Quantification of the affinities of CRISPR-Cas9 nucleases for cognate protospacer adjacent motif (PAM) sequences

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Running Title: Affinities of Cas9 nucleases for PAM sequences

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Abstract

The CRISPR/Cas9 nucleases have been widely applied for genome editing in various organisms. Cas9 nucleases complexed with a guide RNA (Cas9/gRNA) find their targets by scanning and interrogating the genomic DNA for sequences complementary to the gRNA. Recognition of the DNA target sequence requires a short protospacer adjacent motif (PAM) located outside this sequence. Given that the efficiency of target location may depend on the strength of interactions that promote target recognition, here we sought to compare affinities of different Cas9 nucleases for their cognate PAM sequences. To this end, we measured affinities of Cas9 nucleases from Streptococcus pyogenes, Staphylococcus aureus, and Francisella novicida complexed with gRNAs (SpCas9/gRNA, SaCas9/gRNA, and FnCas9/gRNA, respectively) and of three engineered SpCas9/gRNA variants with altered PAM specificities for short, PAM-containing DNA probes. We used a “beacon” assay that measures the relative affinities of DNA probes by determining their ability to competitively affect the rate of Cas9/gRNA binding to fluorescently labeled target DNA derivatives called “Cas9 beacons.” We observed significant differences in the affinities for cognate PAM sequences among the studied Cas9 enzymes. The relative affinities of SpCas9/gRNA and its engineered variants for canonical and suboptimal PAMs correlated with previous findings on the efficiency of these PAM sequences in genome editing. These findings suggest that high affinity of a Cas9 nuclease for its cognate PAM promotes higher genome-editing efficiency.

The type II CRISPR-Cas9 bacterial adaptive immune systems that use a single Cas9 protein programmed with a guide RNA (gRNA) to identify and degrade target DNA have been harnessed for a vast range of genome editing applications (1-6). The type IIA Cas9 nucleases from Streptococcus pyogenes (SpCas9) and Staphylococcus aureus (SaCas9) are commonly used to introduce desired mutations in eukaryotic cells. SaCas9 is smaller than SpCas9, which makes its delivery to cells using viral vectors more convenient (7,8). In contrast to SpCas9 and SaCas9, the type IIB Cas9 from Francisella novicida (FnCas9) cleaves the target DNA in a staggered pattern to leave 4-nucleotide 5'-overhangs (9). It was also reported that FnCas9 exhibits significantly higher cleavage specificity as compared with SpCas9 (10). Cas9 derivatives that lack endonuclease activity but can bind to targets...
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specifies gRNA (dCas9) are used for transcriptome modulation, base-specific genome editing and visualization of genomic loci in live cells (11-13).

Both recognition and cleavage of target DNA by Cas9/gRNA complexes require a short protospacer adjacent motif (PAM) located near the targeted sequence (14). The PAM sequences are diverse among orthologous Cas9 nucleases. The requirement for PAM sequences constrains the choice of sites amenable for genome editing. To address this issue, engineered Cas9 mutants with altered PAM specificities have been developed, but many often show editing activities lower than that of the wild-type protagonists (15-19). Crystal structures of SpCas9 and SaCas9 bound to target DNAs have revealed that these enzymes employ a major-groove PAM recognition mechanism involving direct and water-mediated hydrogen bonding interactions with cognate canonical PAMs (5'-NGG and 5'-NNGRRT for SpCas9 and SaCas9, respectively) (20,21). Genome editing experiments have showed that in vivo, PAM specificities of SpCas9 and its engineered variants are not quite strict. One well-known example is SpCas9 recognition of NAG and NGA sequences as PAM sites that can support cleavage, although significantly less efficiently than the consensus NGG PAM (22,23). The recognition of non-canonical PAM sequences should be taken into account when designing target sequences to minimize off-target effects (22). Factors that determine efficiencies of various suboptimal PAMs are not fully understood.

