Cas9, the RNA-guided DNA endonuclease from the CRISPR-Cas (clustered regularly interspaced short palindromic repeat–CRISPR-associated) system, has been adapted for genome editing and gene regulation in multiple model organisms. Here we characterize a Cas9 ortholog from *Streptococcus thermophilus* LMG18311 (LMG18311 Cas9). In vitro reconstitution of this system confirms that LMG18311 Cas9 together with a trans-activating RNA (tracrRNA) and a CRISPR RNA (crRNA) cleaves double-stranded DNA with a specificity dictated by the sequence of the crRNA. Cleavage requires not only complementarity between crRNA and target but also the presence of a short motif called the PAM. Here we determine the sequence requirements of the PAM for LMG18311 Cas9. We also show that both the efficiency of DNA target cleavage and the location of the cleavage sites vary based on the position of the PAM sequence.

A promising tool for genome manipulation (1–14) and regulation (15–18) in a wide variety of organisms has recently been identified in the RNA-guided DNA endonuclease activity of the CRISPR-Cas (clustered regularly interspaced short palindromic repeat–CRISPR-associated) system. CRISPR-Cas, an inheritable prokaryotic immune system, protects bacteria and archaea against mobile genetic elements via RNA-guided target silencing. CRISPR-Cas systems consist of an array of short direct repeats interspersed by variable invader-derived sequences (spacers) (19–21) and a *cas* operon. During invasion, small fragments of the invading DNA from phage or plasmids (proto-spacers) are incorporated into host CRISPR loci, transcribed, and processed to generate small CRISPR RNAs (crRNA). The invading nucleic acid is then recognized and silenced by Cas proteins guided by the crRNAs. There are three types of CRISPR-Cas system each characterized by the presence of a signature gene (22).

Programmed DNA cleavage requires the fewest components in the type II CRISPR-Cas system, requiring only crRNA, a trans-activating crRNA (tracrRNA), and the Cas9 endonuclease (23, 24), the signature gene of the type II system. The system can be further simplified by fusing the mature crRNA and tracrRNA into a single guide RNA (sgRNA) (23). In addition to its role in target cleavage, tracrRNA also mediates crRNA maturation by forming RNA hybrids with primary crRNA transcripts, leading to co-processing of both RNAs by endogenous RNase III (25). Cas9 contains two nuclease domains that together generate a double-strand (ds) break in target DNA. The HNH nuclease domain cleaves the complementary strand, and the RuvC-like nuclease domain cleaves the noncomplementary strand (23, 24).

A short signature sequence, named the protospacer adjacent motif (PAM), is characteristic of the invading DNA targeted by the type I and type II CRISPR-Cas systems. The PAM serves two functions. It has been linked to the acquisition of new spacer sequences, and it is necessary for the subsequent recognition and silencing of target DNA, reviewed in Ref. 26. The sequence, length, and position of the PAM vary depending on the CRISPR-Cas type and organism. PAMs from type II systems are located downstream of the protospacer and contain 2–5 bp of conserved sequence. A variable sequence, of up to 4 bp, separates the conserved sequence of the PAM from the protospacer. This variable region is often included in the definition of the PAM sequence, but for simplicity, we refer to this variable region as the linker and the conserved sequence as the PAM. To date, Cas9 from *Streptococcus pyogenes*, Cas9 from *Streptococcus thermophilus* DGCC7710, and Cas9 from *Neisseria meningitidis* have been employed as tools for genome editing or regulation. For these Cas9 orthologs, the PAMs are GG, GGNG, and GATT, and the linkers are 1, 1, and 4 bp, respectively (23, 27, 28).

The simplicity of sgRNA design and sequence-specific targeting means the RNA-guided Cas9 machinery has great potential for programmable genome engineering. Cas9 can be employed to generate mutations in cells by introducing dsDNA breaks. The capabilities of Cas9 can be expanded to various genome engineering purposes, such as transcription repression...
or activation, with its nickase (generated by inactivating one of its two nuclease domains) or nuclease null variants (15, 17, 18, 29). Another appealing possibility for the Cas9 system is to target different Cas9-mediated activities to multiple target sites, for example transcriptional repression of one gene but activation of another (30). To achieve this, multiple Cas9 orthologs will need to be employed as a single ortholog cannot concurrently mediate different activities at multiple sites (30). Therefore to broaden our understanding of Cas9 proteins, we have characterized the Cas9 ortholog from *S. thermophilus* LMG18311, which we refer to as LMG18311 Cas9. We chose to investigate Cas9 from this organism not only to increase the repertoire of Cas9 orthologs but also because it utilizes a PAM distinct from those previously characterized and its small gene size is compatible with the standard viral vectors used for delivery into exogenous systems *in vivo* (30).

