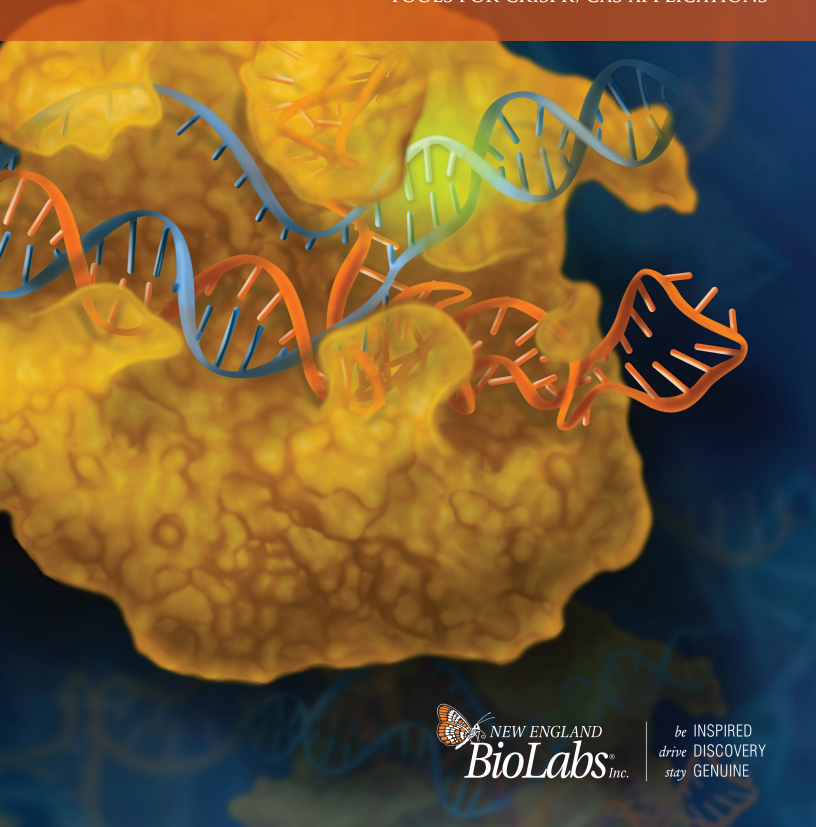


## Genome Editing

TOOLS FOR CRISPR/CAS APPLICATIONS

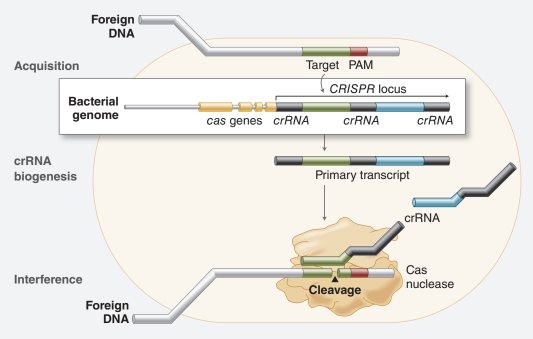


# Genome Editing: Tools for CRISPR/Cas Applications

Genome editing is enabled by the development of tools to make precise, targeted changes to the genome of living cells. Recent approaches to targeted genome modification – zinc-finger nucleases (ZFNs) and transcription-activator like effector nucleases (TALENs) – enable researchers to generate mutations by introducing double-stranded breaks to activate repair pathways. These approaches are costly and time consuming to engineer, limiting their widespread use, particularly for large scale, high-throughput studies. Recently, methods based on a bacterial CRISPR-associated protein-9 nuclease (Cas9) from *Streptococcus pyogenes* have generated considerable excitement.

CRISPR (Clustered Regularly Interspaced Short Palindromic Repeats) and CRISPR-associated (Cas) genes are essential for adaptive immunity in select bacteria and archaea, enabling the organisms to respond to and eliminate invading genetic material.

### CRISPR/Cas in vivo: Bacterial Adaptive Immunity



In the acquisition phase, foreign DNA is incorporated into the bacterial genome at the CRISPR locus. The CRISPR locus is then transcribed and processed into crRNA during crRNA biogenesis. During interference, Cas endonuclease complexed with crRNA cleaves foreign DNA containing a crRNA complementary sequence adjacent to the PAM sequence. (Figure not drawn to scale.)

