilution. As a control, an equal amount of DNA as supercoiled plasmid was added to the lowest MNase dilution. Each reaction was incubated for 11 minutes at room temperature, then stopped in a solution with final concentrations of 20mM EGTA, 200mM NaCl, 1% SDS, 20 mg/mL GlycoBlue (Thermo Fisher Scientific, Waltham, MA, USA), and 13.4 mg/mL Proteinase K (Roche, Basel, Switzerland) and
incubated at 37°C for 30 minutes. DNA was precipitated with 0.5X Phenol/Chloroform/Isoamyl Alcohol solution, washed successively with 100% and 70% ethanol, and run on a 1.3% agarose gel with 0.5X TBE. 3μg 123bp DNA ladder (Thermo Fisher Scientific, Waltham, MA, USA) was used as a standard.

**Restriction enzyme accessibility assays**

Chromatin assembly or supercoiled plasmid with equivalent amounts of DNA were added to restriction enzyme master mixes for final concentrations of 1X CutSmart buffer (New England Biolabs, Ipswich, MA, USA), 6 ng/μL DNA, and 1:50 of the indicated restriction enzyme (New England Biolabs). Reactions were incubated for 1 hour at NEB recommended temperatures. Reactions were stopped in final concentrations of 15mM EDTA, 0.75% SDS, 150mM NaCl, and 15mg/mL Proteinase K and incubated at 37°C for 30 minutes. DNA was extracted with phenol/chloroform as above, resuspended in 1X CutSmart buffer, and linearized by digesting with 1:20 DraIII-HF (New England Biolabs) at 37°C for 1 hour. The full reactions were run on an agarose gel and quantified as above.

**Chromatin remodeling assays**

Reactions were set up with 40mM NaCl, 0.1 mg/mL BSA, 25mM Tris-Acetate pH 7.5, 10mM Mg-Acetate, 1mM DTT, 1.2mM rATP, and 6 ng/ μL chromatinized or supercoiled DNA. For restriction enzyme accessibility, the indicated restriction enzyme was added at 0.5 units/mL. For Cas9 accessibility, Cas9/sgRNA ribonucleoproteins assembled as described above were added at 15.62nM. Chromatin remodeling enzyme *yChd1* ΔNC (gift of Dr. Ashok Patel and Dr. Gregory Bowman) was added to the indicated reactions at 0.2 μM final concentration. Reactions were incubated at 27°C for 1 hour, then processed as with the restriction enzyme accessibility assay above.
Acknowledgments

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Supplemental Material

- Figure 2 – figure supplement 1
- Figure 3 – figure supplement 1
- Figure 4 – figure supplement 1
- Figure 4 – figure supplement 2
- Figure 4 – figure supplement 3
- Supplementary File 1
- Supplementary File 2


26. Canver MC, Smith EC, Sher F, Pinello L, Sanjana NE, Shalem O, Chen DD, Schuppg PG,


**Figure 1. CRISPRi activity anti-correlates with nucleosome occupancy**

(A) Workflow for generating CRISPRi activity scores from pooled genetic screens. The resulting values are distributed around 0 for inactive sgRNAs and around 1 for highly active sgRNAs. (B) Average CRISPRi activity and MNase-seq signal relative to the TSS. Green line represents average CRISPRi activity score of all sgRNAs within a 50bp window around each position. Purple line represents the K562 MNase-seq signal at each position averaged across all genes analyzed. (C) Average CRISPRi activity and MNase-seq signal for genes grouped by expression value. Genes were grouped into lower expression (light lines; N=240), higher expression (heavy lines; N=368), and medium expression (omitted for clarity; N=930), and analyzed as in (B). Expression values were obtained as fragments per kilobase million (FPKM) from ENCODE K562 RNA-seq data. Average activity at positions with fewer than 10 sgRNAs within the 50bp window was not calculated. (D) Quantification of the amplitude of periodic CRISPRi activity. Peak and trough coordinates were obtained by calculating the local maxima and minima of the activity traces from analyses in (B) and (C). Peak 0 was defined as the local maximum closest to the TSS, peak 1 was defined as the next maximum downstream of peak 0, and troughs were defined as the minima immediately downstream of the respective peaks. (E) CRISPRi activity and target site nucleosome occupancy for individual sgRNAs. Target site nucleosome occupancy was calculated from the average MNase-seq signal at all genomic coordinates across the length of the sgRNA protospacer and the protospacer adjacent motif (PAM). sgRNAs were then binned by the target site nucleosome occupancy, displayed as box-and-whisker plots, and labeled according to the minimum value within the bin except where indicated. P-values were calculated by a two-tailed Mann-Whitney test comparing each bin to the =0.0 bin. (F) Linear regression for CRISPRi activity. The squared Pearson correlation was calculated for the sgRNA activity scores compared to the indicated individual parameters (bars 1, 2, and 4) or linear fits of multiple parameters (bars 3 and 6). sgRNA activity scores were corrected for sequence and length features (bar 5) by subtracting the linear fit of those two features.

