Massively Parallel Biophysical Analysis of CRISPR-Cas Complexes on Next Generation Sequencing Chips

Cheulhee Jung,1,8 John A. Hawkins,2,8 Stephen K. Jones, Jr.,1,8 Yibei Xiao,3 James R. Rybarski,1 Kaylee E. Dillard,1 Jeffrey Hussmann,2 Fatema A. Saifuddin,1 Cagri A. Savran,4 Andrew D. Ellington,1,5 Ailong Ke,3 William H. Press,2,6,7 and Ilya J. Finkelstein1,5,9,*
1Department of Molecular Biosciences and Institute for Cellular and Molecular Biology, The University of Texas at Austin, Austin, TX 78712, USA
2Institute for Computational Engineering and Science, The University of Texas at Austin, Austin, TX 78712, USA
3Department of Molecular Biology and Genetics, Cornell University, 253 Biotechnology Building, Ithaca, NY 14853, USA
4School of Mechanical Engineering, Birck Nanotechnology Center, Purdue University, 1205 West State Street, West Lafayette, IN 47907, USA
5Center for Systems and Synthetic Biology, The University of Texas at Austin, Austin, TX 78712, USA
6Institute for Cellular and Molecular Biology, The University of Texas at Austin, Austin, TX 78712, USA
7Department of Integrative Biology, The University of Texas at Austin, Austin, TX 78712, USA
8These authors contributed equally
9Lead Contact
*Correspondence: ifinkelstein@cm.utexas.edu
http://dx.doi.org/10.1016/j.cell.2017.05.044

SUMMARY

CRISPR-Cas nucleoproteins target foreign DNA via base pairing with a crRNA. However, a quantitative description of protein binding and nuclease activation at off-target DNA sequences remains elusive. Here, we describe a chip-hybridized association-mapping platform (CHAMP) that repurposes next-generation sequencing chips to simultaneously measure the interactions between proteins and ~10^7 unique DNA sequences. Using CHAMP, we provide the first comprehensive survey of DNA recognition by a type I-E CRISPR-Cas (Cascade) complex and Cas3 nuclease. Analysis of mutated target sequences and human genomic DNA reveal that Cascade recognizes an extended protospacer adjacent motif (PAM). Cascade recognizes DNA with a surprising 3-nt periodicity. The identity of the PAM and the PAM-proximal nucleotides control Cas3 recruitment by releasing the Cse1 subunit. These findings are used to develop a model for the biophysical constraints governing off-target DNA binding. CHAMP provides a framework for high-throughput, quantitative analysis of protein-DNA interactions on synthetic and genomic DNA.

INTRODUCTION

Clustered regularly interspaced palindromic repeats (CRISPR) and a CRISPR-associated (cas) operon provide bacteria and archaea with adaptive immunity against invading phages and other foreign nucleic acids (Sorek et al., 2013; Wright et al., 2016). To provide adaptive immunity, cells assemble a CRISPR RNA (crRNA)-guided nucleoprotein complex that recognizes specific foreign DNA targets. After target DNA recognition, a CRISPR-specific nuclease degrades the foreign nucleic acids. CRISPR systems also confer immunity against future infections by acquiring foreign DNA sequences as new spacers into the CRISPR locus (Amitai and Sorek, 2016). The ability to program and multiplex DNA/RNA targeting with diverse CRISPR-Cas systems has enabled leveraging of this microbial immune strategy for use in diverse biotechnological and medical applications (Hsu et al., 2014; Sander and Joung, 2014).

Intense interest in emerging CRISPR-Cas systems has driven the development of high-throughput methods for characterizing crRNA-guided binding and cleavage activities. Deep sequencing is frequently used to identify off-target binding (e.g., chromatin immunoprecipitation sequencing [ChIP-seq]) and cleavage (e.g., Digenome-Seq, BLESS) (Kim et al., 2015; O’Geen et al., 2015; Ran et al., 2015). Alternative strategies include in vivo fluorescent reporters for CRISPR-Cas protein binding or for the repair of resulting DNA double strand breaks (Kim et al., 2015; Wu et al., 2014). These methods frequently detect off-target binding and cleavage activities but have several limitations as well (reviewed for Cas9 in Bolukbasi et al., 2016). For example, both GFP production and DNA break repair efficiency may vary with cell-cycle stage and genomic context. Similarly, pull-down methods can be influenced by antibody quality, the degree of chemical crosslinking, and the chromatin state of a given target. Most of these strategies are also limited to identifying genomic off-target DNA cleavage sites thereby making it difficult to place the results in a quantitative biophysical framework. These methods aim to identify off-target sites in vivo but are not optimal for probing the molecular mechanisms underlying CRISPR-Cas activities.

Here, we describe a chip-hybridized association-mapping platform (CHAMP) for comprehensively profiling protein-nucleic
acid interactions on sequenced next generation sequencing (NGS) chips. The most widely adopted NGS sequencers fluorescently image clusters of DNA molecules covalently affixed to the surface of a microfluidic chip. CHAMP leverages these chips—that would normally be discarded after sequencing—to quantitatively measure protein-DNA interactions. Importantly, CHAMP does not require any hardware or software modifications to older NGS sequencers, as has been reported previously (Buenrostro et al., 2014; Tome et al., 2014; Nutiu et al., 2011). Instead, it uses modern and ubiquitous Illumina instruments to generate chips and sequencing data. Protein-DNA profiling experiments are then performed independently on a standard fluorescence microscope. In short, NGS sequencing provides information about the position and identities of millions of different DNA molecules, while the microscopy experiments quantitatively measure binding interactions of the proteins to a library of DNA molecules.

We used CHAMP to quantitatively profile interactions between the T. fusca type I-E CRISPR-Cas (Cascade) effector complex and a diverse library of genomic and synthetic target DNA molecules. Type I systems comprise ~50% of bacterial CRISPRs and have been used to control gene expression and cell fate (Luo et al., 2015; Makarova et al., 2015; Ciliandro and Voigt, 2015). CHAMP profiling revealed that Cascade recognizes an extended, 6-nt protospacer adjacent motif (PAM). Quantitative profiling of off-target DNA-binding sequences reveals a 3-nt periodicity in Cascade-DNA interactions, observed in synthesized libraries and human genomic DNA. Cas3 recruitment was sensitive to the identity of the PAM and PAM-proximal DNA-RNA mismatches, establishing a novel DNA-guided proofreading mechanism. These results were used to develop a predictive biophysical framework that accurately reproduced in vivo interference experiments. Using CHAMP, we also profiled CRISPR-Cas binding in human genomic DNA, paving the way for rapid and quantitative determination of off-target binding sites in patient-specific genomes. More broadly, this study provides an experimental and computational framework for comprehensive analysis of protein-DNA interactions for diverse CRISPR systems and other DNA-binding proteins on both synthetic and genomic DNA libraries.

RESULTS

A CHAMP for Profiling CRISPR-Cas DNA Interactions

CHAMP leverages used MiSeq chips that are generated via the Illumina sequencing pipeline (Figure 1A). At the end of a DNA sequencing run, the surfaces of these chips are decorated with ~20 million spatially registered, unique DNA clusters. CHAMP uses high-throughput fluorescence imaging to measure the association between fluorescently labeled protein complexes and each DNA cluster (Figure 1A). The MiSeq sequencer is ubiquitous in nearly all NGS cores and genomics labs, produces long (~300 bp) reads, and the MiSeq chips also contain integrated microfluidic ports. To prepare chips for CHAMP, the DNA clusters are first regenerated to remove any fluorescent nucleotides that can otherwise confound imaging (Figure S1). A fluorescent oligonucleotide primer is then hybridized to a subset of the DNA clusters and used as an alignment marker in the downstream image-processing pipeline (Figure 1A). Next, fluorescently labeled proteins are incubated in the chip and imaged using a total internal reflection fluorescence (TIRF) microscope. The images are then analyzed using the CHAMP software pipeline, which maps each fluorescent cluster to the underlying DNA sequence, as reported by the Illumina sequencer (Figure S2; STAR Methods). CHAMP’s strength lies in its platform independence and its software pipeline, which quantifies protein association with each DNA sequence (Figure 1; STAR Methods).

Using CHAMP, we profiled the PAM specificity and off-target binding affinity of the thermophilic T. fusca type I-E CRISPR-Cas (Cascade) complex (Figure 1B). Experiments were carried out on regenerated MiSeq chips that contained a synthetic oligonucleotide library encoding substitutions within the PAM and the target DNA sequence. DNA binding was imaged at 11 Cascade concentrations ranging from 63 pM to 630 nM (see the STAR Methods). At each concentration, the thermophilic Cascade complex was first incubated in the chip at 60 °C to promote DNA binding. Next, unbound complexes were flushed out of the chip, and DNA-bound Cascade was rapidly cooled to room temperature and labeled in situ with fluorescent anti-FLAG antibodies (Figures 1A and S1). The T. fusca Cascade complex included a triple FLAG epitope on the C terminus of the Cas6 subunit. The epitope tag did not alter DNA binding by the T. fusca Cascade, as reported previously for the E. coli complex (Szczelkun et al., 2014). We did not observe significant Cascade loss or photobleaching during image collection (~15 min per protein concentration) (Figure S2F). Apparent Kd values were determined by fitting the fluorescence intensities of each DNA cluster at the 11 Cascade concentrations to the Hill equation (Figure 1D; STAR Methods). Non-specific DNA binding was observed via a random DNA sequence that was also included in the chip. This negative control sequence had an apparent Kd that was lower than our highest measured concentration (Figure S2D, dashed curve). We used these fits to define apparent binding affinity (ABA), the difference in apparent ΔG between the negative control sequence and a sequence of interest. Positive values indicate stronger binding, and negative values were discarded as non-specific DNA binding. DNA sequences with at least five unique fluorescent clusters were included in the analysis. This cutoff established an average error of ~0.2 kBT for the apparent binding affinity, where kBT is the Boltzmann constant and T is the temperature (Figure S4B). We sequenced ~16 million target DNA sequences, giving complete coverage of all possible 6-nt PAM variants, as well as all single- and double-nucleotide substitutions along the entire target DNA (Figures 1E and 1F). Paired-end reads of linearly amplified synthetic oligonucleotide libraries were used to minimize biases and errors from library construction, synthesis, and sequencing. To avoid chip-specific biases, we performed experiments on two independent MiSeq chips, which recapitulated the measured ABAs (r = 0.88) (Figure 1G). This CHAMP dataset resulted in ~36,000 unique DNA sequences with ABAs that were above the non-specific DNA binding threshold (Figure 1H). With this dataset, we next set out to define the principles guiding Cascade-DNA interactions.