The Cas9/gRNA effector complexes locate a target site by scanning and interrogating the genomic DNA (24-28). The Cas9 target search in eukaryotic cells is a slow process that may limit the efficiency of genome editing, particularly when the concentration of Cas9/gRNA complexes is low to ensure specificity of target recognition (27,28). Single-molecule and bulk experiments have shown that the DNA interrogation process is triggered by Cas9/gRNA binding to PAM that leads to destabilization of the DNA duplex close to PAM and allows initial pairing of crRNA with complementary DNA bases (24,29). The initial Cas9-PAM interaction is often a target for anti-CRISPR proteins and synthetic small inhibitors (30,31). Since the rate of a target location by DNA binding proteins may depend on the strength of interactions that initiate target recognition (32), the affinity of a CRISPR-Cas effector for PAM might be one of the factors determining the efficiency of genome editing. Given the diversity of DNA-targeting CRISPR-Cas systems, it is thus of interest to compare affinities of both native and engineered CRISPR-Cas effectors for cognate PAM sequences. Here, we report a comparison of affinities of SpCas9/gRNA, SaCas9/gRNA, FnCas9/gRNA, and three engineered SpCas9/gRNA variants with altered PAM specificities (Cas9-VQR, xCas9 and Cas9-NG) for various PAMs in the context of short non-target DNA substrates bearing cognate PAM sequences. We find that affinities of the studied Cas9 effector complexes for their optimal PAM sequences differ significantly, with the highest affinity observed for SaCas9. The results show that the relative affinities of SpCas9/gRNA and its engineered variants for consensus and suboptimal PAM sequences correlate with reported effectiveness of these PAMs in genome editing, suggesting that strong binding to PAM may lead to higher genome editing efficiency. The data also suggest a difference between modes of FnCas9 and SpCas9 interaction with PAM-proximal DNA during target location.

Results

Experimental setup

Affinity of SpCas9/gRNA for canonical NGG PAM has been previously assessed by measuring SpCas9/gRNA binding to ~50 bp-long model DNA substrates containing no protospacer matching the gRNA spacer but bearing
multiple NGG sites (24,29,31). As expected, in contrast to target DNA sequences, such non-target substrates bound to Cas9/gRNA relatively weakly (24,29,31). We previously found that non-target DNA substrates bearing short duplex segments non-complementary to gRNA upstream of PAM bind to SpCas9/gRNA significantly stronger than similar substrates bearing longer upstream segments (29). The difference in the binding strength was suggested to result from obstructing the entry of duplex DNA beyond a certain threshold distance from PAM into the Cas9/sgRNA interior by an element of the effector complex. In this work, we used ~20 bp-long DNA substrates (probes) that contained a single PAM sequence and a short upstream segment which was either non-complementary or complementary to the gRNA spacer as a tool to compare affinity of Cas9/gRNA complexes for various PAM sequences. We expected that comparatively high affinity of such probes for Cas9/gRNA may facilitate detection of interactions of the Cas9 effectors with suboptimal PAM sequences. The relative affinities of DNA probes were determined by measuring their ability to competitively affect the rate of Cas9/gRNA binding to “Cas9 beacons”, fluorescently labeled target DNA derivatives containing a protospacer complementary to spacer part of gRNA and a functional PAM (33,34). Schematic representation of the beacon assay is shown in Fig. 1. Beacon 1 consists of two fully complementary oligonucleotides, while beacon 2 consists of three oligonucleotides: its nontarget strand contains a discontinuity. Cas9/gRNA binding to a beacon results in a readily measurable increase in fluorescence intensity. Beacons composed of three oligonucleotides show the same kinetics of fluorescence increase upon binding to Cas9/gRNA and dCas9/gRNA, but the kinetics of fluorescence increases observed upon Cas9/gRNA and dCas9/gRNA binding to beacons composed of two oligonucleotides may differ (33,34). The rate of beacon binding and maximal increase in fluorescence intensity depend on the beacon sequence and structure, which can be varied easily (29,35). The advantages of the beacon assay are that it reports only on functional effector complexes capable of specific binding to the beacon and allows detection of low affinity binding of a competitor to the effector complex (34). The assay also allows one to calculate the dissociation constant of Cas9/gRNA with a DNA competitor probe, provided that nearly complete beacon binding to Cas9/gRNA in the presence of competitor is reached on a suitable time scale (minutes to a few hours) (29). In practice this condition is fulfilled when the competitor concentration is not too high and its affinity for Cas9/gRNA is not too strong. Beacons used in this work were designed to have binding rates in the absence of competitors on the scale of several minutes. Sequences of competitors were adjusted to avoid formation of hairpin structures. Structures of all beacons and competitor probes are shown in Figs. S2 and S3. Competitors were preincubated with Cas9/gRNA complexes for 15 min prior to the addition of beacon.