Here we demonstrate that requirements for DNA cleavage *in vitro* and *in vivo* by LMG18311 Cas9 are the same as other Cas9 orthologs. We also reveal the sequence and linker length requirements of the PAM for LMG18311 Cas9. Finally, we show that the HNH and RuvC-like nuclease domains of Cas9 select the location of their cleavage sites via different mechanisms. The HNH domain catalyzes cleavage of the complementary strand at a fixed position, whereas the RuvC-like domain catalyzes cleavage of the noncomplementary strand using a ruler mechanism.

**EXPERIMENTAL PROCEDURES**

Identification of the PAM—Natural target sequences were found using the program BLAST. A single mismatch was allowed between the spacer and target sequences. Allowing more mismatches did not increase the number of sequences found. Sequences were considered unique if they were from distinct target genomes.

Cloning and Mutagenesis—The sequence encoding full-length Cas9 was PCR-amplified from *S. thermophilus* LMG18311 genomic DNA (American Type Culture Collection) and inserted into the pMAT expression vector (31, 32). The resulting construct encodes Cas9 fused to an N-terminal hexahistidine-maltose-binding protein (His6-MBP) tag. Cas9 mutants were created using the QuikChange site-directed mutagenesis method (Stratagene). To generate plasmid targets and RNA encoding vectors, synthetic oligonucleotides, bearing the appropriate sequence, were annealed and ligated into the pACYCDuet-1 (Novagen), pRSFDuet-1 (Novagen), or pMK (GeneArt). Primers and oligonucleotides are listed in Table 1. All constructs were verified by DNA sequencing.

Protein Expression and Purification—Cas9 was overexpressed in *E. coli* BL21 (DE3) with plasmids encoding Cas9 (pMAT) and sgRNA (pRSFDuet-1) or empty vectors. All plasmids, including the targets, had unique selection markers and origins of replication. The transformation assay was performed using the CaCl2 heat-shock procedure described in Ref. 33 with minor changes. The recipient cells were transformed with 5 ng of plasmid DNA and recovered in LB medium containing 0.2 mM IPTG at 37 °C for 1 h and plated on LB agar containing appropriate antibiotics and 0.2 mM IPTG. Reported transformation efficiencies are the average of at least three biological replicates. All target plasmids used in this study transformed into control recipient cells with the same efficiency (~200 colony-forming units per 5 ng of DNA).

**RNA Preparation**—RNAs were generated by *in vitro* transcription using T7 RNA polymerase. Plasmid templates were linearized overnight with EcoRI and then purified by phenol:chloroform extraction and ethanol precipitation. 0.5 μg of linear plasmid template was incubated with 0.1 mg/ml T7 RNA polymerase and 5 mM each of CTP, GTP, ATP, and UTP in reaction buffer (25 mM Tris–HCl, pH 8.0, 150 mM KCl, and 1 mM EDTA). The final fractions containing Cas9 were concentrated to ~16 mg/ml. Purified proteins were >95% pure as judged by SDS-PAGE and Coomassie Blue staining (see Fig. 1A). The mutant variants of Cas9 were expressed and purified in the same manner as the wild-type protein (see Fig. 1A).

**In Vivo Transformation Assay**—The recipient cells were prepared by co-transforming *E. coli* BL21 (DE3) with plasmids encoding Cas9 (pMAT) and sgRNA (pRSFDuet-1) or empty vectors. All plasmids, including the targets, had unique selection markers and origins of replication. The transformation assay was performed using the CaCl2 heat-shock procedure described in Ref. 33 with minor changes. The recipient cells were transformed with 5 ng of plasmid DNA and recovered in LB medium containing 0.2 mM IPTG at 37 °C for 1 h and plated on LB agar containing appropriate antibiotics and 0.2 mM IPTG. Reported transformation efficiencies are the average of at least three biological replicates. All target plasmids used in this study transformed into control recipient cells with the same efficiency (~200 colony-forming units per 5 ng of DNA).