### **DOWNLOAD THE NEB AR APP\***



### CRISPR/Cas Genome Editing

The simplicity of the CRISPR nuclease system (nuclease and guide RNA), makes this system attractive for laboratory use. Breaks activate repair through error prone Non-Homologous End Joining (NHEJ) or Homology Directed Repair (HDR). In the presence of a donor template with homology to the targeted locus, the HDR pathway may operate, allowing for precise mutations to be made. In the absence of a template, NHEJ is activated, resulting in insertions and/or deletions (indels), which disrupt the target locus (1,2).

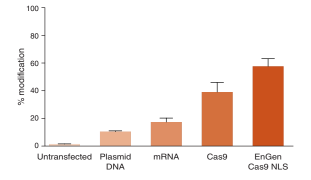
### **TOOLS & RESOURCES**

Visit www.neb.com/GenomeEditing to find our up-to-date listing of products and protocols to support this application.

### Direct Introduction of Cas RNP Complexes

The highest efficiency strategy for genome engineering with CRISPR/Cas is direct introduction of Cas9/guide RNA complexes (3–8) or Cas12a (Cpf1)/guide RNA complexes. This method further simplifies CRISPR/Cas workflows and has been reported to increase mutagenic activity (3–5) and reduce off-target editing events (3,4). NEB\* provides purified Cas9 Nuclease, *S. aureus* and *S. pyogenes* variants, and Cas12a nuclease (Cpf1), *Lachnospiraceae* bacterium ND2006 with nuclear localization signals (NEB #M0654, NEB #M0667, NEB #M0646, NEB #M0653) as standalone enzymes to support direct introduction of Cas RNP complexes.

### Increased genome editing efficiency using Cas9 RNP delivery

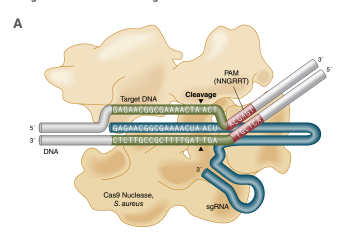


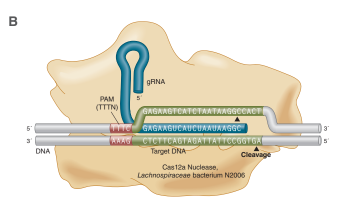
### **Ordering Information**

| PRODUCT                     | NEB #      |  |
|-----------------------------|------------|--|
| EnGen Spy Cas9 HF1          | M0667T/M   |  |
| Cas9 Nuclease, S. pyogenes  | M0386S/T/M |  |
| EnGen Spy Cas9, NLS         | M0646T/M   |  |
| EnGen Spy Cas9 Nickase      | M0650S/T   |  |
| EnGen Spy dCas9 (SNAP-tag®) | M0652S/T   |  |
| EnGen Sau Cas9              | M0654S/T   |  |
| EnGen Lba Cas12a (Cpfl)     | M0653S/T   |  |

Cas9 and sgRNA targeting a human gene were delivered to HEK293 cells by transfection. Transfected plasmid DNA contained expression cassettes for 2x NLS (N- and C-terminal) Cas9 and sgRNA. Plasmid DNA was delivered using TransIT-X2 (Mirus). Transfected mRNA was modified with pseudouridine and 5-methylcytosine and encoded 2x NLS (N- and C-terminal) Cas9. sgRNA was co-transfected with the mRNA using TransIT-mRNA. Cas9 RNPs were delivered in reverse transfections using Lipofectamine RNAiMAX (Life Technologies) using 10 nanomolar final concentration of ribonucleoprotein (RNP). Cas9 has no NLS in the protein sequence. EnGen Cas9 has N- and C-terminal NLSs. The efficiency of editing was determined using T7 Endonuclease I assay and is expressed as % modification.

### Schematic representation of Cas9 Nuclease, S. aureus (A) and Lba Cas12a, Lachnospiraceae bacterium N2006 (B) sequence recognition and DNA cleavage

