Integrase-mediated spacer acquisition during CRISPR–Cas adaptive immunity

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Bacteria and archaea insert spacer sequences acquired from foreign DNAs into CRISPR loci to generate immunological memory. The Escherichia coli Cas1–Cas2 complex mediates spacer acquisition in vivo, but the molecular mechanism of this process is unknown. Here we show that the purified Cas1–Cas2 complex integrates oligonucleotide DNA substrates into acceptor DNA to yield products similar to those generated by retroviral integrases and transposases. Cas1 is the catalytic subunit and Cas2 substantially increases integration activity. Protopspacer DNA with free 3'-OH ends and supercoiled target DNA are required, and integration occurs preferentially at the ends of CRISPR repeats and at sequences adjacent to cruciform structures abutting AT-rich regions, similar to the CRISPR leader sequence. Our results demonstrate the Cas1–Cas2 complex to be the minimal machinery that catalyses spacer DNA acquisition and explain the significance of CRISPR repeats in providing sequence and structural specificity for Cas1–Cas2-mediated adaptive immunity.

Prokaryotic adaptive immunity relies on clustered regularly interspaced short palindromic repeats (CRISPRs) together with CRISPR associated (Cas) proteins to detect and destroy foreign nucleic acids1–7. CRISPR loci contain an AT-rich leader sequence followed by repetitive sequence elements flanking ~30 base pair (bp) spacer segments that are transcribed to produce precursor CRISPR RNAs (pre-crRNAs)3–5. Spacers are frequently virus- or plasmid-derived, although ‘self’-derived spacers from the host chromosome are present in some CRISPR loci6. After pre-crRNA processing and assembly with Cas proteins, the resulting surveillance complexes target and cleave foreign nucleic acids bearing sequences complementary to the crRNA spacer sequence7–12. How spacer DNA sequences, termed protospacers, are acquired into the host CRISPR locus remains unknown.

Overexpression of Cas1 and Cas2 nucleases, the only Cas proteins found in all CRISPR–Cas systems, leads to the site-specific acquisition of 33 bp protospacers at the leader end of the CRISPR locus in E. coli13–15. Furthermore, Cas1 and Cas2 function as a complex in vivo16, suggesting that the Cas1–Cas2 complex might possess DNA recombination activity. We reconstituted CRISPR spacer acquisition using purified Cas1 and Cas2 proteins, protospacers and acceptor plasmid DNA, revealing an elegant mechanism in which both the sequence and structural elements of the CRISPR repeats specify spacer integration sites.

Protopspacer DNA integration by Cas1–Cas2

To test whether the Cas1–Cas2 complex is sufficient to catalyse DNA recombination in vitro, assays were conducted using purified Cas1–Cas2 complex, 33 bp protospacer DNA and an acceptor ‘target’ plasmid consisting of the pUC19 backbone with an inserted CRISPR locus (pCRISPR) (Fig. 1a). Co-incubation of these reagents converted the supercoiled plasmid into three main products: relaxed and linear plasmid species and a fast-migrating species we term band X (Fig. 1b, c and Extended Data Fig. 1a). Product formation required Cas1, Cas2 and the protospacer DNA (Extended Data Fig. 1b–d), and was consistent with previous divergent metal ion-dependent and sequence-nonspecific in vitro activity requirements of Cas1 (refs 17–19) and Cas2 (refs 20–22). Product DNA migration was not affected by treatment with EDTA, EDTA and phenol–chloroform extraction or proteinase K in the presence of EDTA and detergent (Extended Data Fig. 1e), indicating that product DNAs are unlikely to be bound to Cas1 and/or Cas2. Consistent with product DNA resulting from covalent integration of protospacer DNA into the plasmid, the relaxed and linear forms of pCRISPR became radiolabelled in reactions containing 32P-labelled protospacer DNA (Fig. 1d and Extended Data Fig. 2). Although Cas1 alone catalysed a low level of protospacer integration in the presence of Mn2+, the reaction was enhanced substantially by the presence of Cas2 (Extended Data Fig. 2b).