**Plasmid Cleavage Assay**—Cas9 (25 nM), tracrRNA (25 nM), and crRNA (25 nM) were incubated in a cleavage buffer (20 mM HEPES, pH 7.5, 150 mM KCl, 10 mM MgCl2) at 37 °C for 30 min. The reactions were initiated by adding plasmid targets (4 nM), incubated at 37 °C for 30 min, and quenched with phenol. The aqueous layer was extracted and separated on a 0.8% agarose gel. Gels were stained by soaking in 1X Tris-Acetate-EDTA buffer supplemented with 5 μg/μl ethidium bromide for 1 h and then for a further hour in 1X Tris-Acetate-EDTA buffer. Bands were visualized using an FLA-7000 (Fuji) and quantified with ImageGauge (Fuji). To account for the different binding affinity of ethidium bromide to linear and supercoiled DNA, control samples with equal amounts of DNA in both forms by centrifugation at 18,000 rpm at 4 °C for 45 min, and the supernatant was loaded on a 5-ml immobilized metal chromatography column (Bio-Rad) charged with nickel sulfate. The column was washed with lysis buffer, and bound protein was eluted with lysis buffer containing 250 mM imidazole. The elution was run on a HiLoad 26/60 S200 size exclusion column (GE Healthcare) pre-equilibrated with gel-filtration buffer A (20 mM Tris–HCl, pH 8.0, and 500 mM NaCl). Fractions containing His6-MBP tagged Cas9 were collected and treated with tobacco etch virus protease overnight at 4 °C to remove the His6-MBP tag. Samples were reapplied to immobilized metal affinity chromatography resin to remove the His-tagged tobacco etch virus protease, free His6-MBP, and any remaining tagged protein. The flow-through was collected, concentrated using an Ultra-cel 10K centrifugal filter unit (Millipore), and further purified by size exclusion chromatography in gel-filtration buffer B (20 mM Tris–HCl, pH 8.0, 200 mM KCl, and 1 mM EDTA). The final fractions containing Cas9 were concentrated to ~16 mg/ml. Purified proteins were >95% pure as judged by SDS-PAGE and Coomassie Blue staining (see Fig. 1A). The mutant variants of Cas9 were expressed and purified in the same manner as the wild-type protein (see Fig. 1A).
### TABLE 1
Primers and oligonucleotides used in these studies

**Characterization of S. thermophilus LMG18311 Cas9**

| Primers for LMGB18311 Cas9 gene amplification: | Sequence (5’ to 3’) |
|-----------------------------------------------|---------------------|
| Forward                                       | GTGTGGTCCATGGGAAAGTGACTTAGTTAGTTAGGACTTTG |
| Reverse                                       | GTGTGGTCTCAGGTAAATCTAGCTTGGGCTTATC       |

| Primers for site directed mutagenesis: | Sequence (5’ to 3’) |
|---------------------------------------|---------------------|
| D9A forward                           | GTGACTTAGTTTTAGGACTTGCTAATCGGTATAGGTTCAGT |
| D9A Reverse                           | CAACAGCACATCTACCGATAGCAAGGTCTCAAACTAAGTCAC |
| H599A forward                         | CCTAACATGTTTGAAGTAGATGCTATTTTACCTTTTACCTTAC |
| H599A Reverse                         | GTGATAGAAGAGGGTAAATAAGCAGATCTACTTAAAAGATTAGG |

| Oligonucleotides used to construct sgRNA plasmid: | Sequence (5’ to 3’) |
|---------------------------------------------------|---------------------|
| Forward                                           | CATGCTAACCCCGTATGTCAGAGAGGTTTTGTAATCTCGAAATCTTGGC |
| Reverse                                           | GAAAGCTACAAGGTAAGGGCTTCATGCGGAAATC |

| Oligonucleotides used as the template for in vitro transcription of sgRNA: | Sequence (5’ to 3’) |
|---------------------------------------------------------------------------|---------------------|
| S1 Forward                                                                 | GAAAATTAATACGACTCAGATTTAGGTACCGGTATGTCAGAGAGGTTTTT |
| S1 Reverse                                                                 | GATTTCGCGCATAGGGCTCTATCTTTGTAGCTTTGCAAGAGTTTTTTT |
| S2 Forward                                                                 | GAAAATTAATACGACTCAGATTTAGGTACCGGTATGTCAGAGAGGTTTTT |
| S2 Reverse                                                                 | GATTTCGCGCATAGGGCTCTATCTTTGTAGCTTTGCAAGAGTTTTTTT |