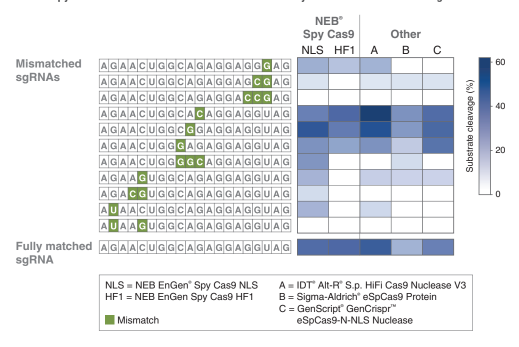




### Reducing Off-Target DNA Cleavage

EnGen Spy Cas9 HF1 is a high-fidelity, quadruple substitution (N497A/R661A/Q695A/Q926A) variant of EnGen Spy Cas9 NLS from *Streptococcus pyogenes* with reduced non-specific DNA cleavage. Spy Cas9 is an RNA-guided endonuclease that catalyzes site-specific cleavage of double stranded DNA. The single guide RNA (sgRNA) targets Cas9 to the region immediately upstream of a 5′-NGG-3′ protospacer adjacent motif (PAM) producing a double stranded break 3 bases upstream of the PAM (9). EnGen Spy Cas9 HF1 contains Simian virus 40 (SV40) T antigen nuclear localization sequence (NLS) on the N- and C-termini of the protein.

EnGen Spy Cas9 HF1 demonstrates increased sensitivity to mismatches between quide RNA and DNA targets in vitro



Comparison of the tolerance of mismatches between the guide RNA sequence and target DNA sequence of EnGen Spy Cas9 NLS, EnGen Spy Cas9 HF1, and other commercially available high fidelity Cas9 variants. One of several guide RNAs encoding a single, double, or triple mismatch with a fluorescently labeled dsDNA substrate were allowed to form a ribonucleoprotein (RNP) complex with each of five Cas9 variants. A fully matched guide RNA was included as a control. The RNPs were incubated with the substrate at a 2:1 ratio at 37°C for 5 minutes. The percent substrate cleavage for each RNP complex was measured by capillary electrophoresis. Results were graphed as a heat map with white representing no cleavage and increasing intensity of blue indicating increasing percent cleavage. The guide RNA sequence is indicated in each row, with mismatches denoted in green. The DNA protospacer sequence is 5′ – AGAACTGGCAGAGGGTAG – 3′ and the protospacer adjacent motif (PAM) is 5′ – TGG – 3′. EnGen Spy Cas9 HF1 demonstrates increased sensitivity to mismatches by showing the greatest ratio of on-target cleavage to average cleavage of off-targets.

## Rapid Generation of sgRNA for Spy Cas9

The EnGen sgRNA Synthesis Kit simplifies the generation of microgram quantities of custom sgRNAs in an hour or less by combining template synthesis and transcription. The single-tube reaction is easy to set up and requires a single ~55 nt ssDNA target-specific oligonucleotide, which is combined with the Reaction Mix and Enzyme Mix included in the kit. sgRNAs are suitable for use in downstream applications, including CRISPR/Cas9-based genome editing and *in vitro* DNA cleavage. This single-reaction format offers ease-of-use and eliminates separate DNA amplification and template clean up steps. This kit is compatible with EnGen Spy Cas9 NLS, Cas 9 Nuclease, *S. pyogenes*, EnGen Spy Cas9 Nickase, EnGen Spy dCas9 (SNAP-tag) and EnGen Spy Cas9 HF1..

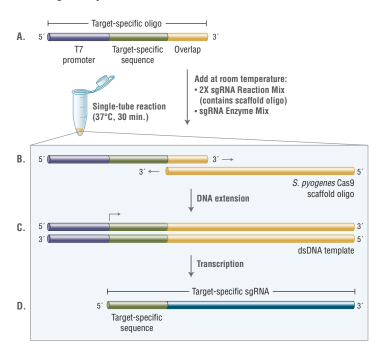
### **Ordering Information**

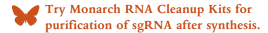
| PRODUCT                   | NEB #  |
|---------------------------|--------|
| EnGen sgRNA Synthesis Kit | E3322S |



Try our EnGen sgRNA Template
Oligo Designer (accessible through
NEBioCalculator\* at NEBiocalculator.neb.com)

### EnGen sgRNA Synthesis Kit overview





Learn more at

www.neb.com/MonarchRNACleanup.

This kit is really easy to use and will save us plenty of time in making sgRNAs! Thanks for the streamlined method!