Bacteria expressing Cas1 active-site mutants, but not Cas2 active-site mutants, are incapable of acquiring new spacers in vivo, demonstrating the catalytic role of Cas1 during spacer acquisition13,14,16. Consistent with these data, Cas1 active site mutants H208A and D221A were defective for protospacer integration in vitro, whereas the Cas2(E9Q) active-site mutant supported integration (Fig. 1c, e and Extended Data Fig. 3). The Cas2 C-terminal (Δβ6–β7) deletion mutant, which is defective for complex formation with Cas1 and spacer acquisition in vivo, failed to support Cas1-mediated integrase activity (Fig. 1c, e). We conclude that our in vitro assay recapitulates the in vivo functions of Cas1 and Cas2 during spacer acquisition.

Integration and disintegration products

We tested whether the reaction products of Cas1–Cas2-mediated DNA integration resemble those formed by the strand transfer activity of retroviral integrases and cut-and-paste transposases23–26. These enzymes generate two main products in vitro corresponding to half-site and full-site integration events (Fig. 2a). We observed similar gel mobility of the slowly migrating DNA product generated by Cas1–Cas2 and Nb.BbvCI nickase-digested pCRISPR, consistent with the slow-migrating relaxed DNA species corresponding to half-site products and/or products resulting from full-site integration of one protospacer molecule (Extended Data Fig. 1a). Digestion with EcoRI, which cuts pCRISPR once, converted the reaction products to linear DNAs (Fig. 2b, compare lane 4 to lane 2, and Fig. 2c). We therefore conclude that both the relaxed and band X DNA products comprise unit-sized pCRISPR circles.

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We observed that band X did not become radiolabelled in reactions conducted with 32P-labelled protospacer DNA. A time-course analysis revealed relaxed DNA product formation within the first minute, followed by accumulation of band X between 10 and 30 min (Fig. 2d). To determine the properties of band X, the purified product was analysed in two different types of agarose gels—one pre-stained with ethidium bromide after electrophoresis (pre-stained) (Extended Data Fig. 3a), and another stained agarose gel analysis was performed on the entire integration reaction (Fig. 2b, c). The Cas1–Cas2 complex accommodated various protospacer lengths in vitro (Fig. 3a, b). The Cas1–Cas2 complex integrated DNA substrates with 9-OH protospacer ends and 9-OH on the target DNA phosphodiester bonds. We found that phosphorylation of both 3′-ends of the protospacer ablated integration, whereas phosphorylation of only one 3′ end strongly limited integration (Fig. 3a, b). By analogy to known integrase enzyme mechanisms, DNA integration could proceed by Cas1-catalysed direct nucleophilic attack of the substrate 3′-OH on the target DNA, or by formation of a Cas1–DNA intermediate, as occurs in the serine and tyrosine families of recombinases. Four tyrosine residues in the vicinity of the Cas1 active site could be involved in forming such a covalent intermediate (Extended Data Fig. 7a, b). Purified Cas1 mutant proteins in which each tyrosine was individually changed to alanine were non-viable (Extended Data Fig. 6b). In contrast to retroviral integrases, substrates with 5′-overhangs up to 5 nt in length (Extended Data Fig. 6b). In contrast to retroviral integrases, substrates with 5′-overhangs were non-viable (Extended Data Fig. 6b).

Integration requires 3′-OH protospacer ends

We next investigated the DNA protospacer and target DNA requirements for integration. Single-stranded protospacer DNA failed to support the reaction (Fig. 3a, b). The Cas1–Cas2 complex accommodated various protospacer lengths in vitro despite the strict 33 bp requirement for spacer acquisition in vivo (Extended Data Fig. 4a), suggesting that protospacer length is pre-determined before integration in vivo by an unknown mechanism. The Cas1–Cas2 complex integrated DNA substrates with blunt-ends or with 3′-overhangs up to 5 nt in length (Extended Data Fig. 6b). In contrast to retroviral integrases, substrates with 5′-overhangs were non-viable (Extended Data Fig. 6b).

Retroviral integration and transposition reactions proceed via nucleophilic attack of DNA 3′-OH groups at target DNA phosphodiester bonds. We found that phosphorylation of both 3′-ends of the protospacer ablated integration, whereas phosphorylation of only one 3′ end strongly limited integration (Fig. 3a, b). By analogy to known integrase enzyme mechanisms, DNA integration could proceed by Cas1-catalysed direct nucleophilic attack of the substrate 3′-OH on the target DNA, or by formation of a Cas1–DNA intermediate, as occurs in the serine and tyrosine families of recombinases. Four tyrosine residues in the vicinity of the Cas1 active site could be involved in forming such a covalent intermediate (Extended Data Fig. 7a, b). Purified Cas1 mutant proteins in which each tyrosine was individually changed to alanine were non-viable (Extended Data Fig. 6b). In contrast to retroviral integrases, substrates with 5′-overhangs were non-viable (Extended Data Fig. 6b).