Quantitative Profiling of the Protoscaler Adjacent Motif

In all CRISPR-Cas systems, the protoscaler adjacent motif (PAM) flanks target DNA that is complementary to the crRNA.
The PAM is crucial for facilitating interrogation of the target DNA by the Cascade complex. Diverse PAMs can also bias CRISPR-Cas systems toward DNA degradation (interference) or spacer acquisition (adaptive immunity) (Heler et al., 2015; Horvath et al., 2008; Marraffini and Sontheimer, 2010). Early studies proposed that Cascade recognizes a 3-nt PAM (Marraffini, 2015; Semenova et al., 2011). However, recent structural and sequencing studies of the E. coli Cascade complex suggested...
that Cse1 is sensitive to an extended PAM (Hayes et al., 2016; Leenay et al., 2016). Thus, we used CHAMP to determine the apparent binding affinity of Cascade toward 6-nt PAMs when the target DNA is fully complementary to the corresponding crRNA (Figure 2A).

CHAMP profiling of all 4,096 unique 6-nt PAMs resulted in 950 sequences that had a non-zero ABA. In order visualize the complete set of all PAM preferences, we adapted sequence specificity landscapes (called PAM landscapes here, Figure 2B) (Carlson et al., 2010). The PAM landscape displays all PAM-dependent ABAs as a series of concentric rings (Figure 2B, top). The highest-affinity sequence for the first three PAM positions (A/A/A/G) is included in the center of the concentric rings. This innermost dataset displays the ABAs for all 6-nt PAM sequences that contain a perfect match to the highest affinity 3-nt “minimal” PAM (N/N/N/A/A/A/G) for T. fusca Cascade: 64 unique sequences). The height and color of each bar on the individual rings corresponds to the ABA. A gray line above each peak represents the SD of each measurement, as determined by bootstrap analysis. The vertical bars are sorted
from the highest to lowest affinity sequences for each minimal PAM. When paired with AAG, variation in the –6 to –4 position contributes minimally to the ABA. The next ring in the landscape shows ABAs for 6-nt PAMs that vary from A₅A₂G₁ by a single nucleotide in the first three positions (e.g., N₃N₂N₁C₃A₂G₁). The final ring shows PAMs that vary from A₅A₂G₁ by 2 nt (e.g., N₃N₂N₁C₃A₂G₁). We did not detect any measurable binding affinity to PAMs with three substitutions relative to A₅A₂G₁, and these PAMs are thus not displayed in Figure 2B. This representation gives a high-level overview of the entire PAM sequence space, reducing the high-dimensionality of CHAMP datasets for rapidly comparing the binding affinity to various PAMs.

We determined the relative importance of each base in the extended PAM by computing the maximum change in the ABA when only that base was varied (Figure 2C). For example, a single data point in the violin plot for the PAM₄₋₂ position plots the maximum difference in ABAs for the four A₅A₄A₃A₂N₁C₂G₁ PAMs. The violin plot extends this comparison for all possible PAMs at each of the six PAM positions and shows the maximum effects of a single base change in varying PAM contexts. The PAM₄₋₂ position is the most critical for defining the highest affinity T.fusca PAM. In contrast, the closely related E.coli Cascade complex has promiscuous recognition at the PAM₄₋₂ position (Hayes et al., 2016). Both PAM₁₋₁ and PAM₃₋₅ make similar contributions to the ABA. Subsequent positions in the extended PAM typically contribute less to ABA (PAM₄₋₂ > PAM₁₋₁ = PAM₃₋₅ > PAM₄₋₄ > PAM₅₋₅ > PAM₆₋₃). These results also highlight that PAMs with intermediate ABAs are the most sensitive to the identity of nucleotide positions –4 to –6. For example, for NNNN, the ABA increases over 60% from 2.7 kBT for GGAGAG to 4.4 kBT for CACAG. Our data highlight additional sequence preferences, including enrichment of C₆ and G₆ in the highest affinity extended PAMs. The PAM₄₋₄ position is likely decoded by direct interactions with Cse1, as reported for the E.coli Cascade structure (Hayes et al., 2016). Contributions of PAM₅₋₅ and PAM₆₋₆ may be due to indirect effects such as changes in the shape of the DNA minor groove.

We next compared the CHAMP results with in vitro electrophoretic mobility shift assays (EMSAs) and in vivo interference assays. EMSAs showed excellent agreement with the CHAMP datasets (r = 0.96) over three orders of magnitude in concentration (Figures S3A and S3B). As expected, purified Cascade complex lacks the Cse1 subunit did not exhibit any target DNA binding via EMSAs or CHAMP (data not shown). Next, we carried out a plasmid-based interference assay and compared the results to those obtained via CHAMP for a variety of PAM sequences. In this assay, T.fusca Cascade, along with Cas3 nuclease, is induced in cells that also harbor a target plasmid that is degraded by the Cascade-Cas3 complex (Figure 2D). After a brief outgrowth without antibiotics, interference efficiency is scored as the relative number of antibiotic-resistant colonies (Huo et al., 2014). The results showed a strong correlation (r = 0.89), indicating that CHAMP-derived binding affinities are also predictive of interference activity in vivo (Figure 2E). Moreover, our observations also help to explain how T.fusca avoids self-targeting its two type I-E CRISPR loci. The first locus has a repeat that contains a 5’-A₅₃C₃₂G₁ sequence adjacent to the CRISPR spacer elements, whereas the second repeat is 5’-T₄C₃₂C₁. Here, we show that these sequences strongly disfavor Cascade binding and thus limit auto-immunity at the CRISPR locus. In sum, CHAMP profiling recapitulates DNA binding affinities measured via EMSAs in vitro and is highly correlated with in vivo interference activity.

### Profiling Off-Target CRISPR-Cas DNA Binding on Synthetic DNA Libraries

To delineate the sequence determinants that influence Cascade-DNA interactions, we next analyzed the ABA for all DNA molecules with single or double substitutions along a 35-nt region that includes the first three positions of the PAM and the target DNA (Figure 3). CHAMP profiling yielded information for all possible single-base substitutions with an average 3,000-fold coverage (Figure 3A). As expected, substitutions in the PAM region reduce the ABA substantially, with the second position being most critical for Cascade binding (Figure 3A). Prior structural and biochemical studies have established that every sixth nucleotide is not paired with the crRNA and flipped out in the type I-E Cascade-DNA complex (Hayes et al., 2016; Jackson et al., 2014; Zhao et al., 2014). A clear signature for these flipped-out base positions is also evident in the CHAMP profiling data (Figure 3A). Surprisingly, Cascade revealed that Cascade affinity was increased when thymidine replaced the complimentary cytosine as the third flipped-out base (position 18). We confirmed a preference for thymidines over cytosines at the flipped-out positions via EMSA assays (Figures S3C–S3E). In line with these observations, a structural study proposed that flipped-out bases interact with a molecular relay of Cse2-encoded arginines (van Erp et al., 2015). Taken together, these results suggest that flipped-out and mismatched DNA bases likely interact with Cascade, further stabilizing partially mismatched crRNA-DNA complexes during both interference and primed acquisition.

We developed a simple model to better quantify how substitutions along the PAM and the target DNA affect Cascade binding (Figures 3B–3D). This model considers a position-dependent penalty for all single base substitutions (Figure 3C) and a position-independent weight that accounts for the identities of each target and substituted base (Figure 3D). This model has fewer parameters than position weight matrices (Stormo and Zhao, 2010), but nonetheless described ~90% of the variance in the experimental data (Figure 3B). To further constrain this model, we acquired a second CHAMP dataset with a second crRNA-Cascade complex targeting a different DNA sequence. The model accurately described both independent CHAMP datasets acquired with two different crRNAs and corresponding DNA libraries (r = 0.92) (Figure 3B). Analysis of the position-specific penalties clearly highlights the importance of the PAM, as well as the PAM-proximal nucleotides (i.e., seed region) in modulating the affinity of Cascade for DNA. The overall substitution penalties decrease with increasing distance from the PAM (Figure 3C). This pattern has been recently observed for other CRISPR-Cas systems, (Hsu et al., 2013) and likely reflects the initiation and directional formation of an R-loop proceeding from the seed region (Blosser et al., 2015; Rutkuskas et al., 2019).
Figure 3. Comprehensive Profiling of Cascade-DNA Interactions

(A) The change in ABA for all possible single-base substitutions along the minimal PAM and the target DNA. Negative values indicate a reduced ABA relative to the best PAM and perfectly paired DNA target. Error bars, SD obtained via bootstrapping.

(B) CHAMP profiling was performed on two distinct DNA libraries (blue and red dots). The resulting data were used to construct a minimal binding model shown in (C) and (D) that accurately describes the data obtained from both CHAMP datasets. (C and D) Position-dependent substitution penalties (C) and position-independent nucleotide preferences (D) obtained from the binding model.

(E) The change in ABA for all dinucleotide substitutions. The triangular matrix represents the average of CHAMP measurements acquired on two independent chips. The PAM is in the upper left-hand corner. Gray regions indicate insufficient data. As an example, the inset shows an enlarged 3 x 3 dinucleotide substitution matrix showing all possible substitutions for positions A12 and C9.