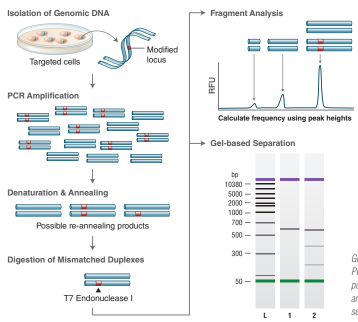
Postdoctoral Researcher,
 Harvard University

- A. The target-specific oligo contains the T7 promoter sequence, ~20 nucleotides of targetspecific sequence and a 14-nucleotide overlap sequence complementary to the S. pyogenes Cas9 Scaffold Oligo, supplied in the reaction mix. Target-specific oligos are mixed with the EnGen 2X sgRNA Reaction Mix and the EnGen sgRNA Enzyme Mix at room temperature.
- B. At 37°C the two oligos anneal at the 14-nucleotide overlap region of complementarity.
- C. The DNA polymerase contained in the EnGen sgRNA Enzyme Mix extends both oligos from their 3´ ends, creating a dsDNA template.
- D. The RNA polymerase contained in the EnGen sgRNA Enzyme Mix recognizes the dsDNA of the T7 promoter and initiates transcription, resulting in a target-specific sgRNA.
- All steps occur in a single reaction during a 30-minute incubation at 37°C.

## Evaluating Targeting Efficiency with the EnGen Mutation Detection Kit and T7 Endonuclease I

A widely used method to identify mutations is the T7 Endonuclease I mutation detection assay (10,11). This assay detects heteroduplex DNA that results from the annealing of a DNA strand, including desired mutations, with a wild-type DNA strand. The EnGen Mutation Detection Kit provides optimized reagents for performing robust T7 Endonuclease-based detection of genome editing events.

### **Workflow for EnGen Mutation Detection Kit**



### Ordering Information

| PRODUCT                                      | NEB #    |
|--|----------|
| EnGen Mutation Detection Kit                 | E3321S   |
| T7 Endonuclease I                            | M0302S/L |
| Q5® Hot Start High-Fidelity<br>2X Master Mix | M0494S/L |



Need to determine targeting efficiencies over 50%?

Visit www.neb.com/Cas9locusmod to find out how.

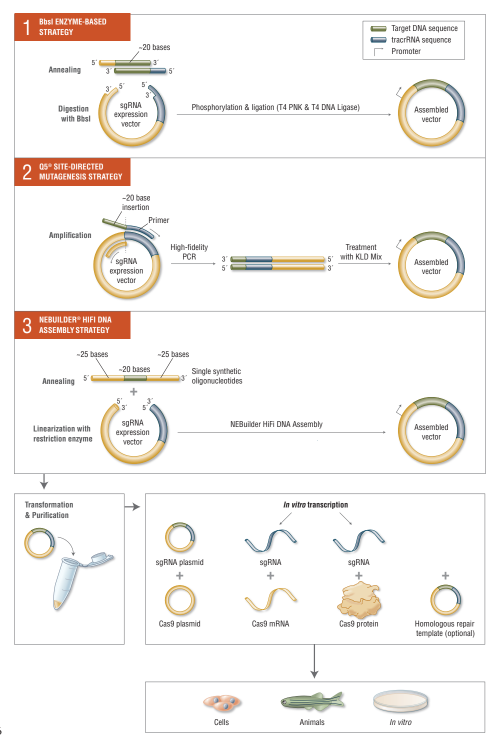


Learn more about genome editing vocabulary through the NIST Genome Editing Lexicon www.nist.gov/programs-projects/nist-genome-editing-lexicon

Genomic DNA is amplified with primers bracketing the modified locus. PCR products are then denatured and re-annealed yielding three classes of possible structures. Duplexes containing a mismatch greater than one base are digested by T7 Endonuclease I. The DNA is then electrophoretically separated and fragment analysis is used to estimate targeting efficiency.

## sgRNA Template Construction for CRISPR/Cas Genome Editing

Cas9 experiments require the introduction of guide RNAs, in addition to Cas9 nuclease. sgRNAs contain a 20-base sequence, specific to the target DNA, upstream of an invariant scaffold sequence. sgRNAs can be delivered as an RNA made *in vitro*, or by delivering an expression cassette in which the sgRNA is transcribed from an upstream promoter. For researchers using plasmid-based expression of sgRNA in target cells, or sgRNAs made *in vitro* from plasmid templates, NEB provides tools to support a number of strategies to quickly change the 20-bp targeting sequence of sgRNA templates.