Figure 1 | The Cas1–Cas2 complex integrates protospacers in vitro. a, Schematic of the in vitro integration assay (PDB code 4P6I for Cas1–Cas2). b, The presence of Cas1, Cas2 and a protospacer resulted in the conversion of the supercoiled pCRISPR into relaxed, linear and band X products. c, Neither the Cas1(H208A) active site mutant nor the complex formation-defective Cas2(A66–F7) deletion mutant support the reaction. The Cas2(E9Q) active site mutant is as active as the wild type. d, Salt- and metal-dependence of radiolabelled protospacer integration into pCRISPR. e, Same as c except using radiolabelled protospacers. The data presented in b–e are representative of at least three replicates.
alanine supported protospacer integration in vitro at levels comparable to wild-type Cas1–Cas2 (Extended Data Fig. 7c). Thus, the integration reaction likely proceeds via direct nucleophilic attack of protospacer 3′-OH ends onto the target DNA phosphodiester bonds, a mechanism previously hypothesized to occur in vivo14.

**Supercoiled DNA and CRISPR locus requirements**

Cas1 and Cas2 overexpression leads to site-selective spacer acquisition proximal to the leader end of the CRISPR locus, a result consistent with observations in native populations of CRISPR-containing bacteria12–15,35. To determine what drives such site-specific integration, we first tested various forms of the pCRISPR plasmid to determine target DNA requirements. Integration requires target DNA supercoiling, as neither relaxed nor linear pCRISPR, nor the isolated 1 kb CRISPR locus, supported integration (Fig. 3c and Extended Data Fig. 6). As a control, we tested supercoiled pUC19 DNA, the parental plasmid of pCRISPR that lacks a CRISPR locus, and were surprised to observe integration products upon incubation with Cas1 and Cas2 in the presence of protospacer DNA (Fig. 3c and Extended Data Fig. 6). This finding raised two possibilities: either in vitro spacer integration is non-specific with respect to target DNA sequence or structures and/or sequence(s) favouring integration are present in the pUC19 plasmid. To determine if integration preferentially occurred at the CRISPR locus of pCRISPR, products of radiolabelled reactions were double-digested to separate the CRISPR locus (960 bp) from the pUC19 plasmid backbone (~2.27 kb). Suggestive of CRISPR-specific integration, the 32P-radiolabel migrated solely with the CRISPR locus fragment (Fig. 3d). The same result was observed when the experiment was conducted using a target plasmid containing the CRISPR locus and a different backbone sequence (pACYC) (Fig. 3e).

**CRISPR repeats provide specificity**

To determine the exact sites of protospacer integration in these reactions, we performed high-throughput sequencing of reaction products

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Figure 2 | Half-site, full-site integration and pCRISPR topoisomer products. a, Schematic of half-site and full-site integration products.
b, Linearization of the integration products (lane 4). Lane 3 is the untreated reaction products.
c, Linearization of integration products from radiolabelled protospacer reactions. d, The time course reveals the initial formation of relaxed products, followed by band X. The inset reveals the products detected using 32P-labelled protospacers. e, f, Analysis of gel-purified relaxed and band X on agarose gels pre-stained with ethidium bromide (e) or post-stained after electrophoresis (f). g, Schematic of the disintegration reaction. h, Native polyacrylamide gel analysis of the disintegration reaction. The data presented in b–f, h are representative of at least three replicates.