**Figure 2. Cas9 nuclease activity anti-correlates with nucleosome occupancy**

(A) Cas9 nuclease phenotypes and target site nucleosome occupancy for individual sgRNAs targeting CDS regions. Ricin susceptibility phenotypes for each sgRNA are expressed as a Z
score and are positive if the phenotype matches the expected knockdown phenotype. Target site nucleosome occupancy was calculated as in Figure 1E. sgRNAs were then binned by the target site nucleosome occupancy, displayed as box-and-whisker plots, and labeled according to the minimum value within the bin except where indicated. P-values were calculated by a two-tailed Mann-Whitney test comparing each bin to the =0.0 bin. (B) Linear regression for Cas9 nuclease phenotypes. The squared Pearson correlation was calculated for the sgRNA activity scores compared to the indicated individual parameters (bars 1, 2, and 4) or linear fits of multiple parameters (bars 3 and 6). sgRNA activity scores were corrected for sequence and length features (bar 5) by subtracting the linear fit of those two features.

Figure 2—figure supplement 1. Cas9 nuclease activity anti-correlates with nucleosome occupancy at all target sites

Cas9 nuclease phenotypes and target site nucleosome occupancy for individual sgRNAs. As in Figure 2A, but incorporating all CDS- and non-CDS-targeting sgRNAs in the ricin-susceptibility gene tiling library.

Figure 3. Cas9 nuclease activity is blocked by the presence of a nucleosome in vitro

(A) Schematic of the experimental setup for *in vitro* cleavage assays. Mononucleosomes were assembled by salt gradient dialysis of purified mouse histone octamer with the minimal nucleosome positioning sequence, Widom 601 (147bp). Prior to assembly, DNA was 5’-end-labeled with Cy3 and histone H2B was fluorescently labeled using an introduced cysteine (T115C) coupled to Alexa Fluor 647. Purified Halo-tagged Cas9 with *in vitro* transcribed sgRNAs were added to naked DNA or assembled nucleosomes, and DNA cleavage products were visualized using a denaturing gel imaged for Cy3-DNA fluorescence. See also Figure 3—figure supplement 1A. (B) Confirmation that fully occupied, well-positioned nucleosomes were assembled. After assembly using salt gradient dialysis, the produced nucleosomes were visualized using a native PAGE gel imaged in the Cy3 and Alexa Fluor 647 channels. Full incorporation of the free DNA into a nucleosome occupying a single position on the DNA is indicated by the presence of a single shifted band containing all of the detectable Cy3-DNA and Alexa Fluor 647-H2B signal. (C) Available PAMs and solvent accessibility of the 601 nucleosome positioning sequence. (Above) A schematic of the 601 sequence. The location of the histone octamer when assembled into a nucleosome is indicated by the gray oval. The location of PAMs within the double-stranded sequence are indicated with arrows spanning the 3
bp of the PAM, pointing in the 5’ to 3’ orientation of the NGG motif. The arrows are colored according to solvent accessibility at the center of the PAM as calculated from the crystal structure of the 601 nucleosome (PDBID 3LZ0). (Below) Crystal structure of the 601 nucleosome. For clarity, the surface area of the histones in the crystal structure has been made transparent. The DNA in the crystal structure is colored according to solvent accessibility using the same color scale as the PAMs above. Residues colored teal are less accessible, while residues colored fuchsia are more accessible by solvent. (D) Experimental conditions and timeline for cleavage assays. (E) Denaturing PAGE gel showing results of a cleavage assay targeting the indicated PAMs. Cleavage reactions containing naked DNA were loaded on the left half of the gel, while reactions containing nucleosomes were loaded on the right. The DNA was imaged via a Cy3 fluorophore attached to the 5’ end of the sgRNA-complimentary strand. A negative control was conducted with an sgRNA that had no sequence complementarity to the 601 sequence used (non-sense guide). See also Figure 3—figure supplement 1B,C for additional controls. (F) Quantification of the gel in (D). For each lane, percent cleavage was determined by calculating the percent of the total band signal corresponding to cleaved DNA.