(F) A schematic representation of T. fusca Cascade highlighting contribution of PAM positions 1 to 6 and the 3-nt periodicity.

(G) Models representing the 3-nt periodicity imposed by the protruding Cas7 finger (residues 193–211) (top) and steric clash with adjacent amino acids (R19, M173, D183, and K271; transparent DNA for clarity) (bottom) based on E. coli Cascade (Hayes et al., 2016).

See also Figure S3.
We also analyzed the ABAs for all double nucleotide substitutions along the same 35-nt PAM and target DNA region (Figure 3E). The data highlight the importance of the PAM, position for controlling Cascade binding, as well as the synergistic effects of having any two flipped out bases. In the seed region, single substitutions are already poorly tolerated and reduce ABAs significantly. Therefore, a second mismatch in the seed reduces the ABAs to DNA-binding levels that are like non-specific DNA, while a second mismatch in PAM-distal positions are often tolerated. Two substitutions in the PAM-distal sequence only marginally destabilized the Cascade-DNA complex.

Surprisingly, our data and model also reveal an additional periodicity in base-substitution penalties centered between the flipped-out bases (Figures 3C and 3E). This periodicity results in an overall decrease in mismatch penalties every three nucleotides (e.g., at +3, +6, +9, etc.). A close inspection of the high-resolution E. coli Cascade structure reveals that every third base pair is puckered due to steric clashes between the RNA-DNA duplex and several residues in the Cas7 subunit (Figures 3F and 3G). Six repeats of the Cas7 subunits polymerize along the crRNA to form the backbone of the Cascade complex. These subunits are likely to give rise to the 3-nucleotide periodicity observed in our model and dinucleotide ABA data. Moreover, these residues are highly conserved among divergent type I-E CRISPR-Cas systems, suggesting that they may play a role in Cascade assembly. Overall, our results highlight an unanticipated 3-nt periodicity in Cascade-DNA binding penalties that reduce the overall fidelity of RNA-DNA binding.

Profilng Off-Target CRISPR-Cas Binding in Human Genomic DNA

CHAMP uses a standard Illumina workflow and is immediately compatible with any nucleic acid library, including those derived from genomic preparations. We therefore extended CHAMP to profile CRISPR-Cas binding on human genomic DNA (Figure 4). To enrich for gene-coding regions, exome capture was used in conjunction with paired-end sequencing on an Illumina MiSeq sequencer (Figure 4A). The resulting sequenced MiSeq chip had an average 11-fold coverage for 17,862 human protein-coding regions from 7 million unique high-quality DNA clusters (Figure S4A). This MiSeq chip was used to quantitatively assay off-target CRISPR-Cas binding. Remarkably, 37 genes showed at least one high-affinity CRISPR binding site (defined as ABAs >4 kBT) and ~200 genes showed moderate-affinity ABAs (>3 kBT). The precision of the off-target DNA sequence is defined by both the length distribution of the sheared genome fragments and the depth of coverage at each position (Figures 4B and S4B). Nonetheless, most genes harboring off-target sites showed a single, well-resolved ~200 bp-wide peak (Figure 4C).

The peaks with the highest ABAs represent genomic high-affinity off-target DNA binding sites. A subset of these peaks may also represent a combination of two lower affinity binding sites that are closer than our nominal resolution of 210 bp (Figure S4B). Nonetheless, a logo analysis of all peaks with ABAs >3 kBT revealed a consensus sequence that matches closely with the expected critical determinants of off-target binding observed in our synthetic DNA libraries (Figure 4D). The consensus off-target site had a strong preference for an AAG PAM, with the second adenine giving the strongest signal (compare to Figure 2C). Second, off-target sites were highly enriched for the first eight base pairs of the target DNA sequence. One notable exception is the flipped-out base in the sixth position, which does not base pair with the crRNA (also see Figure 3). Consistent with binding data obtained from synthetic DNA arrays (Figure 3), mismatches are also tolerated at the third base, which has reduced base-pairing with the crRNA. This data also highlights that an 8-nt PAM-proximal “seed” region is necessary for efficient binding, as has been previously observed in vitro and via in vivo interference assays (Fineran et al., 2014; Semo-nova et al., 2011; Wiedenheft et al., 2011; Xue et al., 2015).

Here, we demonstrate that CHAMP can profile off-target CRISPR-Cas binding sites in human genomic DNA, paving the way for rapid and quantitative profiling of off-target binding sites in patient-specific genomes.

Cas3 Recruitment Requires Perfect Base Pairing Near the PAM

CHAMP profiling revealed pervasive off-target DNA binding by Cascade. Therefore, we reasoned that subsequent binding of the Cas3 nuclease may constitute an additional sequence-dependent proofreading mechanism. We investigated this possibility with three-color CHAMP experiments that measured the degree of Cas3 recruitment to DNA-bound Cascade (Figure 5A). Fluorescent Cascade, Cas3, and alignment markers were spectrally separated into three distinct emission channels. After adding alignment markers, Cascade was introduced into the chips at a sufficiently high concentration to bind most DNA clusters that were partially complementary to the crRNA. Next, a saturating concentration of Cas3 was introduced into the same chip and CHAMP data were acquired (Figure 5B). To prevent Cas3-dependent DNA degradation, these assays were conducted with a buffer containing 1 mM AMP-PNP and lacking CoCl₂ (see the STAR Methods). While most clusters had a linear correlation between Cascade and Cas3 signals, a subset of the clusters deviated from this linear correlation with a reduced Cas3 fluorescence (Figure 5B, inset). As expected, we did not see any Cas3 binding to the DNA clusters when Cascade was omitted from the chip, or on clusters that did not bind Cascade. These results suggest that Cas3 is recruited to Cascade in a DNA-sequence-dependent manner.

We analyzed ~646,000 DNA clusters representing 10,810 unique DNA sequences to determine the requirements for efficient Cas3 recruitment. This dataset represented all extended PAM and single-nucleotide substitution variants, as well as 94% of double-nucleotide substitution variants along the target DNA sequence (Figure 1F). Approximately 450 DNA sequences showed a reduced ratio of Cas3 to Cascade fluorescent intensities relative to that of the fully complementary DNA target sequence. To better understand why Cas3 was not recruited at the same level to all DNA clusters, we focused on DNA sequences with single nucleotide substitutions along the PAM and the target DNA (Figure 5C). Comparing the Cas3 and Cascade fluorescent signals indicated that most DNA sequences fell on a diagonal line that indicates stoichiometric Cas3 recruitment, while those below the diagonal line indicate sub-stoichiometric Cas3 to Cascade ratios. As expected, we did not observe any points above the diagonal (Figure 5C).
Cas3 recruitment was partially compromised at nearly all non-AAG PAMs, as well as for target DNAs with a substitution in the first three PAM-proximal positions (Figure 5C). Using this information, we computed how sequence-dependent substitutions in the target DNA impact Cas3 recruitment. These results are expressed as a Cas3 recruitment penalty relative to expected stoichiometric binding (Figure 5D). Surprisingly, our results revealed that mismatches in PAM+1 and +1 target positions strongly compromised Cas3 recruitment (Figure 5D). These data implicate the PAM, as well as the first few nucleotides in the seed region, as critical for Cas3 binding to a Cascade-DNA complex.

Sequence-Specific Loss of Cse1 Decreases the Cascade Interference Efficiency

We next used EMSAs and nuclease assays to further determine the mechanism of DNA-guided Cas3 recruitment (Figure 6). Cascade readily binds target DNA containing an A₃A₂G₁ PAM. Surprisingly, the Cascade-DNA complex migrated as a faster mobility species when either this PAM was changed or when the +1 DNA position was mismatched relative to the crRNA (Figure 6A). Indeed, a DNA:crRNA mismatch in the +1 position converted 80% of the Cascade complexes to the faster-migrating species. These effects were additive, as changing the PAM and the +1 position simultaneously resulted in nearly 100% of the faster-migrating sub-complex. Consistent with previous studies, we confirmed that this faster migrating species represents Cascade lacking the Cse1 subunit (Figure S5) (Huo et al., 2014; Jore et al., 2011). Adding a large excess of free Cse1 could restore the mobility back to that of a complete Cascade complex (Figure S5). Cse1 physically interacts with Cas3 and loads the nuclease onto the target DNA (Huo et al., 2014). Adding excess Cas3 resulted in a super-shift, but only
when Cse1 was part of the Cascade complex (Figures 6A and 6B). As expected, impaired Cas3 recruitment also reduced Cas3 nuclease activity when ATP and Co^{2+} were added to the reaction mixtures (Figures 6C and 6D). Consistent with these in vitro studies, disrupting either the PAM or first few seed nucleotides also caused strong reduction in the plasmid-based in vivo interference assays (Figure 6E). These results reveal that DNA sequence-specific loss of Cse1 abrogates Cas3 recruitment and provides an additional proofreading mechanism for modulating CRISPR interference.

**DISCUSSION**

CHAMP repurposes sequenced and discarded chips from modern next-generation Illumina sequencers for high-throughput association profiling of proteins to nucleic acids. A key difference between CHAMP and prior NGS-based approaches is that it does not require any hardware or software modifications to discontinued Illumina sequencers (Nutiu et al., 2011; Tome et al., 2014; Buenrostro et al., 2014). In CHAMP, all association-profiling experiments are carried out on sequenced MiSeq chips and imaged in a conventional TIRF microscope. CHAMP’s computational strategy uses phiX clusters as alignment markers to align the spatial information obtained via Illumina sequencing with the fluorescent association profiling experiments. This strategy offers three key advantages over previous approaches. First, using a conventional fluorescence microscope opens new experimental configurations, including multi-color co-localization and time-dependent kinetic experiments. The excitation and emission optics can also be readily adapted for FRET (Figure S6) and other advanced imaging modalities. Second, complete fluidic access to the chip allows addition of other protein components during a biochemical reaction. Third, the computational strategy for aligning sequencer outputs to fluorescent datasets is applicable to all modern Illumina sequencers, including the MiSeq, NextSeq, and HiSeq platforms. Indeed, we also used the CHAMP imaging and bioinformatics pipeline to regenerate, image, and spatially align the DNA clusters in a HiSeq flowcell (Figure S6), providing an avenue for massively parallel profiling of protein-nucleic acid interactions on both synthetic libraries and entire genomes. Future extensions will leverage on-chip transcription and translation (e.g., ribosome
display) to facilitate high-throughput studies of RNA or peptide association landscapes. These studies will permit quantitative biophysical studies of diverse protein-nucleic acid interactions.