| RNA Oligonucleotides: | Sequence (5’ to 3’) |
|----------------------|---------------------|
| crRNA                | CUACCCCGUAUGUCAGAGAGGUUUUUGUACUCUACAGAAAUUUA |
| tracrRNA             | AAUCUUUGCAAGAGCUACAAAGAUAAGGCUUCAUGCCGAAAUC |

| Oligonucleotides used to construct the plasmid targets: | Sequence (5’ to 3’) |
|--------------------------------------------------------|---------------------|
| S1 top strand                                          | CATGCATTGGCTACTGCATCAAGGGGCGAGCTTGAGTGGTACGTCAGA |
| S1 bottom strand                                       | AAATTTGCATGAGTATGAGCTACGGCGCGTGATGTGACAGAGT |
| S2 top strand                                          | CATGCATTGGCTACTGCATCAAGGGGCGAGCTTGAGTGGTACGTCAGA |
| S2 bottom strand                                       | AAATTTGCATGAGTATGAGCTACGGCGCGTGATGTGACAGAGT |

| Oligonucleotides used to construct synthetic DNA targets: | Sequence (5’ to 3’) |
|----------------------------------------------------------|---------------------|
| Top strand                                               | CATTTCGCAACTCTGACATCGGACAGCCGAGCTGTGACGTCAGA |
| Bottom strand                                            | TCATGAGTATGAGCTACGAGTCAAGGCGTATAATGTCAGAGTTGCGAATAAGT |
Characterization of *S. thermophilus* LMG18311 Cas9

Electrophoresis Mobility Shift Assay—DNA oligonucleotides were purified on 10% denaturing polyacrylamide gels. dsDNA targets (Table 1) were made by annealing each strand and purified on 12% native polyacrylamide gels containing 1× Tris-borate-EDTA. dsDNA were 5′ end-labeled with [γ-32P]ATP using T4 polynucleotide kinase (New England Biolabs). A fixed concentration (10–100 pm) of labeled dsDNA targets was mixed with an increasing concentration of premixed Cas9Δ9AΔ599A-sgRNA complex. Binding assays, performed in buffer (20 mM HEPES, pH 7.5, 150 mM KCl, 10 mM MgCl2, 0.1 mg/ml BSA, and 10% glycerol), were incubated at 37 °C for 30 min followed by separation on 5% native polyacrylamide gels. Gels were visualized by phosphorimaging (Fuji) and quantified using ImageGauge (Fuji). Fraction of DNA bound was plotted versus concentration of Cas9, and data were fit to a one-site binding isotherm using GraphPad Prism software. Reported *Kd* values are the average of at least three replicates.

RESULTS

Identifying the PAM for LMG18311 Cas9—The genome of *S. thermophilus* LMG18311 contains two CRISPR-Cas systems, of type II-A and III-A, each associated with a CRISPR loci: CRISPR-1 and CRISPR-2, respectively. The first study of PAM sequences identified a putative PAM for *S. thermophilus* as RYAAA (where R is a purine and Y is a pyrimidine) (19). This sequence was found in natural target sequences matching 41 spacers collected from 13 different *S. thermophilus* strains, including LMG18311. Subsequent studies showed that PAM sequences vary greatly, even between different strains (reviewed in Ref. 26). To confirm the PAM sequence for LMG18311 Cas9, we performed BLAST searches to identify potential protospacers in viral and plasmid genomes that matched any of the 33 spacer sequences from CRISPR-1. This search generated 41 unique target sequences, from the genomes of bacteriophage known to infect *S. thermophilus*. We then aligned 50-nucleotide segments from the identified target genomes, inclusive of the 30-nucleotide protospacer and 10-nucleotide flanking regions (Fig. 1B). In agreement with the previous study (19), inspection of this alignment clearly identified a 5-bp PAM with a consensus sequence, GCAAA, invariantly located 2 bp downstream of the protospacer (Fig. 1, B and C). The most commonly observed PAM sequence, found in 7 of the 41 target sequences, was GCAAA.