Figure 3—figure supplement 1. HaloTagged Cas9 activity is indistinguishable from untagged Cas9

(A) Diagram of the CDS region in the Cas9 expression plasmid used in this study. (B) Cleavage assay comparing the HaloTagged Cas9 construct used in this study with an untagged Cas9 commercially purchased from New England Biolabs (NEB). Both forms of Cas9 were incubated with either naked DNA or the same DNA assembled into a nucleosome (see Figure 3A,C). A positive control used the restriction enzyme, StyI-HF (from NEB), to target a sequence at a location within the DNA known to be fully protected upon assembly into a nucleosome. Unless explicitly labeled as NEB, all constructs of Cas9 used were HaloTagged.

Figure 4. dCas9 is unable to bind nucleosomal DNA in vitro

(A) Schematic of the experimental setup for in vitro binding assays. Either naked DNA or assembled nucleosomes were incubated with catalytically dead Cas9 (d-SpyCas9-Halo), and binding was assessed by an electrophoretic mobility shift assay (EMSA). (B) Experimental conditions and timeline for binding assays. (C) A native PAGE gel showing the results of an EMSA in which dCas9 was targeted to the indicated PAMs on either naked or nucleosomal DNA. Gels were scanned for fluorescence from Cy3 on the DNA (green) and Alexa Fluor 647 on histone H2B (magenta). The two color channels were merged to identify the location of intact
nucleosomes (white). See also Figure 4—figure supplement 1 and 3 for reagent preparation and
experimental conditions, and Figure 4—figure supplement 2 for comparison with wtCas9 binding.
(D) Quantification of the gel in (C). Percent bound was determined by calculating the percent of
the total band signal in each lane corresponding to Cas9-bound target as determined by a shift
in mobility within the gel. (E) Summary of binding and cleavage results for each PAM tested.
The ability of Cas9 to bind or cleave nucleosomal DNA at a targeted PAM is displayed as
percent protection by the nucleosome, and was calculated by taking the ratio of binding or
cleavage on nucleosomal DNA to that on naked DNA. While only the largest error bars are
visible, replicates were performed for 15 of the 30 data points and are displayed with error bars
showing standard deviation from the mean. The DNA positions plotted correspond to the three
nucleotides of the targeted PAM. In order to compare percent protection from binding and
cleavage with solvent accessibility, the PAMs are overlaid with a plot of the solvent accessible
surface area for each strand (Watson or Crick) of DNA in the 601 nucleosome structure. The
percent protection at each PAM, as well as the solvent accessibility were plotted so that the 5'
end of each DNA strand begins at the left of the graph, where position 0 indicates the dyad. (F)
Structural assessment of the ability of Cas9 to bind nucleosomal DNA. Superposition of the
Cas9-guideRNA-DNA crystal structure [41] onto the 601 nucleosome crystal structure [40] was
achieved by alignment of the DNA path in both structures. (Left) To better view the alignment of
the Cas9 target DNA with the nucleosomal DNA, the Cas9 protein density has been removed.
(Right) After alignment of the DNA, inclusion of the Cas9 protein density reveals extensive steric
clashes with the histones. Histone surface area was made partially transparent to better reveal
the overlapping densities with Cas9.

Figure 4—figure supplement 1. Quality control
(A) H2B labeling is near complete. An SDS-PAGE gel after refolding and purifying the histone
octamer containing fully labeled versus poorly labeled H2B. The arrow indicates a mobility shift
of H2B corresponding to a fully labeled band. Each histone is present at equimolar ratios as
indicated by PageBlue protein stain. (B) Nucleosome stability in Cas9 binding and cleavage
assays is ensured by including a nonspecific protein in solution. On the left, a native PAGE gel
showing naked DNA and nucleosomes under Cas9 binding and cleavage reaction conditions
without a nonspecific protein in solution. On the right, the same conditions plus inclusion of a
nonspecific competitor protein, insulin.

Figure 4— figure supplement 2. DNA binding by dCas9 is also representative of wtCas9
binding
(A) Two native PAGE gels showing the results of an EMSA binding assay comparing dCas9 and wtCas9. Binding to naked DNA is shown in the gel on the left, while binding to nucleosomes is shown in the gel on the right. Both gels were imaged in the Cy3-DNA channel, while the gel on the right was also imaged in the Alexa Fluor 647 – H2B channel (inset image). Binding conditions for both dCas9 and wtCas9 were identical, and were as described in the methods.

(B) Quantification of the Native PAGE gel in (A).

Figure 4—figure supplement 3. (d)Cas9 purification strategy
(d)Cas9 purification scheme (left). SDS-PAGE gel stained with PageBlue (Life Technologies, Carlsbad, CA) showing the elution fractions from the HiTrap SP-HP column during purification of 6His-dCas9-HaloTag (right).