**Cascade Interrogates an Extended PAM and Recognizes Mismatched DNA Targets**

Using CHAMP, we profiled the biophysical properties governing interactions between target DNA and the type I-E CRISPR-Cas effector complex. Our findings reveal the biophysical parameters governing PAM recognition and DNA-binding at partially complementary target DNAs. *T. fusca* Cascade first identifies an extended PAM, possibly via hydrogen bonds with the PAM-5 nucleotide as suggested by a recent high-resolution structure of the *E. coli* Cascade-DNA complex (Hayes et al., 2016). Further readout of the PAM-4 and PAM-6 positions may be mediated by indirect effects, such as changes in the major and minor groove widths at the PAM-proximal bases. These results are also broadly consistent with recent plasmid-based PAM-profiling experiments, which highlighted that diverse CRISPR-Cas systems—including the *E. coli* type I-E Cascade—all decode an extended PAM (Leenay et al., 2016).

Following PAM recognition and target DNA unwinding, an R-loop extends along the complementary target DNA. Using CHAMP, we probed the effects of multiple sequence substitutions on Cascade-DNA interactions. In addition to identifying the importance of the PAM, “seed,” and flipped-out bases, our analysis and modeling revealed an unanticipated 3-nt periodic interaction that reduced the relative penalty for DNA-RNA mismatches at these positions. A re-analysis of previously reported *E. coli* Cascade plasmid interference assays also shows the same 3-nt periodicity (Fineran et al., 2014). Here, we propose that this is likely a general structural feature shared by other type I-E systems and that it likely arises due to a steric clash between base pairs in the R-loop and residues in each of the six Cas7 subunits. The crRNA is required for assembly of the *E. coli* Cascade complex (Zhao et al., 2014), and we speculate...
that these periodic contacts allow the crRNA to act as a scaffold during Cascade assembly. The crRNA is held in a conformation that maximizes interaction with the target DNA, possibly avoiding secondary structure formation by targets, as has been demonstrated in other RNA-guided nucleases (Jiang et al., 2015; Schirle and MacRae, 2012; Zhao et al., 2014). This periodic mismatch tolerance was also confirmed at off-target sites mapped to the human exome, further highlighting the importance of quantitatively mapping the influence of mismatches on CRISPR-DNA interactions with both synthetic and genomic DNA substrates.

**A DNA Sequence-Dependent Mechanism Underlies Cse1 Loss and CRISPR Interference**

By performing multi-color CHAMP imaging, we uncovered that Cas3 recruitment is dependent on the identity of the PAM, as well as perfect complementarity between crRNA and DNA in the +1 to +3 positions (Figure 6). These nucleotides interact with the Cse1 subunit of the Cascade complex. EMSAs and in vitro nuclease assays revealed that T. fusca Cse1 dissociates from Cascade at intermediate PAMs or when there are mismatches between the crRNA and the first three nucleotides of the target DNA. The functional significance of this position was further confirmed with in vivo plasmid interference assays and also recapitulates previously published in vivo interference results with the E. coli Cascade complex (Fineran et al., 2014).

In addition to identifying foreign DNAs, Cascade and Cas3 also promote primed spacer acquisition, where additional spacers are rapidly acquired from foreign DNAs that already contain a spacer in the CRISPR locus. Spacer acquisition requires the Cas1-Cas2 protein complex, which binds protospacer DNA and uses its integrase activity to insert the protospacer within the CRISPR array. Cascade can promote target acquisition at both perfectly matched spacers and mismatch-containing spacers that do not elicit strong interference (Sashital et al., 2012; Semenova et al., 2016; Staals et al., 2016; Xue et al., 2016). Conformational control of the Cse1 subunit is emerging as a key paradigm for recruiting Cas1-Cas2 and redirecting the Cascade-Cas3 complex toward primed acquisition (Xue et al., 2016). Here, we speculate that Cse1 undergoes a DNA-sequence-dependent conformational change that renders it labile in the absence of Cas1-Cas2 complex. Future CHAMP studies with fluorescent Cas1-Cas2 and FRET-reporters of Cse1 conformational state will shed light on the mechanisms and sequence requirements for primed spacer acquisition.

**Leveraging CHAMP for Mapping Protein-Nucleic Acid Interactions on Human Genomes**

Because CHAMP uses the standard Illumina workflow, it is immediately compatible with any nucleic acid library, including synthetic DNA, RNA, or genomic preparations. However, mapping CRISPR-DNA interactions on sequenced genomes presents additional computational challenges due to the random shearing lengths and uneven sequencing coverage. To address this challenge, we developed a bioinformatics pipeline that successfully identified off-target binding sites within a human exome with an ~200 bp effective resolution at an average 11-fold coverage depth. Higher resolution mapping can be readily achieved by shorter DNA fragments and greater sequencing coverage. Thus, CHAMP can be used to probe off-target CRISPR-Cas binding in any genome prior to performing genome-editing. Further extensions will allow direct observation of both binding and cleavage at these off-target sites. As CRISPR-Cas systems continue to be developed for human gene modification, CHAMP and similar methods may become useful tools for rapidly and quantitatively assaying target specificity on individual patient’s genomes.

**STAR METHODS**

Detailed methods are provided in the online version of this paper and include the following:

- **KEY RESOURCES TABLE**
- **CONTACT FOR REAGENT AND RESOURCE SHARING**
- **METHOD DETAILS**
  - Protein Cloning and Purification
  - Antibodies
  - DNA oligonucleotides libraries
  - Exome preparation and sequencing
  - Chip regeneration and addition of alignment markers
  - Fluorescence microscopy
  - CHAMP assays
  - Electrophoretic mobility shift assay (EMSA)
  - Cas3 nuclease assays
  - Plasmid loss assays
  - Computational Methods
  - Data Analysis
- **DATA AND SOFTWARE AVAILABILITY**

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes six figures, one table, and two data files and can be found with this article online at http://dx.doi.org/10.1016/j.cell.2017.05.044.

An audio PaperClip is available at http://dx.doi.org/10.1016/j.cell.2017.05.044#mmc4.

**AUTHOR CONTRIBUTIONS**


**ACKNOWLEDGMENTS**

We are indebted to our colleague Scott Hunnicke-Smith and members of the Genomic Sequencing and Analysis Facility for valuable insights. We are grateful to members of the Finkelstein laboratory for carefully reading the manuscript. This work was supported by a College of Natural Sciences Catalyst award, CPRIT (R1214 to I.J.F.), the Welch Foundation (F-1808 to I.J.F.), the National Science Foundation (1455358), and the NIH (F32 AG053051 to S.K.J. and R01 GM120554 to I.J.F.). This publication was also made possible through the support of a grant from the John Templeton Foundation (JTA16-000194 to A.D.E). The opinions expressed in this publication are those of the authors and do not necessarily reflect the views of the John Templeton Foundation. I.J.F. is a CPRIT Scholar in Cancer Research. The content is solely the responsibility of
the authors and does not necessarily represent the official views of the National Science Foundation.

Received: September 9, 2016
Revised: April 23, 2017
Accepted: May 26, 2017
Published: June 29, 2017

REFERENCES


Schirle, N.T., and MacRae, I.J. (2012). The crystal structure of human Argonaute2. Science 336, 537–539.