### **Ordering Information**

| PRODUCT  | NEB #  |
|--|--|
| BbsI   | R0539S/L   |
| T4 DNA Ligase  | M0202S/T/L/M   |
| T4 Polynucleotide Kinase   | M0201S/L   |
| PRODUCT  | NEB #  |
| Q5 Site-Directed Mutagenesis Kit   | E0554S   |
| Q5 Site-Directed Mutagenesis Kit<br>(Without Competent Cells)  | E0552S   |
| PRODUCT  | NEB #  |
| Q5 Hot Start High-Fidelity<br>2X Master Mix  | M0494S/L   |
| NEBuilder HiFi DNA Assembly<br>Master Mix  | E2621S/L/X   |
| NEBuilder HiFi DNA Assembly<br>Cloning Kit   | E5520S   |
|  |  |
| PRODUCT  | NEB #  |
| PRODUCT  EnGen sgRNA Synthesis Kit   | NEB #<br>E3322S  |
|  |  |
| EnGen sgRNA Synthesis Kit  NEB 5-alpha Competent <i>E. coli</i>  | E3322S   |
| EnGen sgRNA Synthesis Kit  NEB 5-alpha Competent <i>E. coli</i> (High Efficiency)  NEB 10-beta Competent <i>E. coli</i>  | E3322S<br>C2987P/R/I/H/U   |
| EnGen sgRNA Synthesis Kit  NEB 5-alpha Competent <i>E. coli</i> (High Efficiency)  NEB 10-beta Competent <i>E. coli</i> (High Efficiency)  HiScribe T7 mRNA Kit with   | E3322S<br>C2987P/R/I/H/U<br>C3019I/H                                     |
| EnGen sgRNA Synthesis Kit  NEB 5-alpha Competent <i>E. coli</i> (High Efficiency)  NEB 10-beta Competent <i>E. coli</i> (High Efficiency)  HiScribe T7 mRNA Kit with CleanCap® Reagent AG  | E3322S<br>C2987P/R/I/H/U<br>C3019I/H<br>E2080S                           |
| EnGen sgRNA Synthesis Kit  NEB 5-alpha Competent <i>E. coli</i> (High Efficiency)  NEB 10-beta Competent <i>E. coli</i> (High Efficiency)  HiScribe T7 mRNA Kit with CleanCap® Reagent AG  HiScribe™ T7 ARCA mRNA Kit  HiScribe T7 ARCA mRNA Kit   | E3322S  C2987P/R/I/H/U  C3019I/H  E2080S  E2065S                         |
| EnGen sgRNA Synthesis Kit  NEB 5-alpha Competent E. coli (High Efficiency)  NEB 10-beta Competent E. coli (High Efficiency)  HiScribe T7 mRNA Kit with CleanCap® Reagent AG  HiScribe™ T7 ARCA mRNA Kit HiScribe T7 ARCA mRNA Kit (with tailing)  HiScribe T7 High Yield   | E3322S  C2987P/R/I/H/U  C3019I/H  E2080S  E2065S  E2060S                 |
| EnGen sgRNA Synthesis Kit  NEB 5-alpha Competent <i>E. coli</i> (High Efficiency)  NEB 10-beta Competent <i>E. coli</i> (High Efficiency)  HiScribe T7 mRNA Kit with CleanCap® Reagent AG  HiScribe™ T7 ARCA mRNA Kit (with tailing)  HiScribe T7 High Yield RNA Synthesis Kit  HiScribe T7 Quick High Yield                               | E3322S  C2987P/R/I/H/U  C3019I/H  E2080S  E2065S  E2060S                 |
| EnGen sgRNA Synthesis Kit  NEB 5-alpha Competent E. coli (High Efficiency)  NEB 10-beta Competent E. coli (High Efficiency)  HiScribe T7 mRNA Kit with CleanCap® Reagent AG  HiScribe™ T7 ARCA mRNA Kit (with tailing)  HiScribe T7 High Yield RNA Synthesis Kit  HiScribe T7 Quick High Yield RNA Synthesis Kit  RNA Cap Structure Analog | E3322S  C2987P/R/I/H/U  C3019I/H  E2080S  E2065S  E2060S  E2040S  E2050S |