Figure 3 | Integration requires 3′-OH protospacer ends and supercoiled target DNA. a, b, Integration assays using single-stranded DNAs and either –OH or –PO4 at the 3′ or 5′ ends of unlabelled (a) or radiolabelled (b) protospacers. S1 corresponds to one strand of the protospacer and S2 corresponds to the complementary strand. c, Comparison of protospacer integration into different DNA targets. d, e, Restriction enzyme digestion of pCRISPR, either in a pUC19 (d) or pACYC backbone (e), after the integration assay detects integration into the CRISPR fragment (green arrows). The data presented in a–e are representative of at least three replicates.
that resulted from using either pCRISPR or the parental pUC19 vector as the target of integration (Extended Data Fig. 8a). Of the 7,866 protospacer-pCRISPR junctions retrieved, ~71% mapped to the CRISPR locus (Fig. 4a and Extended Data Fig. 8b). Protospacer insertion occurred at the borders of each repeat, with the most preferred site at the first repeat adjacent to the leader (Fig. 4b). The minus strand of each repeat (the bottom strand in Fig. 4a, b that runs 5' to 3' towards the leader sequence) is also highly preferred, highlighting the role of CRISPR repeats in providing sequence specificity for the Cas1–Cas2 complex (Fig. 4b). Sequence alignment of the integration sites revealed strong preference for sequences resembling the CRISPR repeat on both strands of pCRISPR, further supporting the selection of CRISPR repeat borders by the Cas1–Cas2 complex (Extended Data Fig. 8d–f).

The most frequent integration site in the pUC19 control plasmid mapped to the amp resistance gene adjacent to the AT-rich promoter sequence (~8.8% of 5,524 total retrieved junctions, Fig. 4c and Extended Data Fig. 8c). An inverted repeat sequence with a propensity to form a DNA cruciform occurs 9 nt adjacent to this integration site (plus strand sequence: 5'-TTCAATTATTTGAA-3'), suggesting that potential DNA cruciform formation adjacent to AT-rich sequences is important for protospacer integration. Sequence analysis of pUC19 target sites revealed the propensity for a G nucleotide to occur at the -2 and +1 positions of the protospacer insertion site, similar to the preferred pCRISPR sites (Extended Data Fig. 8g, h). These observations imply that in addition to sequence, pCRISPR repeat selectivity stems from the unique structural features of these sites, such as their ability to form cruciforms (Fig. 4a, b, c).

In E. coli, newly acquired spacers harbour a 5'-G as the first nucleotide flanking the leader-proximal end of the repeats, which originates from the last nucleotide of the AAG protospacer-adjacent motif (PAM) from foreign DNA. Such positional specificity is critical for crRNA-guided interference, as a mutation in this position of the corresponding crRNA disrupts PAM binding and subsequent target destruction. We found that ~73% of all integration events into pCRISPR used the 3'-G end instead of the 3'-C end of protospacer DNA during integration (see Fig. 4b for protospacer sequence), and there was a strong preference for this nucleotide to attack the minus strand of the repeat sequence (Fig. 4b, d, e). A similar nucleotide bias was observed in the pUC19 target plasmid sequence data (Fig. 4d). This preference positions the G at the 5' end of the protospacer substrate as the first nucleotide of the newly integrated spacer in the CRISPR locus (Fig. 5).

When we used protospacer DNAs lacking a 3'-C or bearing 3'-C on both ends, the preference for integration into the minus strand of the CRISPR locus was significantly decreased (Extended Data Fig. 9). Thus, the Cas1–Cas2 complex plays a critical role in correctly orienting the 3'-OH end of protospacer DNA substrates for incorporation within the CRISPR locus.

**Mechanism of protospacer integration**

The results presented here explain the mechanistic basis for foreign DNA acquisition during CRISPR–Cas adaptive immunity (Fig. 5). The Cas1–Cas2 complex catalyses integration of protospacers at the leader-end of

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**Figure 4 | Protospacers are specifically integrated into the CRISPR locus.**

(a) Integration sites along pCRISPR. (b) Magnified view of the integration sites along the ~1 kb CRISPR locus. The cyan peaks represent positions where the 3'-T of the protospacer DNA was integrated whereas the black peaks represent the 3'-OH integration events. The protospacer sequence is depicted above the plot. (c) Integration sites along pUC19. (d) Comparison of C 3'-OH or T 3'-OH selection in the total reads from pCRISPR and pUC19 targets (n = 7,866 reads for pCRISPR and n = 5,524 reads for pUC19, chi-square test, *P < 0.0001). (e) Schematic of DNA cruciform formation of the repeat sequences. The orange arrows depict the cleavage sites.