## STAR★METHODS

### KEY RESOURCES TABLE

<table>
<thead>
<tr>
<th>REAGENT or RESOURCE</th>
<th>SOURCE</th>
<th>IDENTIFIER</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antibodies</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mouse anti-FLAG M2</td>
<td>Sigma-Aldrich</td>
<td>Cat# F3165; RRID: AB_259529</td>
</tr>
<tr>
<td>Rabbit anti-HA</td>
<td>ICL labs</td>
<td>Cat# RHGT-45A-Z</td>
</tr>
<tr>
<td><strong>Bacterial and Virus Strains</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>BL21 star (DE3) cell</td>
<td>Thermo Fisher Scientific</td>
<td>Cat# C6010-03</td>
</tr>
<tr>
<td>BL21 (DE3) cell</td>
<td>New England Biolabs</td>
<td>Cat# C2527H</td>
</tr>
<tr>
<td>BL21-AI competent cell</td>
<td>Huo et al., 2014</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Chemicals, Peptides, and Recombinant Proteins</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>DNase</td>
<td>GoldBio</td>
<td>Cat# D-301-500; CAS: 9003-98-9</td>
</tr>
<tr>
<td>HALT protease inhibitor</td>
<td>Thermo Fisher Scientific</td>
<td>Cat# 78439</td>
</tr>
<tr>
<td>Desthiobiotin</td>
<td>Sigma-Aldrich</td>
<td>Cat# D1411; CAS: 533-48-2</td>
</tr>
<tr>
<td>Alexa Fluor 488 Antibody Labeling Kit</td>
<td>Thermo Fisher Scientific</td>
<td>Cat# A20181</td>
</tr>
<tr>
<td>Alexa Fluor 647 Antibody Labeling Kit</td>
<td>Thermo Fisher Scientific</td>
<td>Cat# A20186</td>
</tr>
<tr>
<td>Bst 2.0 WarmStart DNA polymerase</td>
<td>New England Biolabs</td>
<td>Cat# M0538S</td>
</tr>
<tr>
<td>Tween-20</td>
<td>Sigma-Aldrich</td>
<td>Cat# P9416; CAS: 9005-64-5</td>
</tr>
<tr>
<td>Q5 High Fidelity DNA polymerase</td>
<td>New England Biolabs</td>
<td>Cat# M0491S</td>
</tr>
<tr>
<td>γ32P-ATP</td>
<td>PerkinElmer</td>
<td>Cat# BLU502H500UC</td>
</tr>
<tr>
<td>T4 polynucleotide kinase</td>
<td>New England Biolabs</td>
<td>Cat# M0201S</td>
</tr>
<tr>
<td>AMP-PNP</td>
<td>Sigma-Aldrich</td>
<td>Cat# 10102547001; CAS: 25612-73-1 (free acid)</td>
</tr>
<tr>
<td>CoCl2</td>
<td>Sigma-Aldrich</td>
<td>Cat# 232696; CAS: 7646-79-9</td>
</tr>
<tr>
<td>ATP</td>
<td>Sigma-Aldrich</td>
<td>Cat# 2383; CAS: 34369-07-8</td>
</tr>
<tr>
<td>Proteinase K</td>
<td>New England Biolabs</td>
<td>Cat# P8107S</td>
</tr>
<tr>
<td><strong>Oligonucleotides</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>See Table S1</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Recombinant DNA</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HeLa genomic DNA</td>
<td>New England Biolabs</td>
<td>N4006S</td>
</tr>
<tr>
<td>pRSF (crRNA)</td>
<td>This study</td>
<td>N/A</td>
</tr>
<tr>
<td>pCDF (SUMO-CasA)</td>
<td>This study</td>
<td>N/A</td>
</tr>
<tr>
<td>pET19 (2XFLAG-CasBtoE)</td>
<td>This study</td>
<td>N/A</td>
</tr>
<tr>
<td>pET28b (HA-Cas3)</td>
<td>This study</td>
<td>N/A</td>
</tr>
<tr>
<td>pET28b (Cas3)</td>
<td>Huo et al., 2014</td>
<td>N/A</td>
</tr>
<tr>
<td>pBAD (Cascade)</td>
<td>Huo et al., 2014</td>
<td>N/A</td>
</tr>
<tr>
<td>pACYC-Duet-1 (crRNA)</td>
<td>Huo et al., 2014</td>
<td>N/A</td>
</tr>
<tr>
<td>pCDF-Duet-1 (target DNA)</td>
<td>Huo et al., 2014</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Software and Algorithms</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CHAMP</td>
<td>FinkelsteinLab</td>
<td><a href="https://github.com/finkelsteinlab/champ">https://github.com/finkelsteinlab/champ</a></td>
</tr>
<tr>
<td>mManager</td>
<td>Open Imaging</td>
<td><a href="http://micro-manager.org">micro-manager.org</a></td>
</tr>
<tr>
<td>Source Extractor</td>
<td>Emmanuel Bertin</td>
<td><a href="http://www.astromatic.net/software/sextractor">http://www.astromatic.net/software/sextractor</a></td>
</tr>
<tr>
<td>Trimmomatic</td>
<td>Usadel Lab</td>
<td><a href="http://usadellab.org/cms/?page=trimmomatic">usadellab.org/cms/?page=trimmomatic</a></td>
</tr>
<tr>
<td>Bowtie2</td>
<td>Ben Langmead</td>
<td><a href="https://github.com/BenLangmead/bowtie2">https://github.com/BenLangmead/bowtie2</a></td>
</tr>
<tr>
<td><strong>Other</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MiSeq chips</td>
<td>Illumina</td>
<td><a href="https://www.illumina.com/">https://www.illumina.com/</a></td>
</tr>
<tr>
<td>HiSeq chips</td>
<td>Illumina</td>
<td><a href="https://www.illumina.com/">https://www.illumina.com/</a></td>
</tr>
</tbody>
</table>
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact Ilya J. Finkelstein (finkelstein@cm.utexas.edu).

METHOD DETAILS

Protein Cloning and Purification

*T. fusca* Cascade and Cas3 were overexpressed and purified as described previously with minor modifications (Huo et al., 2014). Briefly, the Cascade complex and crRNA were expressed from pET-based plasmids that were co-transformed into BL21 star (DE3) cells (Thermo-Fisher). Cse1 contained a His6/Twin-Strep/SUMO N-terminal fusion, while Cas6 contained an N-terminal triple FLAG epitope for fluorescent labeling. Single colonies were used to inoculate LB + Kanamycin/Carbenicillin/Streptomycin media. At OD₆₀₀ 0.8, cells were induced with 1 mM IPTG overnight at 25°C. Cells were then lysed in 20 mM HEPES, pH 7.5, 500 mM NaCl, 2 μg mL⁻¹ DNase (GoldBio) and 1x HALT protease inhibitor (Thermo-Fisher), and the clarified lysate was applied to a hand-packed Strep-Tactin Superflow gravity column (IBA Life Sciences) for purification via the Twin-Strep tagged Cse1. The Cascade complex was eluted with 20 mM HEPES, pH 7.5, 500 mM NaCl, 5 mM desthiobiotin, and then concentrated by centrifugal filtration (30 kDa Amicon, Millipore). The concentrate was then incubated overnight at 4°C with 3.3 μM SUMO protease to remove tags from Cse1. The complex was further fractionated over a HiLoad 16/600 Superdex 200 column (GE Healthcare) equilibrated in storage buffer (10 mM Tris-Cl, pH 7.5, 500 mM NaCl, 5 mM DTT). Fractions containing the full Cascade complex were determined by SDS-PAGE, pooled, and concentrated to ~5-10 μM (30 kDa centrifuge concentrators, Millipore). Small aliquots were flash frozen in liquid nitrogen and stored at −80°C. All aliquots were used only once and not refrozen.

Antibodies

Cascade and Cas3 were fluorescently labeled with mouse anti-FLAG M2 (F3165, Sigma) and Rabbit anti-HA (RHGT-45A-Z, ICL labs), respectively. Antibodies were conjugated to Alexa488 or Alexa647 at a ratio of ~1:3 antibody:dye according to the manufacturer’s instructions (Molecular Probes Alexa Fluor antibody labeling kits, Thermo Fisher Scientific). The antibody to dye conjugation ratio was measured using a NanoDrop (Thermo Fisher Scientific) according to the manufacturer-provided protocol. Fluorescent antibodies were stored in PBS buffer (pH 7.2, with 2 mM sodium azide) at −20°C.

DNA oligonucleotides libraries

Oligonucleotides were purchased from IDT or IBA (see Table S1). A synthetic oligonucleotide with six randomized bases was purchased from IDT and used to profile the extended six nucleotide PAM. Two additional synthetic oligonucleotide libraries were designed to measure the effects of mismatches along the entire target DNA sequence. These libraries were made by randomizing the bases along the entire length of the consensus target DNA sequence. In these “doped” libraries, every correct base had a 9% chance of being substituted for each of three other bases (3% each; 9% total). This doping mixture was chosen to provide comprehensive coverage for sequence variants with a Hamming distance less than three on a typical MiSeq chip (representing ~20-25 million unique reads). Pooled custom DNA libraries were also purchased from CustomArray. DNA libraries were sequenced on a MiSeq (Illumina) using a 2x75 or a 2x300 paired end reagent kit (v3).

Exome preparation and sequencing

HeLa genomic DNA (NEB N4006S) was prepared using the TruSeq Exome Library Prep Kit (Illumina), yielding approximately 170 basepair-long DNA fragments. The exome library was then sequenced using the MiSeq Reagent Kit v3 (Illumina, 2x300 paired-end reads). The resulting MiSeq run yielded 9.1 million exome reads.

Chip regeneration and addition of alignment markers

After sequencing, MiSeq chips were kept at 4°C in storage buffer (10 mM Tris-Cl, pH 8.0, 1 mM EDTA, 500 mM NaCl). All imaging and chip regeneration steps were carried out in a custom-built microscope stage adaptor with integrated microfluidic interconnects. An overview of the microscope stage and fluidic interface is summarized in Figure S1. Detailed blueprints of all components are also available via GitHub (https://github.com/finkelsteinlab/). Temperature was controlled by PiWarmer, a home-built Raspberry Pi-controlled heating element. PiWarmer was also used to run the heating and cooling cycles required for on-chip cluster regeneration. Schematics and code for assembling the temperature controller, as well as protocols for chip regeneration are available via GitHub (https://github.com/finkelsteinlab/). The heating element was mounted on the microscope turret to allow for easy and consistent heat application.

All fluidic methods utilized an automated syringe pump (KD scientific) operating at a flow rate of 100 μl min⁻¹ for chip preparation and experimentation. All reagents were added to the flow path through an automated, multi-position valve (Rheodyne MXP9000) containing either a 100 or 700 μl injection loop.

To regenerate the DNA clusters, all DNAs covalently affixed to the MiSeq chip surface were denatured with 500 μl 0.1 N NaOH as it flowed through the chip (5 min) and similarly washed with 500 μl TE buffer. This removed the untethered DNAs strands containing...
residual fluorescent dyes from sequencing (see Figure S1). After denaturation, the chip was heated to 85°C and incubated with 500 nM of the regeneration primer (CJ.RP) in hybridization buffer (75 mM Trisodium Citrate, pH 7.0, 750 mM NaCl, 0.1% Tween-20). CJ.RP was annealed at 85°C for 5 min, followed by ramped linear cooling to 65°C over 10 min, ramped linear cooling from 65°C to 40°C over 30 min, and then washed with 1 mL washing buffer (4.5 mM Trisodium Citrate, pH 7.0, 45 mM NaCl, 0.1% Tween-20) at 40°C (10 min). CJ.RP binds to all user clusters but does not target phiX clusters. CJ.RP was extended at 60°C for 10 min in isothermal amplification buffer (20 mM Tris-HCl, pH 8.8, 10 mM (NH₄)₂SO₄, 50 mM KCl, 2 mM MgSO₄, 0.1% Tween-20) containing 0.08 U/μl of Bst 2.0 WarmStart DNA polymerase (New England Biolabs) and 0.8 mM of dNTPs. The chip was then washed with 500 μL hybridization buffer at 60°C to remove the polymerase (5 min). Finally, a phiX primer labeled with Atto647 or Cy3 (atto647-PCP / Cy3-PCP) was annealed under the same conditions as CJ.RP. The resultant fluorescent phiX clusters were used for aligning the FASTQ points to imaged clusters (see Figure S1 and Computational Methods below). Prepared chips could be used for at least a dozen Cascade-DNA binding experiments before requiring regeneration.