Comparison of affinities of SpCas9 and its engineered variants with altered PAM specificities for optimal and suboptimal PAMs

The reported PAM preferences of SpCas9 and its engineered variants may correlate with their affinities for corresponding PAMs. To test this conjecture, we measured the affinities of dSpCas9, dCas9-VQR, dxCas9, and dCas9-NG effector complexes for short DNA probes bearing PAM sequences of interest using the competition Cas9 beacon assay. The engineered Cas9-VQR variant recognizes the NGAN sequence as PAM (15). The base in the fourth position affects the efficiency of Cas9-VQR genome editing. In human and plant cells the NGAG >NGAT≈NGAA> NGAC relative PAM preferences have been found (15,36). The preference for the
NGAG PAM is consistent with base-specific recognition of a guanine base in the fourth position (37). The relative affinities of dCas9-VQR/gRNA for the NGAN PAMs were compared in the context of competitor probes 1.1-1.8 (Fig. 2A). The probes contained the TGAG, TGAT, TGAA, or TGAC PAM variants and 2-bp segments upstream of PAM, which were either non-complementary (probes 1.1-1.4) or complementary (probes 1.5-1.8) to the gRNA spacer. As expected, probes bearing the complementary 2-bp segments upstream of PAM were more effective competitors than corresponding fully non-complementary probes. Probes 1.1 and 1.5 bearing the TGAG PAM were significantly stronger competitors of beacon binding and thus had higher affinities for Cas9-VQR/gRNA than probes bearing other TGAN PAM variants (Figs. 2B,C and S4). The calculated $K_d$ for probe 1.1 was 6.4±1.3 nM, whereas $K_d$ for probes 1.2-1.4 were about 200 nM. Although we did not observe a difference between competitor strengths of probes 1.2-1.4 (Fig. 2B), experiments with partially complementary probes showed that probes 1.6 and 1.7 containing the TGAA and TGAT PAM sequences, respectively, are stronger competitors than probe 1.8, which bears the TGAC PAM (Fig. 2C). The data thus show that the Cas9-VQR PAM variants known to be more efficient in genome editing have higher affinities for Cas9-VQR/gRNA, suggesting that strong binding to PAM favors Cas9-VQR genome editing.

The relative affinities of dSpCas9/gRNA for various PAM sequences were compared in the context of parent probe 2.1 bearing the canonical TGG PAM and its derivatives 2.2-2.8 in which the TGG PAM sequence was substituted for either suboptimal TGA/TAG, or nonfunctional TGT, TGC, TTG, TCG, and TAA sequences (Fig. S3). The probes bore 2-bp segments complementary to gRNA spacer immediately upstream of PAM in order to increase the strength and specificity of the binding. Probe 2.1 bearing the TGG PAM was a much stronger competitor than all other probes (Fig. 3). The beacon binding rate dropped down by about 400-fold in the presence of 200 nM of probe 2.1, whereas the addition of probes 2.2 and 2.3 with suboptimal TGA and TAG PAMs, correspondingly, caused decrease in the binding rate by only 1.7-fold. Probes 2.4-2.8 bearing non-functional PAM sequences were the weakest competitors and caused less than 25% decrease in beacon binding rate (Fig. 3).