**Figure 5 | Model of protospacer integration during CRISPR–Cas adaptive immunity.** The first nucleophilic attack occurs on the minus strand of the first repeat, distal to the leader, by the C 3'-OH end of the protospacer. After half-site intermediate formation, the second integration event occurs on the opposite strand at the leader-repeat border. The resulting single-stranded DNA gaps are repaired by yet uncharacterized mechanisms and the protospacer is fully integrated with the G as the first nucleotide at its 5' end. The asterisk denotes the duplication of the first repeat, as previously observed in vivo.13–15.
the CRISPR locus and also selects the terminal C 3′-OH as the attacking nuclease, resulting in the 5′ G on the opposite strand of the protospacer becoming the first nucleotide of the newly integrated spacer. This orientation bias, previously observed in vitro49, is a key step during immunity for productive downstream foreign DNA targeting by the Cascade complex and Cas3 effector nuclease (Extended Data Fig. 10). Interestingly, the presence of the complete AAG PAM in the protospacer is not required for in vitro integration, suggesting that a highly specific selection or processing step occurs in vivo to exclude the AA nucleotides from the mature protospacer before integration.

We propose a two-step integration mechanism in which the C 3′-OH first attacks the minus strand of the CRISPR repeat to produce a half-site intermediate (Fig. 5). The 3′-OH on the opposite strand of the integrating DNA then attacks the target DNA 28 bp away on the opposite side of the repeat on the plus strand, leading to full integration of the protospacer (Fig. 5). Our in vitro system predominantly traps the first step of this two-step integration mechanism, suggesting that the second nuclease attack is greatly accelerated in vivo in the presence of cellular factors. This model is consistent with spacer integration intermediates that are observed in vivo, in which protospacers are integrated such that staggered cleavage at each end of the repeat generates single-stranded gaps that ensure repeat duplication44. The in vivo conditions could also promote the high specificity of integration to occur solely downstream of the first repeat of the CRISPR locus in E. coli, instead of at every repeat, as observed in our in vitro assay.

CRISPR spacer integration shares mechanistic similarities with retroviral integration and DNA transposition, where the integrase/transposase enzyme uses donor DNA 3′-OH ends to make a staggered cut at the DNA target site, which concurrently joins the donor DNA to target DNA 5′-phosphates31,32. Completion of the integration reaction requires a DNA polymerase to fill in sequence gaps and a DNA ligase to seal the phosphodiester backbone44. Similar polymerase and ligase functions are required to complete CRISPR spacer acquisition in vivo, although the specific enzymes involved have not yet been identified. Despite these similarities, we note that the Cas1 active site does not get DNA 5′-ends to make a staggered cut and DNA transposition, where the integrase/transposase enzyme uses donor DNA 3′-OH ends to make a staggered cut at the DNA target site, which concurrently joins the donor DNA to target DNA 5′-phosphates31,32. Completion of the integration reaction requires a DNA polymerase to fill in sequence gaps and a DNA ligase to seal the phosphodiester backbone44. Similar polymerase and ligase functions are required to complete CRISPR spacer acquisition in vivo, although the specific enzymes involved have not yet been identified. Despite these similarities, we note that the Cas1 active site does not get DNA 5′-ends.

Our results highlight the fundamental role of repeat sequences at multiple stages of CRISPR–Cas adaptive immunity. In addition to creating structures within nascent CRISPR transcripts that ensure correct RNA processing during crRNA maturation44, the repeats operate at the DNA level to recruit the Cas1–Cas2 complex for sequence- and structure-specific protospacer integration. We envision that this recruitment involves transient DNA cruciform formation within the CRISPR inverted repeats that occurs as a function of target DNA supercoiling44. The observation that a preferred non-CRISPR site of Cas3 CRISPR (band X) in vitro, although the physiological significance of band X production remains unclear.

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Author Contributions J.K.N. performed the biochemical experiments. A.S.Y.L. processed and analysed the high-throughput sequencing data. J.K.N., A.S.Y.L., A.E. and J.A.D. designed the study, analysed the data and wrote the manuscript.

Author Information Sequencing data are deposited in Gene Expression Omnibus under accession number GSE64552. Reprints and permissions information is available at www.nature.com/reprints. The authors declare competing financial interests: details are available in the online version of the paper. Readers are welcome to comment on the online version of the paper. Correspondence and requests for materials should be addressed to J.A.D. (doudna@berkeley.edu).
METHODS