To confirm that the identified PAM was functional, we used a previously described transformation assay in which *E. coli* cells containing an exogenous type II CRISPR-Cas system are resistant to plasmid transformation, whereas cells lacking the system are competent for transformation (33, 35) (Fig. 2A). To generate cells containing the type II CRISPR-Cas system (CRISPR+ cells), compatible vectors encoding either LMG18311 Cas9 or its cognate sgRNA, engineered to contain a 20-nucleotide sequence derived from the first spacer of CRISPR-1 (Fig. 1C), were co-transformed into *E. coli* BL21(DE3). In this overexpression system, the Cas9 and sgRNA genes are under the control of an IPTG-inducible T7 promoter. Control cells lacking the CRISPR-Cas system (CRISPR− cells) were generated by co-transforming compatible empty vectors into *E. coli* BL21(DE3). We constructed a target and two control plasmids. The target plasmid contained protospacer-1 (whose sequence was identical to the first spacer of CRISPR-1), a 2-bp linker, and the identified PAM (GCAAA) (Fig. 1C). The first control plasmid contained only protospacer-1, whereas the second control plasmid lacked both protospacer-1 and PAM. The target and control plasmids were then tested for CRISPR-Cas silencing by transformation into the CRISPR− and CRISPR+ strains in the pres-

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FIGURE 1. The type II CRISPR-Cas system of *S. thermophilus* LMG18311. A, Coomassie Blue-stained SDS-polyacrylamide gel of Cas9, Cas9 D9A, Cas9 D599A, and Cas9 D9A,D599A. B, logo plot revealing the PAM for LMG18311 Cas9. The positions of the protospacer, PAM, and linker are indicated. C, schematic representation of the crRNA (green), tracrRNA (blue), and DNA target (black). The positions of the protospacer, PAM, and linker are indicated. The site at which the crRNA and tracrRNA are fused to generate the sgRNA is indicated with a dotted line.

were loaded on the same gel. The ratios of the fluorescence intensities of linear and supercoiled bands were measured and used to calculate a correlation coefficient *K* (34),

\[
K = \frac{I_{sc}}{I_{lin}} \quad \text{(Eq. 1)}
\]

where *I*<sub>lin</sub> and *I*<sub>sc</sub> are the intensities of the linear and supercoiled bands, respectively. In our case, *K* was determined to be 0.4 ± 0.05 and did not vary significantly between experiments. The percentage of linear product was then calculated as follows (34).

\[
\text{Percentage Linear} = \frac{I_{lin}}{K + I_{lin}} \times 100 \quad \text{(Eq. 2)}
\]
ence of IPTG and the appropriate antibiotics (Fig. 2A). The control plasmids transformed into both strains with similar efficiency (Fig. 2B). The target plasmid failed to transform into the CRISPR\(^{-}\) cells but transformed into the CRISPR\(^{+}\) cells with an efficiency comparable with that of the control plasmids (Fig. 2B). All of the transformation efficiencies were comparable with those previously reported (35). These results indicate that the identified PAM is functional in vivo and that the type II CRISPR-Cas system of S. thermophilus LMG18311 protects E. coli cells from transformation by plasmid DNA.

**Both the PAM Sequence and the Linker Length Are Important for Plasmid Interference**—To investigate the PAM sequence requirements for LMG18311 Cas9, we transformed a series of plasmid targets harboring single-nucleotide mutations throughout the PAM sequence in the CRISPR\(^{-}\) strain (Fig. 2C). Only the plasmid containing a mutation at the position 1 guanosine (that is, the PAM nucleotide closest to the protospacer) was transformed, albeit with a reduced (~66%) transformation efficiency as compared with the intact PAM sequence (Fig. 2C). Plasmids containing single mutations to any of the other four positions were resistant to transformation (Fig. 2C). These results indicate that the guanosine at position 1 is important for PAM function but individually the four other positions have little effect on PAM function.

A 2-bp linker separates the protospacer from the PAM for LMG18311 Cas9 (Fig. 1, B and C). To investigate how linker length affects Cas9 activity, we generated plasmid targets with linkers ranging from 0 to 5 bp in length (Fig. 2D). We then determined the transformation efficiency for these plasmids into the CRISPR\(^{-}\) cells. The CRISPR\(^{-}\) cells were equally resistant to transformation by a plasmid target with a linker length of either 2 bp or 3 bp (Fig. 2D). Plasmids with other linker lengths transformed with efficiencies more similar to the control plasmid (Fig. 2D), suggesting that plasmids with these linkers were able to escape CRISPR-Cas silencing.