The xCas9 and Cas9-NG variants of SpCas9 recognize the relaxed NGN sequence as PAM (though SpCas9-NG was found to be less active at NGC relative to NGD PAMs in human cells) (17,18). In addition to its broader PAM compatibility, xCas9 has considerably reduced off-target activity compared with wild-type SpCas9 (17,38,41). Available data also show that in many cases Cas9-NG is more efficient that xCas9 in inducing mutations at target sites with NGN PAMs (18,38-41). We compared affinities of dxCas9/gRNA and dCas9NG/gRNA for probes 3.1-3.5 containing AGG, AGA, AGT, AGC, and ATG PAM variants and 2-bp segments complementary to the gRNA spacer upstream of PAM (shown in Fig. 4A) using the same beacon. In the absence of competitors, dCas9-NG/gRNA bound beacon about 6-fold faster than dxCas9/gRNA (Figs. 4B and C), consistent with the higher rate of Cas9NG-induced DNA cleavage observed in in vitro assays of Cas9NG and xCas9 activities (18). In the presence of 200 nM of probe 3.1 containing the AGG PAM, the rate of beacon binding to dxCas9/gRNA decreased 7-fold whereas other probes caused ~2-fold decreases in the beacon binding rate (Fig. 4B). Much higher inhibitory effects were observed in similar experiments with dCas9-NG. The rate of beacon binding to dCas9-NG/gRNA decreased ~ 600 and 120-fold in the presence of 200 and 40 nM of probe 3.1, respectively. The calculated $K_d$ for probe 3.1
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binding to dxCas9/gRNA was about 80-fold higher than for dCas9-NG/gRNA (K_d = 30±6 nM and K_d = 0.35±0.11 nM, respectively). We compared the effects caused by probes 3.1-3.5 on beacon binding to Cas9-NG/gRNA at 40 nM probe concentrations. As can be seen from Figs. 4C and S5, the AGG PAM containing probe 3.1 was a slightly stronger inhibitor than probes 3.2 and 3.3 bearing, respectively, the AGT and AGA PAM sequences, while the inhibition effect caused by probe 3.4 (AGC PAM) was weaker than that of probe 3.1 by 4.8-fold. The latter observation is in agreement with lower Cas9-NG editing activity at NGC sites (18). Control probe 3.5 bearing a T at the second PAM position was a weaker inhibitor than probes 3.1-3.4 (Fig. 4C). The data thus show that dCas9-NG binds short DNA probes bearing the relaxed NGN PAM stronger than dxCas9, which suggests that the observed difference between xCas9 and Cas9-NG cleavage efficiencies (18,38-41) may be, at least in part, accounted for by stronger interaction of Cas9-NG with its PAM. Overall, the above data show that there is a correspondence between affinities of the studied Cas9 effectors for PAM sequences and their reported in vivo activities.

Affinity of SaCas9 for its canonical PAM is significantly higher than that of SpCas9

We compared affinities of dSpCas9 and dSaCas9 for their PAMs in the context of non-complementary to sgRNA probes 4.1-4.7 that had the same downstream boundary but different upstream edges at positions +1, -2, -3, -5, and -7 (Fig. 5A). The probes bore overlapping NGG and NNGRRG canonical PAM sequences and thus could be used in competition beacon assay experiments with both SaCas9 and SpCas9. The probes 4.1-4.3 bearing, respectively, no nucleotides and 1 or 2 bp segment upstream of PAM very strongly inhibited dSaCas9/gRNA binding to beacon at 40 nM probe concentration, while further extensions of the upstream edge to -3, -5, and -7 positions in, respectively, probes 4.4, 4.5, and 4.6 led to gradual decrease of the inhibitory effect (Fig. 5B). The substitution of a G for a T at the third PAM position of probe 4.3 almost eliminated the competition effect of resulting probe 4.7, proving that probe 4.3 binding was PAM-dependent (Fig. S6). As can be seen in Fig. 5C, similar dependence of relative probe competition strength on upstream probe boundary was found in experiments with dSpCas9, in agreement with our previous measurements performed under somewhat different conditions (29). However, the inhibition effects caused by the probes on beacon binding to dSaCas9 were much stronger than those observed with dSpCas9 (Figs. 5B and C). Consistently, the K_d for binding of probe 4.5 to dSaCas9/gRNA was 26-fold lower than for binding to dSpCas9/gRNA (3.2±0.5 and 85±21 nM, respectively). The K_d values for probe 4.3 binding to dSpCas9/gRNA and dSaCas9/gRNA were 67±0.7 and <0.4 nM, respectively.