Cas1, Cas2 and DNA preparation. The cas1 and cas2 genes from E. coli K12 (MG1655) were cloned into expression vectors and the proteins were separately purified as previously described. The proteins were stored in 100 mM KCl, 20 mM HEPES-NaOH, 5% glycerol and 1 mM TCEP at -80 °C before use. Single-stranded DNAs were synthesized (Integrated DNA Technologies). Double-stranded DNA protospacers were annealed in 20 mM HEPES-NaOH, pH 7.5, 25 mM KCl, 10 mM MgCl₂ or MnCl₂, 1 mM DTT, 10% DMSO by heating at 95 °C for 3 min and slow cooling to room temperature. The sequence of the 33 bp protospacer used in this study was shown to be the most acquired in vivo in E. coli K12 after M13 bacteriophage infection: strand 1 (5'-GCCCAATTCTACATCTGGTTCTGTTCTCTCGT-3') and strand 2 (5'-ACGAGAAACACCCAGAAGCAGTATAATTGCGGC-3'). The pCRISPR target plasmid was constructed by annealing the CRISPR fragment into a pACYC CRISPR plasmid. In vitro integration assays. The integration reactions were performed in 20 mM HEPES-NaOH, pH 7.5, 25 mM KCl, 10 mM MgCl₂ or MnCl₂, 1 mM DTT and 10% DMSO. There was little difference when DMSO was omitted from the reaction (Fig. 1d), in contrast to its in vitro integration efficiency with HIV-1 integrase. All of the reactions were conducted with MgCl₂ unless otherwise noted. For reactions with the Cas1–Cas2 complex, separately purified Cas1 and Cas2 were pre-incubated for 20–30 min at 4 °C to allow complex formation. The protospacer DNAs were incubated with the protein(s) for 10–15 min at 4 °C, followed by the addition of the target pCRISPR or pUC19 plasmid DNA. The reactions were conducted at 37 °C for 1 h and quenched with DNA loading buffer containing a final concentration of 50 mM EDTA. The products were analysed on 1.5% agarose gel electrophoresis, and the gel was stained with ethidium bromide. All of the reactions, except those shown in Fig. 1 and Extended Data Fig. 1a, were conducted with 75 nM protein, 200 nM protospacers and 7.5 nM pCRISPR to clearly visualize band X from pCRISPR. Reactions in Fig. 1 and Extended Data Fig. 1a, c, e were conducted with 50 nM protospacers. Each integration and disintegration assay was performed a minimum of three times. Radiolabelled protospacer integration assays. Pre-annealed double-stranded protospacer DNA substrates were 5'-radiolabelled using [γ-³²P]-ATP (PerkinElmer) and T4 polynucleotide kinase (New England Biolabs). Protospacers with 3'-PO₄ ends were 5'-radiolabelled using T4 polynucleotide kinase with 3' phosphatase minus activity (New England Biolabs). The reactions were carried out in the same buffer as above. Unless otherwise noted, 200 nM of Cas1–Cas2 was first incubated with 200 nM protospacers at 4 °C for 10–15 min, followed by the addition of 200 ng (~5 nM) of pCRISPR. The reactions were conducted at 37 °C for 1 h and quenched with 25 mM EDTA and 0.4% SDS. The DNA samples were deproteinized with 30 μg of proteinase K for 1 h at 37 °C and ethanol precipitated. The reactions were analysed on 1.5% agarose gels. After electrophoresis, the gels were dried onto positively charged nylon transfer membrane (GE Healthcare) and imaged using Phosphor Screens (GE Healthcare). The restriction enzyme digest experiments were performed by first conducting the integration reaction, followed by addition of the respective enzymes (New England Biolabs), which were allowed to digest for an additional 1 h at 37 °C.

Disintegration assays. The four single-stranded DNA substrates were annealed to form the Y DNA in a stepwise manner: 95 °C for 3 min, 65 °C for 20 min, 50 °C for 20 min, and gradual cooling to room temperature. The annealing reactions were analysed on 15% native polyacrylamide gel to confirm the formation of the Y DNA (Extended Data Fig. 5b). The disintegration assay was performed in the integration reaction buffer with 50 nM protein and 5 nM Y DNA at 37 °C for 1 h. For native polyacrylamide gel analysis, the reaction was quenched with DNA loading buffer with 50 mM EDTA and analysed on 15% polyacrylamide gels. For denaturing polyacrylamide gel analysis, the reactions were quenched with formamide buffer and heated at 95 °C before loading on 15% 8M urea-polyacrylamide gels. The sequences of the four strands are as follows: A (5'-GGCCCCAGTGTCGGAATGAT3'); B (5'-GTGACGCTGTTCTGGTGTTTCTCGTACCGC-3'); C (5'-GCCCAATTCTACATCTGGTTCTGTTCTCTCGTACCGC-3'); and D (5'-ACGAGAAACACCCAGAAGCAGTATAATTGCGGC-3').