Figure 5. Nucleosomes within chromatinized DNA can block cleavage by Cas9, but a chromatin remodeling factor can restore Cas9 access.
(A) Schematic of the experimental setup. Supercoiled plasmid containing the 601 sequence inserted into a pBlueScript II SK (+) backbone (pBSIISK+601) was chromatinized by step gradient salt dialysis in the presence of histone octamer. Purified yeast Chd1 (yChd1) remodeling factor was used to test the effect of ATP-dependent remodeling factors on Cas9 access to nucleosomal DNA. (B) Quality assessment of the chromatinized plasmid used in this study. Titrated amounts of Micrococcal Nuclease (MNase) were incubated with the chromatinized plasmid and the resulting pattern of protection by assembled nucleosomes was visualized on a 1.3% agarose gel post-stained with ethidium bromide (EtBr). As a control, the supercoiled plasmid was also incubated with the lowest concentration of MNase. (C) A restriction enzyme accessibility assay (REAA) was used to assess the occupancy and position of the nucleosome assembled at the 601 sequence within the chromatinized plasmid. A panel of unique restriction enzyme sites spanning the 601 sequence were incubated with either the supercoiled plasmid, or the chromatinized plasmid. Cleavage was stopped, and protein was removed by incubation with proteinase K followed by phenol/chloroform/isoamyl alcohol extraction and ethanol precipitation. (Top) The resulting DNA was then linearized using DraIII, and the level of cleavage by the restriction enzyme panel was visualized on a 1% agarose gel post-stained with EtBr. The label “P” represents supercoiled plasmid, while “C” represents chromatinized plasmid. (Bottom right) The location of the restriction sites used are indicated on a diagram of the plasmid. (Bottom left) After quantification of the gel, the percent protection from cleavage experienced in the chromatinized plasmid was plotted versus the location of the
cleavage sites on the top strand of the 601 sequence. Experiment 1 refers to the REAA experiment shown in the gel above, while experiment 2 refers to the REAA experiment without remodeler shown in Figure 5F. The grey shading indicates the borders of the 601 sequence, and the grey oval represents the corresponding nucleosome. (D) REAA experiment using the frequent cutter, HaeIII, to assess the remodeling activity around the chromatinized plasmid by the purified yChd1 chromatin remodeler. The resulting banding patterns were visualized on a 1.5% agarose gel post-stained with EtBr. Low molecular weight fragments indicate a high degree of HaeIII accessibility, while higher weight bands indicate protection from digestion. (E) Diagram showing the location of the restriction enzyme cleavage sites and the PAMs targeted by Cas9/sgRNA in the experiment shown in F and G. (F) An accessibility assay was performed essentially as in C using either restriction enzymes or Cas9/sgRNAs in the presence or absence of the remodeler yChd1. The level of cleavage by the restriction enzyme panel (left) or Cas9/sgRNAs (right) was visualized on a 1.3% agarose gel post-stained with EtBr. A negative control was conducted with an sgRNA that had no sequence complementarity to the plasmid used (non-sense guide). The concentration of yChd1 used was the same as in panel D. (G) Quantification of the gels shown in F. Percent protection from cleavage of the chromatinized plasmid in the presence or absence of the chromatin remodeler was calculated relative to the percent cleavage in the corresponding supercoiled plasmid control, and plotted at the location of the restriction enzyme cleavage sites or the center of the PAMs with respect to the 601 dyad.
Supplementary Table titles

Supplementary Table S1. CRISPRi sgRNA annotations, activity scores, and target site MNase signal, related to Figure 1

Supplementary Table S2. Ricin tiling library sgRNA annotations, phenotype scores, and target site MNase signal, related to Figure 2
Figure 1

30 CRiSPRi Screens
18,380 sgRNAs targeting
1,539 high-confidence hit genes

sgRNA phenotype scores normalized
to top 3 sgRNAs within each gene
Activity score

CRISPRi activity
(average sgRNA activity score)

Average MNase signal

Distance to TSS (bp)

CRISPRi activity
(average sgRNA activity score)

Peak to trough difference
(sgRNA activity)

Linear regression for CRISPRi activity

Pearson correlation squared (R^2)
Figure 2

(A) Linear regression for CRISPR nuclease activity

(B) Pearson correlation squared ($R^2$)
Figure 4

A. Naked DNA + Alexa Fluor 647

B. molar ratio: target DNA Cas9 sgRNA

C. Naked DNA and Nucleosome

D. % Bound

E. % Protection

F. Nucleosome structure: histones

Gel Shift Assay (EMSA) native gel

D-SpyCas9-halo

Gel Shift Assay (EMSA) native gel

Widom 601

d-SpyCas9-halo

Crick Watson

solvent accessible surface area (Å²)

DNA position (nt)

% Protection

nucleosome structure:

601 DNA

Cas9 structure:

guide RNA

target DNA

Cas9