Next, we conducted competition beacon assay measurements using as a non-target competitor the pUC19 plasmid, which contains multiple (330 NGG and 78 NNGRR) PAMs for both enzymes in various contexts. Consistent with data obtained with short probes, the dSaCas9/gRNA binding to the beacon was strongly competed by 3 nM of pUC19, whereas only about 2-fold decrease in the rate of dSpCas9/gRNA binding to the beacon was observed (Figs. 6A and B). We also measured the inhibition effects caused by probe 4.8 bearing a truncated PAM lacking the non-conserved upstream base-pair in the 5'-TGGAGT PAM segment. As can be seen in Figs. S7A and B, probe 4.8 was a significantly weaker inhibitor than probe 4.1 bearing the full PAM sequence. This observation suggests that sequence non-specific contacts of SaCas9 and SpCas9 with upstream nucleotides of their PAMs seen in crystal structures contribute noticeably to affinities towards PAMs (20,21,42). To verify the results of beacon
assays, the binding of probe bearing a 2 bp segment upstream of PAM and control probe bearing mutated PAM sequences to dSaCas9/gRNA and dSpCas9/gRNA was measured by the EMSA assay. As can be seen from Fig. S8, the probe affinity for dSaCas9/sgRNA is considerably higher than for dSpCas9/sgRNA. Overall, the above data show that dSaCas9/gRNA binds short PAM-containing substrates much stronger than does dSpCas9/gRNA, suggesting that SaCas9 affinity for PAM is significantly higher than that of SpCas9.

**Interaction of FnCas9/gRNA with short DNA probes bearing its PAM sequence**

Since FnCas9 recognizes the 5'-NGG sequence as an optimal PAM (43), we used the same competitor probes that were used with SpCas9 and SaCas9 (Fig. 5A) in competition beacon binding assays with FnCas9. We found that FnCas9/gRNA bound probe 4.1 that bears no nucleotides upstream of PAM quite strongly (calculated $K_d = 3.0\pm0.6$ nM, Figs. 7 and S9). In sharp contrast to results obtained with SpCas9 and SaCas9, extensions of the upstream edge of probe 4.1 by just 1 and 2 bp led to considerable decrease in affinities (by 7 and 30-fold, respectively) (Fig. 7, probes 4.2 and 4.3). Probes 4.4-4.6, which bore longer segments upstream of PAM, as well as probe 4.8 containing truncated PAM were very weak competitors (Figs. 7 and S7C). These data thus show a difference in the modes of FnCas9/gRNA and SpCas9/gRNA interactions with the segment of target proximal to PAM, implying differences in the process of DNA interrogation that requires destabilization of this segment of the target (24, 29).

**Discussion**

High editing efficiency is desired for many research and potential therapeutic applications of genome editing (44). We hypothesized that strong binding of a Cas9 effector to PAM during early steps of DNA interrogation might favor high efficiency of genome editing, for example, through acceleration of target location or facilitation of effector access to partially blocked targets. On the other hand, too high an affinity for PAM might delay the target search by increasing the effector’s dwell time at multiple PAM sites located throughout the genome. To elucidate the extent of variation of Cas9 nucleases affinities for PAM sequences, we compared the ability of several Cas9 nucleases to bind to short DNA probes bearing functional PAM sequences. Overall, our results reveal large differences in the affinity for PAM among the Cas9 enzymes studied. In particular, we find that affinities of probes containing canonical SpCas9 and SaCas9 PAM sequences but noncomplementary to gRNA spacer are about 20-fold lower for SpCas9 effector than for SaCas9. Given that SaCas9 was observed to form less stable R-loop complexes with target DNA than SpCas9 (45), the high affinity of SaCas9/gRNA for cognate PAM is somewhat surprising. These observations are, however, consistent with a previous study showing that R-loop stability is mainly determined by Cas9/gRNA interactions with protospacer, whereas PAM affects primarily the R-loop formation rate (46). A number of earlier studies showed that SaCas9 works with comparable efficiency to SpCas9 (7,8). Yet, recently reported measurements of SpCas9 and SaCas9 genome editing activities found that SaCas9 was noticeably more efficient in inducing mutations than SpCas9 when targeting the same sites at similar conditions (47-49). We suggest that the higher efficiency of SaCas9 observed in these studies may be in part accounted for by its relatively stronger interaction with PAM.