High-throughput sequencing. The integration reaction was performed with 75 nM Cas1–Cas2, 200 nM protospacer and 7.5 nM pCRISPR or pUC19 in 20 mM HEPES, pH 7.5, 25 mM KCl, 10 mM MgCl₂, 10% DMSO and 1 mM DTT. The DNA samples were isolated by phenol–chloroform extraction and ethanol precipitation. The excess protospacers were removed using 100K MWCO Amicon Ultra-0.5 ml centrifugal filters. The resulting integration products were digested into smaller DNA fragments using dsDNA Fragmentase (New England Biolabs) for 75 min at 37 °C and quenched at 65 °C for 15 min. Fragments were end repaired using T4 DNA Polymerase (NEB), Klenow (NEB) and T4 PNK (NEB) and A-tailed with Klenow exonuclease (3’ to 5’ exonuclease minus) (NEB). Adapters were ligated onto fragments using T4 DNA ligase (NEB) and cDNA libraries were amplified by PCR using Phusion (NEB). Libraries were sequenced on an Illumina HiSeq2500 on rapid run mode. The oligonucleotides used were: universal adaptor: 5'-AATGTAGCCGGGGACACCGAATCTCACTCTTCCACGAGAAGCTTCCGATCTGCTCGTATGCT-3' (*phosphothioate bond); indexed adaptor: 5'-S5PhosGATGCGAAGAAGCGAGTTCTGAACTCCAGTCAC-index-ATCTCGATGCGTCCTGTCTC-3'); PCR primers: 5'-AATGTAGCCGGGGACACCGAATCTCACTCTTCCACGAGAAGCTTCCGATCTGCTCGTATGCT-3'; 5'-CAAGCAGAAAGAGGCGATACAGAT-3'.

Computational analysis. For preprocessing, 3’ adapters were removed from raw Illumina reads using Cutadapt (http://code.google.com/p/cutadapt/), discarding reads shorter than 15 nt. Reads containing integrated protospacer were selected using Cutadapt, requiring the presence of at least 10 nt of protospacer sequence with no errors. After creating Bowtie indexes from fasta files of the pUC19 empty and pCRISPR plasmid sequences, these reads were mapped to the respective plasmids using Bowtie, allowing up to 2 mismatches and requiring unique mapping. Sequence motif analysis depicted in Extended Data Fig. 8 were generated using WebLogo, using integration sites that are represented at least ten times in the sequencing data.
Extended Data Figure 1 | The integration reaction is dependent on the presence of protospacers, low salt and divalent metal ions. a, In vitro integration assay alongside EcoRI- and Nb.BbvCI nickase-treated pCRISPR. b, Salt-dependence assay using Cas1 or Cas2 only and Cas1 + Cas2. The titration corresponds to 0, 25, 50, 100 and 200 nM KCl, in addition to the salt carried in from the reaction reagents. c, Integration assays in the presence of 10 mM EDTA, Mg²⁺, Mn²⁺ or no additive. d, Integration assays with increasing protospacer concentrations. e, A comparison of post-reaction treatments as indicated. The data presented in a–e are representative of at least three replicates.
Extended Data Figure 2 | Cas1 requires Cas2 for robust protospacer integration. a, Schematic of the integration assays using 32P-labelled protospacers (PDB code 4P6I for Cas1–Cas2). b, Integration assays in the presence of increasing protein and 10 mM MnCl₂. The titration corresponds to 0, 50, 100 and 200 nM protein. c, Same as b except in the presence of 10 mM MgCl₂. The data presented in b and c are representative of at least three replicates.
Extended Data Figure 3 | The catalytic activity of Cas1 is required for integration. **a**, Close-up view of the Cas1 active site with the conserved residues shown in stick configurations (PDB 4P6I). **b**, Integration assays of purified Cas1 active site mutants complexed with wild-type Cas2. **c**, The same as **b** except using radiolabelled protospacers. The data presented in **b** and **c** are representative of at least three replicates.
Extended Data Figure 4 | Band X corresponds to topoisomers of pCRISPR.  