**Fluorescence microscopy**

All fluorescence images were collected using a Nikon Ti-E microscope in a prism-TIRF configuration equipped with a motorized stage (Prior ProScan II H117) containing the experimental MiSeq chip (Illumina) housed in our custom stage adaptor (Figure S1). The chip was illuminated with 488 nm (Coherent), 532 nm (Ultralasers), or 633 nm (Ultralasers) lasers through a quartz prism (Tower Optical). The laser exposure was controlled with high-speed shutters (LS68220, Vincent Associates) To minimize spatial drift, the microscope was assembled on a floating optical table (TMC). An active feedback system was used to maintain focus across the entire chip surface (Nikon PerfectFocus). Data were collected with a 100 ms exposure through a 60X water-immersion objective (1.2NA, Nikon) paired with (i) a quad-band filter (89401 Chroma), a 638 nm dichroic beam splitter, and either a 600 nm long-pass filter or 500 nm long pass / 600 nm short pass filters (Chroma), or (ii) a dual-band filter (ZET352/660 m Chroma), a 640 nm dichroic beam splitter, and either a 655 nm long-pass filter or ET4585/65 m band pass filter (Chroma), which allowed multi-channel detection through two EMCCD cameras (Andor Ixon DU897, cooled to –80°C). Images were collected using Micro-Manager Open Source Microscopy software (Edelstein et al., 2014) and saved in an uncompressed TIFF file format for later analysis via a custom written image-processing pipeline (see below).

**CHAMP assays**

Increasing concentrations of the Cascade complex (0.063, 0.16, 0.39, 1, 2.5, 6.3, 16, 39, 100, 250, and 630 nM) were injected into a regenerated MiSeq chip and incubated at 60°C for 10 min in imaging buffer (40 mM Tris-HCl, pH 8.0, 150 mM NaCl, 2 mM MgCl₂, 1 mM DTT, 0.2 mg mL⁻¹ BSA, 0.1% Tween-20). After the incubation, excess Cascade was rapidly flushed out of the chip while the remaining proteins were labeled; this was accomplished by washing with 100 μl imaging buffer at 60°C, then 100 μl of 20 nM fluorescently conjugated anti-FLAG antibody in imaging buffer at 25°C, and then an additional 100 μl of imaging buffer at 25°C (3 min total). Control experiments that omitted Cascade indicated that the fluorescent antibodies did not bind to the chip surface.

For each Cascade concentration, we imaged up to 812 fields of view spanning nearly 50% of the total sequenced MiSeq chip surface area. The chip was illuminated with 20, 40 or 30 mW of laser power at 488, 532, or 633 nm, respectively (measured at the front face of the TIRF prism). To prevent photobleaching, the lasers were shuttered between subsequent fields of view during the ~15 min of image acquisition. No appreciable Cascade dissociation or cluster photobleaching occurred during this time (see Figure S2F). In order to avoid pixel saturation at high protein concentrations, ten 100 ms images were captured at each field of view. These images were summed into a final image and stored in hd5 file format by channel and position. Care was taken to minimize experiment-to-experiment variation by acquiring all concentrations of a titration series in a single day. Following each experiment, the MiSeq chips were deproteinized with 32 units of Proteinase K (New England Biolabs) in washing buffer for 30 min at 42°C and the chip showed no sign of degradation even after twelve Proteinase K treatments. The DNA in a chip could be denatured and re-synthesized up to five times using the regeneration protocol described above.

**Electrophoretic mobility shift assay (EMSA)**

All EMSAs were performed with radioactively or fluorescently labeled PCR products containing the indicated PAM and protospacer, as well as flanking sequences used in the CHAMP experiments (i.e., Illumina adapters). PCR was performed using 1 ng of template plasmid containing the desired PAM/protospacer, 500 nM of P5 primer for radioactive-labeling or Cy5-P5 primer for fluorescent-labeling, 500 nM of CJ.RP, 200 μM of dNTPs and 0.5 unit of Q5 high-fidelity DNA polymerase (New England Biolabs) in a 25 μl reaction on an MJ Research PTC-200 Thermal Cycler. The PCR product was purified (PCR purification kit, Qiagen) and quantified on a NanoDrop spectrophotometer (Thermo Fisher Scientific). For radioactive assays, PCR products were labeled with γ³²P-ATP (PerkinElmer) using T4 polynucleotide kinase (New England Biolabs). The labeled PCR products were purified with MicroSpin G-25 columns (GE Healthcare).

Cascade binding assays were performed by incubating 0.1 nM of ³²P-labeled dsDNA with increasing Cascade concentrations (0.025, 0.063, 0.16, 0.39, 1, 2.5, 6.3, 16, 39, 100, 250, 630 nM) for 30 min at 62°C in binding buffer (40 mM Tris-HCl, pH 8.0, 150 mM NaCl, 2 mM MgCl₂, 1 mM DTT, 0.2 mg mL⁻¹ BSA, 0.01% Tween-20). The reactions were resolved on a 2.5% agarose gel run with 0.5X TBE buffer. Gels were dried and DNA was visualized using a Typhoon scanner (GE Healthcare). ImageQuant
software (GE Healthcare) was used to quantify the bound and unbound DNA amounts. The fraction of bound DNA was fit to the Hill equation to obtain $K_r$ values. All experiments were repeated in triplicate.

To observe Cas3 binding, Cascade (39 nM) and target dsDNA (2 nM) were pre-bound for 30 min at 62°C in a binding buffer. Then, Cas3 and AMP-PNP (Sigma) were added into the EMSA reaction for final concentrations of 1.1 μM and 2 mM, respectively and incubated for 10 min at 62°C. The reactions were resolved on a 5% native PAGE gel containing 0.5X TBE buffer and visualized using a Typhoon scanner (GE Healthcare).

**Cas3 nuclease assays**
Cascade (39 nM) was first incubated with Cy5-labeled target dsDNA (2 nM) for 30 min at 62°C in binding buffer. Then, Cas3, CoCl$_2$ (Sigma) and ATP (Sigma) were added into the EMSA reaction at final concentrations of 650 nM, 111 μM and 1.9 mM, respectively and incubated for 30 min at 62°C. The reaction was quenched with 50 mM EDTA and deproteinized with proteinase K. The reactions were resolved on a 10% denaturing PAGE gel containing 0.5X TBE buffer and visualized using a Typhoon scanner (GE Healthcare).

**Plasmid loss assays**
The Cascade expression construct was generated by insertion of the Cascade gene cassette (encoding all protein subunits) into a pBAD (ApR) vector. The pre-crRNA expression cassette containing five identical CRISPR units for target A, was cloned into the pCDF-Duet-1 (SmR) vector. A 127-bp fragment containing a protospacer and a PAM for target A was cloned into the pCDF-Duet-1 (SmR) vector to serve as the target DNA. In vivo assays were performed with *T. fusca* Cascade and Cas3 plasmids as described previously (Huo et al., 2014).

**Computational Methods**
The main challenge for CHAMP is the precise mapping of each individual DNA cluster to an underlying DNA sequence. This is because CHAMP uses images obtained via conventional TIRF microscopy and the information in these images is only partially encoded in the sequencing output generated by all Illumina sequencers. These images are transformed by an arbitrary translation, scaling, and rotation relative to the coordinate system used in the Illumina software. Alignment between the sequencing output and CHAMP images is further confounded by false-positive (e.g., spurious fluorescent signals) and false-negative cluster coordinates (e.g., fluorescent signals that are filtered out by the Illumina sequencing software). CHAMP overcomes this challenge by using alignment markers with known DNA sequences to match the spatial position of all fluorescent clusters to a corresponding record in the sequencing output file (Figure S2A). A library consisting of the bacteriophage PhiX genome was used as the alignment marker because this DNA is included as an internal control and typically comprises 5%–10% of all sequenced DNA clusters on every Illumina chip. This library also contains a unique sequencing adaptor that can be selectively illuminated with a fluorescent primer (Figure S1). Mapping the alignment markers and protein-bound clusters requires two stages: first, a rough alignment using Fourier-based cross correlation methods is performed, followed by a precision alignment using least-squares constellation mapping between FASTQ and de novo extracted clusters (see Figure S2). This is a specialized example of the image registration problem (Zitova and Flusser, 2003), and allows CHAMP to function with any fluorescence-based sequencing platform and TIRF microscope.

**Aligning Fluorescent Images and FASTQ Points: Overview**
To identify the DNA sequence of each cluster, we developed an image-processing pipeline to process images collected by TIRF microscopy. To decode each cluster’s sequence, its position was correlated to the corresponding record in the FASTQ file generated at the end of each MiSeq run. For each identified cluster, the FASTQ file reports the specifying lane, tile, and relative x-y coordinates. However, the FASTQ-supplied spatial information is reported in an arbitrary coordinate system that is scaled, rotated, and translated relative to our fluorescent images. An additional confounding factor is that FASTQ files do not report all fluorescent clusters (e.g., clusters that did not pass Illumina-specified quality control filters). In addition, some Illumina-reported clusters may also not light up in our fluorescent images. This may occur due to errors in the Illumina cluster identification pipeline, or possibly due to incomplete fluorescent labeling of the cluster sequences during our experiments. As such, the mapping problem required finding the rotation, scale, x-offset, y-offset, and chip surface (both surfaces are imaged in a MiSeq chip) which best aligned the FASTQ points and imaged clusters. We accomplished this through two alignment stages: rough alignment and precision alignment, discussed below.