Comparison of affinities of SpCas9/gRNA and its engineered variants for optimal and suboptimal PAM sequences also supports the idea that there is a relationship between the PAM affinity and the efficiency of genome editing. Our data indicate that Cas9-VQR/gRNA has much higher affinity for probes bearing TGAG
PAM than for similar probes with bases other than G at the 4th PAM position, which is consistent with the preference for the NGAG PAM sequence observed in genome editing experiments (15). Further, we find that affinities of SpCas9/gRNA for suboptimal NAG and NGA PAMs are much lower than that for the canonical NGG PAM but higher than for nonfunctional PAM sequences. The finding that Cas9-NG/gRNA binds to short probes bearing the relaxed NGN PAM sequences stronger than xCas9/gRNA is also consistent with the correlation between PAM affinity and genome editing efficiency. Aminoacid substitutions introduced into xCas9 and Cas9-NG are located in different protein domains (Table S1), implying that different structural features account for their relaxed PAM recognition (17,18). The crystal structure of Cas9-NG/gRNA complex with target DNA suggests that the relatively high PAM affinity of Cas9-NG is due to non-base-specific interactions of the PAM duplex with substituted SpCas9 amino acid residues, which compensate the loss of the base-specific interaction with the third PAM nucleobase (18).

Our data show that relative affinities of SpCas9/gRNA and SaCas9/gRNA for probes bearing duplex segments non-complementary to gRNA upstream of PAM gradually decrease with increasing length of the upstream segments in a quite similar way (Figs. 5B and C). In contrast, in the case of FnCas9, we found that probe bearing no nucleotides upstream of PAM had high affinity for FnCas9/gRNA but the affinities of probes bearing even short (2-3 bp) segments upstream of PAM dropped sharply (Fig. 7). These observations suggest a difference in the modes of interaction of these effectors with the adjacent to PAM DNA duplex during DNA interrogation. In the crystal structure of FnCas9/sgRNA-DNA R-loop complex, the PAM duplex is sandwiched between the WED and PI domains which are connected by the WED–PI linker loop. As in SpCas9 and SaCas9, the backbone phosphate group between nucleotides at the +1 and -1 positions in the target DNA strand interacts with the phosphate lock loop, thereby stabilizing target DNA immediately upstream of PAM in an unwound conformation (43). The structure also shows that Arg1474 in the WED-PI linker forms a stacking interaction with the most upstream PAM base in the target DNA strand without hydrogen bond formation (43). Such stacking interaction is not observed in structures of SpCas9 and SaCas9 ternary complexes. Consideration of the FnCas9/sgRNA-DNA R-loop complex structure suggests that a 1-bp duplex extension of the PAM segment in the upstream direction would likely result in a steric clash with Arg1474. We thus suggest that the observed obstruction of DNA binding to FnCas9/gRNA by an adjacent to PAM short duplex segment may be at least in part due to its clash with the WED-PI linker, although the involvement of other neighboring segments is also possible. This suggestion is in line with a recent study on specificity of FnCas9 binding to and cleavage of DNA, which proposes that FnCas9 may possess a distinct mode of DNA interrogation and rejecting off-targets (10).

**Experimental procedures**

**Purification of Cas9 proteins**

SpCas9 variants Cas9-VQR, xCas9 3.7 and SpCas9-NG used in this work were prepared essentially as described previously (15,17,18). Details of the purification procedure are described in SI Experimental procedures. dSaCas9 protein (saCas9 Null Mutant Protein) was purchased from Applied Biological Materials, Inc. FnCas9 protein was purchased from Sigma-Aldrich. The commercial proteins showed a high batch-to-batch reproducibility in the beacon assay experiments. The concentrations of SpCas9 and its variants, of SaCas9, and of FnCas9 were determined spectrophotometrically at 280 nm using calculated extinction coefficients of
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120450, 116900, and 199750 M⁻¹ cm⁻¹, respectively.