a, Agarose gel of purified relaxed and band X integration products. b, Analysis of the total reaction products, after phenol chloroform extraction and ethanol precipitation, on a pre-stained agarose gel. c, Same as b except ethidium bromide staining was performed after electrophoresis. d, PCR amplification products of various segments of pCRISPR using the relaxed, band X or pCRISPR template shown in a. The laddering effect of minor products using CRISPR locus primers likely reflects the propensity of CRISPR repeats to form DNA hairpins. The data presented in a–d are representative of at least three replicates.
Extended Data Figure 5 | Cas1 catalyses the disintegration of half-site integrated protospacers. 

a, Schematic of the four strands constituting the Y DNA substrate used in the disintegration assays. 

b, Native polyacrylamide gel analysis of the annealing products with either strand A or strand C radiolabelled.

c, Native polyacrylamide gel analysis of disintegration assay products using Y DNA substrates with strand A labelled.

d, Denaturing gel analysis of the disintegration assay products with strand A labelled.
Extended Data Figure 6 | Cas1–Cas2 can integrate various lengths of double-stranded DNA with blunt- or 3’-overhang ends into a supercoiled target plasmid. 

a, Integration assays using the indicated lengths of protospacer DNA. 

b, Integration assays using varying 5’ or 3’ overhang lengths. 

c, d, A comparison of integration assays using pCRISPR or Nb.BbvCI-nicked pCRISPR target. 

e, Integration assay using different target plasmids with or without a CRISPR locus. The green arrows correspond to the relaxed product of each target and the cyan arrows correspond to the band X product. 

The data presented in a–e are representative of at least three replicates.
Extended Data Figure 7 | Cas1 tyrosine mutants support integration activity in vitro. a, A close-up of the Cas1 active site with the tyrosine residues labelled in blue. b, Structure-based sequence alignment of Cas1 proteins, highlighting the tyrosine residues mutated to alanine in this study. c, Radiolabelled protospacer integration assay of Cas1 tyrosine mutants complexed with wild-type Cas2. The gel presented in c is representative of at least three replicates.
Extended Data Figure 8 | High-throughput sequencing of integration products reveals sequence-specific integration. a, Schematic of the workflow for high-throughput sequencing analysis of the integration sites. b, Raw map of the total reads along pCRISPR before collapsing into single peaks of protospacer–pCRISPR junctions depicted in Fig. 4. c, Same as b, except for the pUC19 target. d, Sequence of the leader-end of the CRISPR locus in *E. coli*. e, f, WebLogo analysis from the −5 to +5 positions surrounding the protospacer integration sites on the plus (e) and minus (f) of pCRISPR. The arrow points to the nucleotide that is covalently joined to the protospacer. g, h, Same as e, f, except for the pUC19 target.
Extended Data Figure 9 | Cas1–Cas2 correctly orients the protospacer DNA during integration. 

a–f, Mapped integration sites along the CRISPR locus of pCRISPR when using protospacer DNA with nucleotide ends 'wild-type' 3' C and 3' T (a), 3' A and 3' T (c), and 3' C and 3' C (e). The red arrow in c and e points to the nucleotide change in the protospacer DNA compared to the 'wild-type' sequence in a. The protospacer DNA 3' nucleotide and the CRISPR locus strand biases in a, c, e are plotted in b, d and f, respectively, as percentages of integration events within the CRISPR locus. The black and clear bars represent the (−) and (+) strands of the CRISPR locus, respectively. NS corresponds to not significant and *P < 0.0001 by chi-square test. The n values for b, d and f are 5,623, 5,685 and 12,453 reads along the CRISPR locus, respectively.
Extended Data Figure 10 | Model of the CRISPR–Cas adaptive immunity pathway in *E. coli*. Mature double-stranded protospacers bearing a 3′-C-OH are site-specifically integrated into the leader-end of the CRISPR locus. Correct protospacer integration (left) results in the 5′G/3′C as the first nucleotide of the spacer, proximal to the leader. After transcription of the CRISPR locus and subsequent crRNA processing, foreign DNA destruction is initiated by strand-specific recognition of the 3′-TTC-5′ PAM sequence in the target strand by the crRNA-guided Cascade complex. Incorrect protospacer integration (right) cannot initiate foreign DNA destruction due to the inability for the crRNA to recognize the strand with the 3′-TTC-5′ PAM. Thus, foreign DNA interference during CRISPR–Cas adaptive immunity relies on the Cas1–Cas2 complex for correctly orienting the protospacer during integration.