**In Vitro Reconstitution Recapitulates in Vivo Activity**—To further investigate the requirements of PAM sequence and linker length, we reconstituted the activity of LMG18311 Cas9 in vitro. LMG18311 Cas9 was expressed and purified from E. coli (Fig. 1A). A 42-nucleotide tracrRNA mimicking the processed tracrRNA and a 42-nucleotide crRNA containing the sequence derived from first spacer of CRISPR-1 (Fig. 1C) were chemically synthesized. Plasmid targets were incubated with Cas9, tracrRNA, and crRNA and then analyzed by electrophoresis through agarose gels and ethidium bromide staining. As observed for other Cas9 orthologs, cleavage of the plasmid target occurred in the presence of Cas9, tracrRNA, crRNA, and Mg\(^{2+}\) (Fig. 3A). Cleavage also occurred when an sgRNA was
substituted for the tracrRNA and crRNA (Fig. 3B). As expected, cleavage was dictated by the sequence of the sgRNA (Fig. 3C). Cas9 variants with active site mutations in either the RuvC-like domain (D9A) or the HNH domain (H599A) nicked the plasmid targets, whereas a variant with a double mutation (D9A,H599A) displayed no activity (Fig. 3D). Cleavage assays using short oligonucleotide substrates confirmed that the HNH domain cleaves the strand complementary to the guide RNA, whereas the RuvC-like domain cleaves the noncomplementary strand (Fig. 3E). Mapping the location of the cut sites revealed that, as seen with other Cas9 orthologs (23, 24, 36, 37), cleavage of both strands occurs within the protospacer, 3 bp from its PAM proximal end, producing a blunt-end dsDNA break (Fig. 3F).

We next wished to confirm that either mutations in the PAM or changes in linker length had the same effect on DNA interference in vitro as they did in vivo. Therefore we monitored cleavage of these variant plasmids by recombinant LMG18311 Cas9. The fraction plasmid cleaved was calculated using the procedure detailed under “Experimental Procedures,” which accounts for the different binding affinity of ethidium bromide to linear and supercoiled DNA. Consistent with the in vivo results, mutation of the guanosine at position 1 had the greatest effect, and individual mutations to the other four positions of the PAM had only a modest effect on plasmid cleavage (Fig. 3G).
Characterization of *S. thermophilus* LMG18311 Cas9

Cleavage of plasmid targets with different linker lengths was optimal at 2 or 3 bp and then decreased steadily with increasing or decreasing lengths (Fig. 3G).

**Metal Dependence of DNA Cleavage by Cas9**—To evaluate whether other divergent cations besides Mg$^{2+}$ can activate DNA cleavage by Cas9, we performed plasmid cleavage assays in the presence of one of the following divergent cations: Ca$^{2+}$, Mn$^{2+}$, Co$^{2+}$, Ni$^{2+}$, and Cu$^{2+}$. Reactions containing Ca$^{2+}$ yielded nicked, instead of linear plasmid (Fig. 4A), suggesting that Ca$^{2+}$ activates only one of the Cas9 nuclease domains. To identify which domain was activated, we assayed the single active site mutants of Cas9 (D9A or H599A) in a reaction buffer containing Ca$^{2+}$. We observed little cleavage with the HNH mutant (H599A) but robust cleavage with the RuvC-like mutant (D9A) (Fig. 4B), suggesting that the HNH but not the RuvC-like domain was activated by Ca$^{2+}$. None of the other divergent cations tested activated either nuclease domain of Cas9 (Fig. 4A).

*Both the PAM Sequence and the Linker Length Are Important for Target Binding*—Previous studies indicate that mutations within the PAM impair DNA cleavage by Cas9 due to weakened binding (23, 24, 38). To determine the effect of PAM sequence and linker length on binding of LMG18311 Cas9 to DNA targets, we determined the binding affinity ($K_d$) of the Cas9-sgRNA complex to 5’-end labeled dsDNA targets using native gel electrophoresis (Fig. 5A). Binding experiments were conducted with the nuclease-deficient mutant of Cas9 (D9A,H599A) in the presence of Mg$^{2+}$. Fixed concentrations of the dsDNA targets were incubated with increasing concentrations of the Cas9-sgRNA complex (Fig. 5A). A target containing a complementary protospacer, a 2-bp linker, and a functional PAM bound to Cas9-sgRNA with an affinity of 0.94 ± 0.27 nM (Fig. 5B). We were unable to detect binding to a target containing a noncomplementary protospacer or to a target that lacked a PAM. Mutation of the guanosine at position 1 of the PAM resulted in an ~100-fold increase in $K_d$ (Fig. 5B), whereas mutations at positions 2 through 5 did not significantly alter the affinity (all within ~4-fold on the consensus PAM) (Fig. 5B). Changes in linker length had a larger effect on binding affinity (Fig. 5C). Under the conditions tested, we failed to detect binding to plasmid targets containing linker lengths of 0, 4, or 5 bp ($K_d > 1000$ nM), whereas linkers of 1 and 3 bp reduced the affinity by ~400- and ~20-fold, respectively (Fig. 5C).