**Preparation of guide RNAs and DNA probes**

Single guide RNAs (shown in Fig. S1) were in vitro transcribed using HiScribe T7 High Yield RNA Synthesis Kit (NEB) according to manufacturer’s protocol. The transcribed sgRNAs were purified by Bio-Gel P-30 spin-column (Bio-Rad). SpCas9 dual guide RNA (dgRNA) was formed by mixing equimolar amounts of SpCas9 tracrRNA (Sigma-Aldrich) and crRNA (Integrated DNA Technologies (IDT)) in a buffer containing 20 mM Tris, pH 7.5, 100 mM NaCl; heating for 30 s at 90 °C and slowly cooling the reactions to 20 °C. In experiments with SpCas9 and its variants, we used dgRNA since it could be readily prepared in large quantities. Control beacon assay experiments with SpCas9 sgRNA and dgRNA yielded identical results. All gRNAs used in this work contained the same spacer segment (Fig. S1). Cas9 beacon constructs (Fig. S2) and competitor DNA probes (Fig. S3) were prepared from unmodified and chromophore-labeled DNA oligonucleotides synthesized by IDT by mixing equimolar amounts of synthetic complementary strands in a buffer containing 20 mM Tris (pH 7.5) and 100 mM NaCl heated for 2 min at 90 °C, with the reactions slowly cooled to 20 °C. The PAM-distal ends of the beacon target and nontarget strands were labeled with fluorescein and Iowa Black R-FQ, respectively.

**Fluorometric measurements**

Fluorescence measurements were carried out at 25 °C using a QuantaMaster QM4 spectrofluorometer (PTI) in binding buffer [20 mM Tris HCl (pH 7.5), 100 mM NaCl, 5% glycerol, 0.1 mM DTT and 1 mM MgCl₂] containing 0.02% Tween-20. The MgCl₂ concentration used was chosen since it is close to estimates of intracellular concentration of free Mg²⁺ (50). Final assay mixtures (800 μL) contained 5-10 nM Cas9 protein, gRNA in 2-fold excess over Cas9, 0.5-1 nM Cas beacon and competitor DNA probes at various concentrations. The DNA probes were preincubated with Cas9/gRNA complexes for 15 min at 25 °C prior the beacon addition. The fluorescein fluorescence intensities were recorded with an excitation wavelength of 498 nm and an emission wavelength of 520 nm. Time dependent fluorescence changes were monitored using manual mixing; the mixing dead-time was 15 s. Competition experiments were analyzed with Felix software (PTI), Kd values were calculated as described previously (29,34). Details of the Kd calculation are also described in SI Experimental procedures. The Kd values were determined as averages obtained from at least three independent experiments performed on different days, the standard deviations were less than 30% of the corresponding mean values. Typical variations among replicate kinetic traces are shown in Fig. S5.

**Electrophoretic mobility shift assay**

DNA probe duplexes were formed by mixing 5′-[³²P]-labeled target strand (110 nM) and unlabeled nontarget strand (125 nM) in Nuclease-Free Duplex Buffer (IDT), heating for 2 min at 95 °C and slowly cooling to 20 °C. Reconstitution of the Cas9-gRNA complexes was carried out by mixing stock solutions of a dCas9 protein and dgRNA or sgRNA in 1:1.5 ratio and incubating at room temperature for 10 minutes. Target DNA binding assays were carried out in 10 μL assay buffer (20 mM Tris-HCl, pH 8.0, 25 mM NaCl, 5 mM MgCl₂, 0.25 mM TCEP) containing 10 nM DNA probe and 8-400 nM dCas9/gRNA. Binding reactions were incubated for 10 min at 37 °C, and resolved by native 6% PAGE at 4 °C (0.5×TBE buffer with 2 mM MgCl₂). DNA probes were visualized by phosphorimaging and quantified with ImageQuant (GE Healthcare).

**Data availability**
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All data generated or analyzed during this study are included in this published article and its supporting information.

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Footnotes

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This article contains SI Experimental procedures, Figs. S1-S9, Tables S1-S3.