For the purposes of internal calibration, Illumina requires a percentage of each MiSeq run, typically 5%–10% of all clusters, to be DNA from the small, thoroughly characterized phiX bacteriophage genome. Separate adaptor chemistry is used for this phiX library, which can be accurately and specifically illuminated on any chip using complementary oligonucleotides. The phiX clusters do not contain a run-specific index barcode and are thus not demultiplexed as normal reads, but can be determined by mapping reads to the phiX genome. These phiX clusters provide a convenient resource for a variety of purposes, including alignment, categorization, and intensity training, and as a control. We illuminated the phiX clusters by hybridizing them to a dye-conjugated oligo (Atto647-PCP or Cy3-PCP) during cluster re-generation and used the resulting fluorescent signals to align our fluorescent images with the corresponding FASTQ records.

**Stage 1: Rough Alignment**
The rough alignment was performed through cross-correlation of FASTQ points and images using fast Fourier methods (Press et al., 2007). Briefly, each FASTQ tile was converted to an image, each cluster represented as a radially symmetric Gaussian with $\sigma$ of 0.25 μm, a typical cluster size. Cross-correlation was then performed via the formula

$$K_r = \frac{\sum_{i}^{n} (x_i - \mu_x)(y_i - \mu_y)}{\sqrt{\sum_{i}^{n} (x_i - \mu_x)^2 \sum_{i}^{n} (y_i - \mu_y)^2}}$$
Cross correlation = |$F^{-1}((FF)^*F T)$|

with zero-padding enough to accommodate any offset, where $F$ and $F^{-1}$ are the fast forward and inverse 2D Fourier transforms, $*$ is the complex conjugate, $F$ is the FASTQ image, and $T$ is the TIRF image. This allowed consideration of all x-y offsets (translation) in a computationally efficient manner, though did not inherently consider rotation or scale. For each TIRF image, the maximum cross-correlation was first found against two FASTQ tiles known from their position to not overlap the TIRF image in order to measure background noise level, after which correlations above a signal-to-noise cutoff of choice, 1.4 in the current work, indicated a good alignment. In order to achieve our first alignment, we exhaustively sampled the parameter space around initial estimates of rotation, scale, and parity. The first rough alignment established the approximate rotation and scale, and was performed on each MiSeq chip to account for small deviations in their mounting within the custom-built stage adaptor. With reasonable estimates for these parameters, the Fourier-based alignment can be performed within 45 s on a desktop computer.

Stage 2: Precision Alignment

Following rough alignment in the alignment marker image channel, we performed precision alignment via constellation mapping in all channels. Our algorithm aimed to maximize the number of matches between FASTQ points and fluorescent clusters, forming the same “constellation” in each space. The mapping parameters were then quickly determined using linear least-squares fitting.

First, cluster location information was extracted from the TIRF images. We used the astronomy software Source Extractor to fit two-dimensional Gaussian functions to the fluorescent clusters (Bertin and Arnouts, 1996). Next, we found the nearest neighbors of FASTQ points in imaged cluster space and vice-versa using kd-trees (Maneewongvatana and Mount, 1999). Two points which were nearest neighbors of each other in both directions were termed a mutual hit. Due to accrued noise — missing data in FASTQ space, missing data in imaged cluster space, and imperfect Gaussian calling — mutual hits were not by themselves high-confidence mappings. We further subcategorized mutual hits by the statuses of other nearby clusters. If cluster A and FASTQ point B were mutual hits and no other cluster X or FASTQ point Y consider A or B nearest neighbors, then the mutual hit was termed an exclusive hit. If there was another cluster X whose nearest neighbor was FASTQ point B, or another FASTQ point Y whose nearest neighbor was cluster A, then the status of hit AB was determined by the distance to the closest such X or Y. If the closest such X or Y was more than 1.25 microns away — the diameter of a typical cluster — AB was termed a good mutual hit; otherwise AB was called a bad mutual hit. Using exclusive hits and good mutual hits, we then performed linear least-squares fitting to determine the final alignment. The precision alignment process, including both constellation identification and least-squares fitting, is typically performed within 2.5 s on a desktop computer.

Calculating Cluster Intensity

Machine-learned linear weighting of pixels was used to calculate the fluorescent intensity of each cluster (see Figure S2). For training, we used an experiment with only phiX clusters illuminated and restricted the analysis to exclusive and good mutual hits. Seven by seven pixel squares were extracted around each of these FASTQ points and linearized into feature vectors. Linear Discriminant Analysis (LDA) was then used to find pixel weights that best capture the intensity of a given cluster and penalize the intensity of neighboring clusters. The positive weights were used to calculate raw cluster intensities. To correct for variation in laser intensities across fields of view, cluster intensities were normalized within each run. The mode of pixel intensities of each image was calculated, and the intensity calculations in each image were normalized by the mode of the given image divided by the median of all modes.

Data Analysis

Calculating the apparent dissociation constant

Calculation of the apparent $K_d$ value was performed for each sequence via curve fitting to the Hill equation (without cooperativity):

$$I_{\text{obs}} = \frac{I_{\text{max}} - I_{\text{min}}}{1 + \frac{K_d}{[X]}}$$

where $I_{\text{min}}$ is the background intensity, $I_{\text{max}}$ is the intensity of a fully saturated cluster, and the concentration values $X$ and cluster intensity values $I_{\text{obs}}$ are derived from the concentration gradient experiment. $I_{\text{min}}$ is calculated as the median intensity of negative control clusters in the lowest concentration point. $I_{\text{max}}$ is determined separately for each concentration to normalize small differences in fluorescence intensities across the entire flowcell and between concentrations. At higher concentrations, DNA sequences that are perfectly complementary to the crRNA-Cascade complex become saturated and can be used as a reference to normalize between concentrations. To this end, $I_{\text{max}}$ is calculated in two steps, using only clusters of the perfect target sequence. First, the $K_d$ and a temporary, constant $I_{\text{max}}$, call it $I_{\text{max, const}}$, are fit jointly on the perfect target sequence clusters using information from all concentrations. Second, for each concentration where median $I_{\text{obs}}$ is greater than 90% of the fit $I_{\text{max, const}}$, $I_{\text{max}}$ is solved for from the above equation, using the observed median cluster intensity as $I_{\text{obs}}$. At all preceding concentrations, $I_{\text{max, const}}$ is used. These values of $I_{\text{min}}$ and $I_{\text{max}}$ are then used to fit $K_d$ for all other sequences. Error bars indicate the standard deviation of bootstrap $K_d$ values.
**Position-Transition Model**

The position transition model for change in apparent binding affinity (ΔABA) can be written as:

$$
\Delta ABA = \sum_{j=1}^{35} p_j t(r, s_j)
$$

where $p_j$ is the penalty, $r_j$ is the reference base, and $s_j$ is the sequenced base in the $j^{th}$ position, and $t(x, y)$ is the position-independent transition weight from $x$ to $y$. The summation is carried out over all 35 positions in the minimal three-nucleotide PAM and the protoscaler.

For computational efficiency, we cast this in matrix form. We represented each sequence as a 35-by-12 indicator matrix $S$ with rows representing each sequence position and columns representing each non-identity transition. The position penalties and transition weights were represented as vectors $p$ and $t$. Then the above is written as

$$
\Delta ABA = S : (p \otimes t)
$$

where: is the Frobenius inner product and $\otimes$ is the outer product. This was linearized and concatenated into multiple-sequence sparse matrices and fit using non-linear least-squares. Having multiple reference sequences and normalizing the transition vector to have mean value one, obviated model degeneracy.

**Cas3 Penalties**

The slope of stoichiometric Cascade/Cas3 intensity was fit to all single-mismatch data with a mismatch in the fourth target position or greater. Cas3 penalties were then calculated as the observed Cas3 average intensity minus the expected stoichiometric intensity given average Cascade intensity, such that points furthest from the line represented sequences with the greatest difference in Cas3 versus Cascade occupancy. Error bars are the SEM of intensity values.

**Exome dataset analysis**

Exome reads were first trimmed with Trimmomatic 0.32 to remove Illumina adaptor sequences (Bolger et al., 2014). Trimmed reads were then mapped to the human genome using Bowtie2 2.2.3 (Langmead and Salzberg, 2012). We filtered for read quality and mapped phred score above 20, resulting in seven million high quality mapped reads, or an average 11-fold coverage in regions of interest. For each position with at least five overlapping imaged reads, intensity information from all reads was used to measure ABA, following the same procedure as with the synthetic libraries. This results in a flat signal across most of the genes, with peaks at off-target sites with high ABAs. The peak width reflects both the distribution of read lengths and coverage depth across the library. Below, we demonstrate that this results in a triangle-shaped function.

For our observed read length distribution, this is approximately true for $|x - x_b| < 100$ bp (Figure S4A). This accounts for the top > 60% of the peak, so the theoretical peak shape is approximately triangular (Figure S4B). If all reads had the same length, this would result in a perfectly triangular peak. Due to library size-selection, read lengths were relatively focused around the mean length (Figure S4A), so the resulting theoretical peak shape is approximately triangular (Figure S4B). Using the observed read length distribution results in theoretical peaks with a full width at half maximum (FWHM) of 162 bp. The experimental peak shape was determined by summing the normalized peak shapes from the top thirty high-affinity DNA binding sites. Remarkably, this result is in near quantitative agreement with the theoretical calculations with an observed FWHM of 210 bp. Deviation from the theoretical shape is due to finite coverage, bias in shearing sites, and the non-linear map from reads included to measured ABA. We therefore used the more conservative estimate of 210 bp as our cutoff for determining the underlying consensus motif. This motif was determined by searching a 210 bp window around the peak of the ABA curves for the presence of a high-affinity PAM and crRNA-complementary DNA. The results were plotted as a logo using WebLogo (Crooks et al., 2004).