**HNH and RuvC-like Domains Determine the Location of Their Cut Sites Using Different Mechanisms**—Previous studies reported that Cas9 cleaves both DNA strands within the protospacer, 3 bp from its PAM proximal end, producing a predominantly blunt-end dsDNA break (23, 24, 36, 37). To determine whether linker length has any effect on where the Cas9 nuclease domains cut, we mapped the location of the cut sites in plasmids containing protospacer-1 and different lengths of linker. Following cleavage by Cas9 (programmed with an sgRNA complementary to protospacer-1), the linear plasmid products were purified by agarose gel electrophoresis and sequenced. Sequencing data revealed that the position of the cleavage site on the noncomplementary strand, but not on the complementary strand, depended on linker length (Fig. 6A). Cleavage of the complementary strand always occurred 3 nucleotides from the 5’ end of the protospacer sequence, independent of the linker length (Fig. 6A). In contrast, cleavage of the noncomplementary strand occurred predominantly 5 nucleotides from the 3’ end of the PAM with linker lengths of 2 or more bp or at 4 and 5 nucleotides from the 3’ end of the PAM with a linker length of 1 bp (Fig. 6A). The site of cleavage on both strands of the DNA target was also found to be independent of spacer sequence. The location of Cas9 cut sites in plasmids containing protospacer-2 was found to be identical to plasmids containing protospacer-1 for all linker lengths investigated (Fig. 6B). We were unable to generate enough cleaved DNA from the plasmid target with a linker length of zero for sequencing.

**DISCUSSION**

Cas9, the RNA-guided endonuclease from the type II CRISPR-Cas system, has the potential to revolutionize our ability to manipulate the genomes of a wide variety of organisms (1–18). Targeting Cas9 to specific genomic sites relies on the presence of a PAM and complementarity between the sequence of its crRNA and the protospacer. A remarkably diverse set of PAM sequences is recognized by Cas9 orthologs (30). To date, PAM recognition and DNA cleavage have been experimentally studied in only a handful of Cas9 orthologs (23, 24, 28, 30).
Characterization of additional orthologs is expected to improve our mechanistic understanding of Cas9 and likely expand our engineering capabilities. Here we present characterization of the Cas9 protein from *S. thermophilus* LMG18311.

We demonstrate LMG18311 Cas9 is active in vivo through transformation assays (Fig. 2) and *in vitro* by monitoring plasmid cleavage (Fig. 3). We also confirm that the PAM for LMG18311 Cas9 identified by sequence alignments is functional (Fig. 1 B). As observed for other Cas9 orthologs, LMG18311 Cas9 activity requires tracrRNA, crRNA, and Mg^{2+} (Fig. 3). Metal ion substitution studies also reveal that Ca^{2+} likely activates the HNH but not the RuvC-like domain of LMG18311 Cas9 (Fig. 4 B). Here however, we cannot rule out the possibility that the observed activation of the HNH domain may be due to trace Mg^{2+} contamination in the Ca^{2+} solution.

Cas9 orthologs have been reported to cleave target DNA with a wide range of mutations in the PAM sequences (30). However, in natural targets, PAM sequences are highly conserved.

**FIGURE 6.** Mapping the Cas9 cleavage sites in plasmid targets with different linker lengths. A and B, direct sequencing electropherograms for plasmid-sp1 (A) and plasmid-sp2 (B) from complementary strand primer (bottom left) and noncomplementary strand primer (bottom right) are shown. Termination of primer extension in the sequencing reactions reveals the positions of the cleavage site (red lines). The positions of the protospacer, PAM, and linker are indicated. The 3'-terminal A addition, indicated by *asterisk*, is an artifact of the sequencing reaction.
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apparent discrepancy may arise from the dual function of the PAM (26, 30). The stringency on the PAM sequence is greater for spacer acquisition than for DNA cleavage by Cas9. Consistent with this, our results show that although the PAM for LMG18311 Cas9 is conserved (Fig. 1B), the nuclease activity of LMG18311 Cas9 tolerates a broad range of mutations in the PAM of the target DNA. Mutations to the guanosine at position 1 impair Cas9 activity, whereas individual mutations at positions 2 through 5 have little effect. The PAM for N. meningitidis Cas9 also contains a single guanosine important for Cas9 activity. In addition, two recent in vivo studies show that an AG sequence can partially replace the consensus PAM, GG, for S. pyogenes Cas9 (13, 39). Thus, despite the varying sequence of PAM, Cas9 proteins from LMG18311, S. pyogenes, and N. meningitidis all contain a guanosine that appears essential for DNA silencing in vivo.