The abbreviations used are: CRISPR, clustered regularly interspaced short palindromic repeat; Cas, CRISPR associated; PAM, protospacer adjacent motif; gRNA, guide RNA; crRNA, CRISPR RNA; Sp, *Streptococcus pyogenes*; Sa, *Staphylococcus aureus*; Fn, *Francisella novicida*; EMSA, electrophoretic mobility shift assay; $K_d$, dissociation constant.
Affinities of Cas9 nucleases for PAM sequences

Figure 1. Principle of the Cas beacon assay and structure of SpCas9 beacon. Beacon 1 consists of two fully complementary oligonucleotides, while beacon 2 has a discontinuity in its nontarget strand. The PAM-distal ends of the beacon target and non-target strands are labeled with a fluorophore (F) and fluorescence quencher (Q), respectively. The baseline fluorescence intensity of beacon is low because of proximity of the fluorophore with the quencher. The Cas9/gRNA effector complex binds to beacon in a way that mimics their binding to target DNA, which leads to separation of the fluorophore and the quencher and a readily detectable increase in fluorescence intensity. The sequence of SpCas9 crRNA and beacon used in experiments with SpCas9 and are shown below the panel. The crRNA spacer and beacon protospacer bases complementary to the spacer sequence are highlighted in blue, the mismatched protospacer bases are highlighted in pink, the PAM sequence is highlighted in yellow.
Affinities of Cas9 nucleases for PAM sequences

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**Figure 2.** Competition assays with dCas9-VQR. A, Structures of competitor DNA probes. The probes bore 2 bp segments upstream of PAM sequences that were either non-complementary (probes 1.1-1.4) or complementary (probes 1.5-1.8) to gRNA. The PAM sequences are highlighted in yellow, 2-bp segments matching the spacer sequence are highlighted in blue. B,C, Effect of competitor probes 1.1-1.4 (B) and 1.5-1.8 (C) that bore indicated PAM sequences on the kinetics of dCas9-VQR/gRNA binding to beacon. The concentrations of competitors were 200 and 40 nM in experiments shown in panels B and C, respectively.
Figure 3. Competition assays with dSpCas9 using DNA probes bearing canonical, suboptimal or nonfunctional PAM sequences. Effect of competitor DNA probes 2.1-2.5 (shown in Fig. S3) bearing indicated PAM sequences on the kinetics of dCas9/gRNA binding to beacon. The concentration of all competitor probes was 200 nM.
Affinities of Cas9 nucleases for PAM sequences

Figure 4. Competition assays with dxCas9 and dCas9-NG. A, Structures of competitor DNA probes. The probes bore 2 bp segments upstream of PAM sequences (highlighted in yellow) that were complementary to gRNA. B,C, Effect of competitor probes bearing indicated PAM sequences on the kinetics of dxCas9/gRNA (A) and dCas9-NG/gRNA (B) binding to beacon. The concentrations of competitor DNA probes were 200 and 40 nM in experiments shown in panels B and C, respectively.
Affinities of Cas9 nucleases for PAM sequences

Figure 5. Competition assays with dSpCas9 and dSaCas9 using DNA probes with upstream edges at different positions. A, Structures of competitor DNA probes. All probes bore no gRNA guide sequence complementarity. The SpCas9 PAM sequence is highlighted in yellow. B,C, Effect of probes 4.1–4.6 on the kinetics of dSaCas9/gRNA (B) and dSpCas9/gRNA (C) binding to beacons. Positions of upstream edges of the probes are indicated in the panels. The concentration of all competitor probes was 40 nM.
Affinities of Cas9 nucleases for PAM sequences

Figure 6. Competition assays with pUC19 DNA vector. Effect of pUC19 (3 nM) on the kinetics of dSpCas9/gRNA (A) and dSaCas9/gRNA (B) binding to beacons.
Figure 7. Competition assay with FnCas9 using DNA probes with upstream edges at different positions. Effect of competitor DNA probes 4.1-4.6 (shown in Fig. 5A) on the kinetics of FnCas9/gRNA binding to beacon. Positions of upstream edges of the probes are indicated in the panels. The concentration of all competitors was 200 nM.
Quantification of the affinities of CRISPR-Cas9 nucleases for cognate protospacer adjacent motif (PAM) sequences
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