**DATA AND SOFTWARE AVAILABILITY**

The source code for cluster identification, spatial registration, and binding affinity calculations is available via GitHub (https://github.com/finkelsteinlab/).
Supplemental Figures

Figure S1. Overview of the CHAMP Experimental Platform, Related to Figure 1

(A) MiSeq chips are imaged via prism-based total internal reflection fluorescence (TIRF) microscopy on a custom-built microscope stage. Three lasers are used to excite the fluorophores. Exposure times are controlled by three computer-controlled shutters (S1-S3). Neutral density filters (F1-F3) are used to control the laser intensity, long-pass dichroic mirrors (DM1-DM2) combine the laser beams into a single path, mirrors (M1-M2) direct the beams through a prism to generate an evanescent excitation field for TIRF imaging. The reflected beams are blocked at a beam stop (BS). The emitted photons pass through the objective and a computer-controlled filter wheel (FW) that removes residual laser excitation. A dichroic mirror (DM3) separates spectrally distinct fluorophore emissions, which are directed toward two electron-multiplying charge coupled device cameras (EM-CCDs) for wide-field imaging. Reagents are delivered to the microfluidic chip via a computer-controlled syringe pump. Temperature is controlled via a custom-built Raspberry Pi-based controller (plans available at http://github.com/finkelsteinlab). (B) A diagram of the MiSeq chip adaptor. The MiSeq chip is inserted into the chip holder and secured to the base plate in combination with the tubing holder. Microfluidic tubing is fit into the tubing holder, passed between the tubing guide and pressure plate, and mated with the MiSeq chip. Blueprints for this assembly are available via GitHub (https://github.com/finkelsteinlab). (C) Regenerating DNA clusters on a sequenced MiSeq chip. After sequencing, the chip contains residual fluorescence in all emission channels (left). The residual fluorescence and sequenced DNA strands are stripped with NaOH and the DNA is regenerated (middle two panels). PhiX clusters are labeled with a fluorescent oligonucleotide (magenta) for downstream image alignment. Cascade is incubated in the chip and binds a subset of the DNA clusters. Cascade can be visualized after the addition of fluorescent anti-FLAG antibody, (fifth panel, green). After chip regeneration, all fluorescent signals are sensitive to DNase I treatment, indicating that these signals originate from DNA clusters.
Figure S2. Related to Figure 1
(A) Flow chart for cluster identification. (B) A representative alignment. The first image (green) shows the alignment marker coordinates, each represented by a radially symmetric Gaussian. These coordinates are found by mapping all reads against the PhiX genome, and aligning the mapped reads with a TIRF microscope image with fluorophores attached to all alignment markers (magenta, middle). The third image shows the overlap of the synthetic and experimental images (overlap seen as white). (C) Example 7x7 pixel images centered on aligned FASTQ points for targeted and non-targeted clusters. (D) Linear discriminant analysis (LDA) was used to train pixel weights using sub-images as in (C) from sequences known to be on or off. Shown are the trained weights. 7x7 pixels sub-imaged were found to be optimal. To calculate intensity scores for K_a calculations, these weights, with negative values set to zero, are multiplied by the corresponding pixel values and summed. (E) The ROC (receiver operating characteristic) curve using LDA scores from (D) for classification of a test set of approximately 75,000 points. Perfect target A sequences were used as ground-truth positive values, and non-target sequences as ground-truth negative values when calculating the true- and false-positive rates (TPR, FPR). The extremely high area under the curve (AUC) of 0.999 indicates both very good alignment of the sequence coordinates and microscope images, as well as high fidelity of the chemistry in illuminating the correct clusters and only the correct clusters. (F) Fluorescent signal intensity remains constant throughout the CHAMP experiment. Cascade (10 nM) was incubated on an NGS chip for 10 min at 60°C, then washed and labeled with

(legend continued on next page)
anti-FLAG Alexa488 antibody. Images were then collected every five minutes for one hour. The graph above represents the mean intensity of all clusters containing the perfectly basepaired target DNA sequence. Error bars: SEM. The normalized data were fit to an exponential decay curve to estimate the half-life (dashed line). (G) Estimating the error in the ABA. Bootstrap ABA values were calculated for the perfect target sequence with all numbers of clusters between 3 and 100. Shown are the average errors (blue points) and 90% confidence intervals of error (red points), using the ABA fit with 2,000 clusters as reference. The gray dotted line shows a cutoff of 5 clusters, with average ABA error of approximately 0.2 k_BT. Solid lines indicate a fit to the data.
Figure S3. Electrophoretic Mobility Shift Assays for Different PAMs and Flipped-Out Bases, Related to Figures 2 and 3

(A) Electrophoretic mobility shift assays (EMSA) were performed with eight different PAM sequences and the K_d values were calculated by fitting the fraction of Cascade-dsDNA complexes at each Cascade concentration to the Hill equation. Error bars: SD from three replicates. (B) The K_d values obtained from EMSA were plotted with ABAs derived from a CHAMP experiment for the eight PAMs shown in (A). Horizontal error bars: SEM of the ABAs. (C) Schematic of the DNA sequences and (D) EMSAs with radiolabeled dsDNAs containing an “AAG” PAM and the three different groupings of flipped-out bases. Radiolabeled dsDNA (0.1 nM) was incubated with Cascade (lane 1: 0.033 nM, 2: 0.083 nM, 3: 0.21 nM, 4: 0.53 nM, 5: 1.3 nM, 6: 3.3 nM, 7: 8.3 nM, 8: 21 nM, 9: 53 nM) and resolved on a 2.5% agarose gel. (E) K_d values were obtained by fitting the data from three replicates to a Hill equation. Error bars: SD.
Figure S4. Exome Sequence Length Distribution and Expected Peak Shape, Related to Figure 4

(A) The distribution of exome sequence lengths. The DNA was sheared and sized to a nominal DNA fragment length of approximately 150 bp. The observed mean DNA length and coefficient of variation were 170 bp and 22%, respectively. (B) The resolution of measuring a DNA binding site in a randomly sheared DNA sample depends on the fragment length distribution and the coverage depth of each fragment. The shear lengths from (A) were used to calculate the probability that a random read covering a nearby base would also cover a target binding site (red dashed curve, see Methods). In the limit of infinite coverage and perfectly random shearing, this gives the range of influence a binding site has on measurements for nearby bases, and hence provides an estimate for the resolution of this method. In the current experiment, the full width at half maximum (FWHM) of this peak is 162 bp. The observed resolution was calculated by normalizing and averaging the thirty highest-affinity binding peaks (blue curve). The experimentally observed FWHM was 210 bp and was used to define the resolution for this experiment. Deviations from the expected peak shape (red) are due to finite coverage, bias in shearing sites, and the non-linear map from reads included to measure ABA.
Figure S5. Cse1 Dissociates from the Cascade Complex, Related to Figure 6

(A) EMSA of a DNA with a “TTAC” PAM and perfectly paired target DNA. Cse1 is dissociated from 50% of the Cascade-DNA complexes (lane 2). Adding excess Cse1 can drive its re-association with the nucleoprotein complex. (B) Quantification of three replicates. Light gray: Cascade/dsDNA, gray: Cascade without Cse1, dark gray: free double-stranded DNA. Error bars indicate SD [DNA]: 2 nM, [Cascade]: 39 nM, additional [Cse1]: 0, 0.11, 0.22, 0.43 and 0.87 µM. All components were incubated together at 62°C for 10 min.
Figure S6. Adapting CHAMP for Advanced Imaging Modalities and HiSeq Sequencers, Related to Figure 5 and the STAR Methods

(A) FRET-CHAMP on a MiSeq chip. A subset of DNA clusters was hybridized with an oligonucleotide containing either a Cy3 dye (top), or a Cy3 and Cy5 dyes separated by 16 nucleotides (bottom). (B) Cy3 was illuminated with a 532 nm laser (15 mW intensity at the prism face) and fluorescent images were simultaneously collected in both the Cy3 and Cy5 channels. (C) Mean FRET efficiency from at least 100 clusters computed from five different fields-of-view. Error bars: SD. (D) Photograph of a HiSeq microfluidic chip. The HiSeq chip has eight separate lanes. We used the HiSeq 4000, which typically generates ~1-5 billion unique DNA clusters per chip. (E) A subset of fluorescent PhiX clusters imaged in a 0.26 x 0.87 mm region of the fourth lane using TIRF microscopy. This composite image is assembled from eight fields-of-view. The CHAMP image analysis pipeline was used to identify these clusters in the corresponding HiSeq sequencing (FASTQ) file. (F) An expanded view of the PhiX clusters (magenta), the aligned FASTQ coordinates image (green), and the merged image of the two (right). The aligned FASTQ coordinates are depicted as Gaussian convolutions to mimic the diffraction-limited fluorescent spots seen in TIRF microscopy. (G) Maximum cross-correlation of the TIRF image in (F) with HiSeq FASTQ tiles shows strong signal for correct alignment. Maximum cross-correlation was calculated for FASTQ tiles that neighbor the region imaged in (E). Maximum correlation of the TIRF image with incorrect FASTQ tiles is primarily a function of the density of the alignment markers and size of the tiles, and therefore relatively constant for tiles in the same lane. The signal-to-noise ratio (SNR) of the correct alignment in the correct tile (shown in red) is nearly 3, well above our relatively conservative SNR threshold of 1.4 (shown as gray background). The background noise level (SNR = 1) was determined by using the maximum cross correlation value of tiles in the same lane known not to contain the image (E).