A previously unexplored aspect of target binding and cleavage by Cas9 is the length of the linker between the PAM and protospacer. The 41 natural targets of LMG18311 Cas9 we identified in our sequence searches all contain a 2-bp linker. However, we found that DNA containing a 3-bp linker was silenced with the same efficiency as that with a 2-bp linker (Figs. 2D and 3F). Further lengthening or shortening of the linker eliminates CRISPR-Cas silencing and inhibits plasmid cleavage (Figs. 2D and 3F). Thus, our results on the type II system of S. thermophilus LMG18311 suggest that the requirements for the length of the linker appear to be less stringent for DNA silencing than for spacer acquisition, a pattern similar to that observed for requirements on PAM sequence.

Recognition of target DNA by either Cas9 or effector complexes from the type I CRISPR-Cas systems is thought to be a multistep process (23, 24, 38, 40, 41). First, cellular DNA is scanned for PAM sequences. Once a PAM is identified, the adjacent DNA duplex is destabilized, enabling Cas9 to probe sequence complementarity on the target strand. Target recognition is completed if this adjacent sequence contains a protospacer that can base-pair with the crRNA, stabilizing the complex. If this sequence lacks a protospacer, then the crRNA-DNA heteroduplex fails to form and Cas9 dissociates. We found the affinity of LMG18311 Cas9-sgRNA for its target sequence is ~1.0 nM (Fig. 5), which is similar to the K_d of ~0.5 nM reported for S. pyogenes Cas9 (38) and comparable with the affinity of the type I effector complexes for their DNA targets (42–44). Targets lacking a PAM had no detectable affinity for Cas9. As expected (23, 24), the impaired nuclease activity of LMG18311 Cas9 observed when PAM sequences are mutated arises from the weakened binding affinity between Cas9 and target DNA (Fig. 5B). Further analysis also revealed that the inhibition of cleavage of targets with different linker lengths was also due to weakened affinity (Fig. 5C). Although both PAM and linker mutations result in reduced target affinity, they likely affect different steps in binding. PAM mutations inhibit the initial recognition of a target sequence, whereas altering linker length likely impairs the efficiency of base-pairing between crRNA and the protospacer, thus destabilizing the complex.

The length of the linker between the PAM and protospacer affects both the efficiency of DNA target cleavage and the position of the cleavage sites. This suggests that the two nuclease domains of Cas9 select their cleavage sites by different mechanisms. The HNH domain cleaves the complementary strand at a fixed position, whereas the RuvC-like domain, employing a ruler mechanism, cleaves the noncomplementary strand at a position measured from the PAM (Fig. 6). These observations suggest that the relative positions of the Cas9 nuclease domains are highly flexible.

While this manuscript was in preparation, crystal structures of Cas9 from S. pyogenes and Actinomyces naeslundii (45) and Cas9 from S. pyogenes in complex with sgRNA and its ssDNA target (46) were reported. The domain organization observed in these structures is consistent with our data showing that the two nuclease domains of Cas9 select their cleavage sites by different mechanisms. These structures reveal that Cas9 adopts a bilobed architecture composed of target recognition and nuclease lobes. The target recognition lobe is essential for binding the sgRNA and the complementary strand of the DNA target. The nuclease lobe contains a C-terminal domain implicated in PAM binding (45, 46) as well as the HNH and RuvC-like nuclease domains. The position of the RuvC-like domain is fixed relative to the position of the PAM binding domain, supporting our observation that cleavage of the noncomplementary strand by the RuvC-like domain occurs at a fixed distance from the PAM (Fig. 7). In contrast, the position of the HNH domain is variable among the current structures (45, 46). In the structure of Cas9-sgRNA bound to ssDNA, which is in an inac-
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