### Featured NEB Products Supporting CRISPR Workflows

| PRODUCT NAME   | CRISPR/CAS9 APPLICATION  | NEB #         | SIZE              |
|--|--|---------------|-------------------|
| <b>NEW</b> EnGen Spy Cas9 HF1                                      | In vitro cleavage of dsDNA. Genome engineering by direct introduction of active nuclease complexes.                                | M0667T/M      | 400/2,000 pmol    |
| EnGen Spy Cas9 NLS   | In vitro cleavage of dsDNA. Genome engineering by direct introduction of active nuclease complexes.                                | M0646T/M      | 400/2,000 pmol    |
| EnGen Mutation Detection Kit                                       | Determination of the targeting efficiency of genome editing protocols  | E3321S        | 25 rxns           |
| EnGen sgRNA Synthesis Kit  | Generation of microgram quantities of custom sgRNA   | E3322S        | 20 rxns           |
| EnGen Spy Cas9 Nickase   | In vitro nicking of dsDNA. Genome engineering by direct introduction of active nuclease complexes.                                 | M0650S/T      | 70/400 pmol       |
| EnGen Spy dCas9 (SNAP-tag®)  | <i>In vitro</i> binding of DNA. Compatible with SNAP-tag substrates for visualization and enrichment.                              | M0652S/T      | 70/400 pmol       |
| EnGen Sau Cas9   | In vitro cleavage of dsDNA. Genome engineering by direct introduction of active nuclease complexes. Recognizes 5'-NNGRRT-3' PAM.   | M0654S/T      | 70/400 pmol       |
| EnGen Lba Cas12a (Cpfl)  | <i>In vitro</i> cleavage of dsDNA. Genome engineering by direct introduction of active nuclease complexes. Recognizes 5´-TTTN PAM. | M0653S/T      | 70/2,000 pmol     |
| Cas9 Nuclease, <i>S. pyogenes</i>                                  | In vitro cleavage of dsDNA. Genome engineering by direct introduction of active nuclease complexes.                                | M0386S/T/M    | 70/400/2,000 pmol |
| Q5 Site-directed Mutagenesis Kit (with or without competent cells) | Insertion of target sequence into the Cas9-sgRNA construct and modification of HDR templates                                       | E0554S/E0552S | 10 rxns           |
| Q5 High-fidelity DNA Polymerases                                   | High-fidelity construct generation for use with CRISPR workflows   | Multiple*     | Multiple*         |
| NEBuilder HiFi DNA Assembly Master Mix                             | Single-tube, isothermal generation of the Cas9-sgRNA construct and HDR templates   | E2621S/L/X    | 10/50/250 rxns    |
| NEBuilder HiFi DNA Assembly Cloning Kit                            | Single-tube, isothermal generation of the Cas9-sgRNA construct and HDR templates   | E5520S        | 10 rxns           |
| HiScribe T7 ARCA mRNA Kit (with or without tailing)                | Generation of Cas9 mRNA with ARCA cap  | E2060S/E2065S | 20 rxns           |
| HiScribe T7 High Yield RNA Synthesis Kit                           | Generation of sgRNA and Cas9 mRNA  | E2040S        | 50 rxns           |
| HiScribe T7 Quick High Yield RNA Synthesis Kit                     | Generation of sgRNA and Cas9 mRNA  | E2050S        | 50 rxns           |
| HiScribe T7 mRNA Kit with CleanCap® Reagent AG                     | Generation of Cas9 mRNA with CleanCap Reagent AG   | E2080S        | 20 rxns           |
| T7 Endonuclease I  | Determination of the targeting efficiency of genome editing protocols  | M0302S/L      | 250/1,250 units   |
| Monarch RNA Cleanup Kit  | Cleanup of sgRNA and Cas9 mRNA   | T2040S/L      | 10/100 preps      |

<sup>\*</sup> Visit Q5PCR.com for ordering information.

### VISIT NEB.COM TO FIND:



### **Protocols** for

applications such as sgRNA synthesis and direct introduction of sgRNA/Cas9 complexes

### **EnGen sgRNA Template Oligo Designer**

(accessible through **NEBioCalculator®** at NEBiocalculator.neb.com)



**NEBuilder®** for DNA Assembly (NEBuilder.neb.com)



**NEBaseChanger®** for the Q5 Site-Directed Mutagenesis Kit

(NEBaseChanger.neb.com)

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### Featured Tools



For help with configuring target-specific DNA oligos, try our EnGen sgRNA Template Oligo Designer (accessible through NEBioCalculator\* at NEBiocalculator.neb.com)



For help with designing primers for DNA assembly, try NEBuilder\* DNA Assembly Tool (NEBuilder.neb.com)



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