

Includes the latest products and tools for DNA Assembly

Molecular Cloning

TECHNICAL GUIDE

FREE Next Day Delivery (UK mainland)



Molecular Cloning Overview

Molecular cloning refers to the process by which recombinant DNA molecules are produced and transformed into a host organism, where they are replicated. A molecular cloning reaction is usually comprised of two components:

- 1. The DNA fragment of interest to be replicated.
- 2. A vector/plasmid backbone that contains all the components for replication in the host.

DNA of interest, such as a gene, regulatory element(s), operon, etc., is prepared for cloning by either excising it out of the source DNA using restriction enzymes, copying it using PCR, or assembling it from individual oligonucleotides. At the same time, a plasmid vector is prepared in a linear form using restriction enzymes (REs) or Polymerase Chain Reaction (PCR). The plasmid is a small, circular piece of DNA that is replicated within the host and exists separately from the host's chromosomal or genomic DNA. By physically joining the DNA of interest to the plasmid vector through phosphodiester bonds, the DNA of interest becomes part of the new recombinant plasmid and is replicated by the host. Plasmid vectors allow the DNA of interest to be copied easily in large amounts, and often provide the necessary control elements to be used to direct transcription and translation of the cloned DNA. As such, they have become the workhorse for many molecular methods such as protein expression, gene expression studies, and functional analysis of biomolecules.

During the cloning process, the ends of the DNA of interest and the vector have to be modified to make them compatible for joining through the action of a DNA ligase, recombinase, or an *in vivo* DNA repair mechanism. These steps typically utilize enzymes such as nucleases, phosphatases, kinases and/or ligases. Many cloning methodologies and, more recently kits have been developed to simplify and standardize these processes.

This technical guide will clarify the differences between the various cloning methods, identify NEB^{*} products available for each method, and provide expert-tested protocols and FAQs to help you troubleshoot your experiments and **Clone with Confidence**^{*}.

Visit CloneWithNEB.com



- Technical tips and FAQs
- Videos and animations
- Much more...

TABLE OF CONTENTS

- 3 Online Tools
- 4-5 Cloning Workflow Comparison

6–10 DNA Assembly

- 6 Overview
 - 6 Product Selection
 - 7 NEBridge[®] Golden Gate Assembly Kits 7 Optimization Tips
 - 8 Technical Tips for Optimizing NEBridge Golden Gate Assembly Reactions
 - 9 NEBuilder[®] HiFi DNA Assembly 10 Protocol/Optimization Tips

11–12 Cloning & Mutagenesis

11 NEB PCR Cloning Kit

- 12 Q5[®] Site-Directed Mutagenesis Kit
 - 12 Protocols/Optimization Tips
- 13–25 DNA Preparation
 - 13 Nucleic Acid Purification
 - 13 Overview
 - 14 cDNA Synthesis

14 Overview

15 Restriction Enzyme Digestion

- 15 Overview
- 15 Protocol/Optimization Tips

16–22 Performance Chart

23 Activity in rCutSmart Buffer

24 PCR/Amplification

- 24 Overview
- 24 Product Selection
- 25 Protocols/Optimization Tips

26-28 Common DNA End Modifications

26 Overview

- 26 Phosphorylation
 - 26 Protocol/Optimization Tips
- 26 Dephosphorylation
 - 26 Product Selection
 - 26 Protocol/Optimization Tips

27 Blunting/End-repair

- 27 Product Selection
- 27 Protocol/Optimization Tips
- 28 A-tailing
 - 28 Product Selection
- 28 Protocol

29–30 DNA Ligation

29 Vector and Insert Joining

- 29 Overview
- 30 Product Selection
- 30 Protocol/Optimization Tips
- 31 Transformation
 - 31 Overview
 - 31 Product Selection
 - 31 Protocol/Optimization Tips
- 32 DNA Analysis

32 DNA Markers and Ladders

32 Overview32 Product Selection

- 33-34 Traditional Cloning Quick Guide
- 35-37 Troubleshooting Guide
- 38-39 Ordering Information

Online Tools for Cloning

Competitor Cross-Reference Tool



Use this tool to select another company's competent cell product and find out which NEB strain is compatible. Choose either the product name or catalog number from the available selection, and this tool will identify the recommended NEB product and its advantages. A link to the product

page where you can also order the product is provided.

DNA Sequences and Maps Tool



With the DNA Sequences and Maps Tool, find the nucleotide sequence files for commonly used molecular biology tools, including plasmid, viral and bacteriophage vectors.

Double Digest Finder



Use this tool to guide your reaction buffer selection when setting up double-digests, a common timesaving procedure. Choosing the right buffers will help you to avoid star activity and loss of product.

Enzyme Finder



Use this tool to select restriction enzymes by name, sequence, overhang or type. Enter your sequence using single letter code, and Enzyme Finder will identify the right enzyme for the job.

NEBridge® Ligase Fidelity Tools



Visualize overhang ligation preferences, predict high-fidelity junction sets, and split DNA sequences to facilitate the design of high-fidelity Golden Gate assemblies.

NEBridge® Golden Gate Assembly Tool



Use this tool to assist with in silico DNA construct design for Golden Gate DNA assembly. It enables the accurate design of primers with appropriate Type IIS restriction sites and overlaps, quick import of sequences in many

formats and export of the final assembly, primers and settings. The latest version also incorporates ligase fidelity information.

NEBaseChanger[®]



NEBaseChanger can be used to design primers specific to the mutagenesis experiment you are performing using the Q5® Site-Directed Mutagenesis Kit. This tool will also calculate a recommended custom annealing temperature based on the sequence of the primers by taking into

account any mismatches. The latest version enables multi-site mutagenesis using NEBuilder HiFi DNA Assembly.

NEBcloner[®]



Use this tool to find the right products and protocols for each step (digestion, end modification, ligation and transformation) of your next traditional cloning experiment. Also, find other relevant tools and resources

to enable protocol optimization.

NEBcutter[®] V3.0



Identify restriction sites within your DNA sequence using NEBcutter. Choose between Type II and commercially available Type III enzymes to digest your DNA. NEBcutter V3.0 indicates cut frequency and

methylation sensitivity.

NEBioCalculator®



NEBioCalculator is a collection of calculators and converters that are useful in planning bench experiments in molecular biology laboratories.

NEBuilder[®] Assembly Tool



NEBuilder Assembly Tool can be used to design primers for your NEBuilder HiFi DNA Assembly and Gibson Assembly reaction, based on the entered fragment sequences and the polymerase being used for amplification.

NEBuilder[®] Protocol Calculator



Use the NEBuilder Protocol Calculator to calculate the optimal amounts of input DNA sequences for the NEBuilder HiFi DNA Assembly reaction given the length and concentration of each input fragment.

PCR Selector



Use this tool to help select the right DNA polymerase for your PCR setup. Whether your amplicon is long, complex, GC-rich or present in a single copy, the PCR selection tool will identify the perfect DNA polymerase

for your reaction.

REBASE[®]



Use this tool as a guide to the ever-changing landscape of restriction enzymes. REBASE, the Restriction Enzyme DataBASE, is a dynamic, curated database of restriction enzymes and related proteins.

Tm Calculator



Use this tool when designing PCR reaction protocols to help determine the optimal annealing temperature for your amplicon. Simply input your DNA polymerase, primer concentration and your primer sequence and the

Tm Calculator will guide you to successful reaction conditions.

MOBILE APPS



NEB Tools for iPhone®, iPad® or Android®

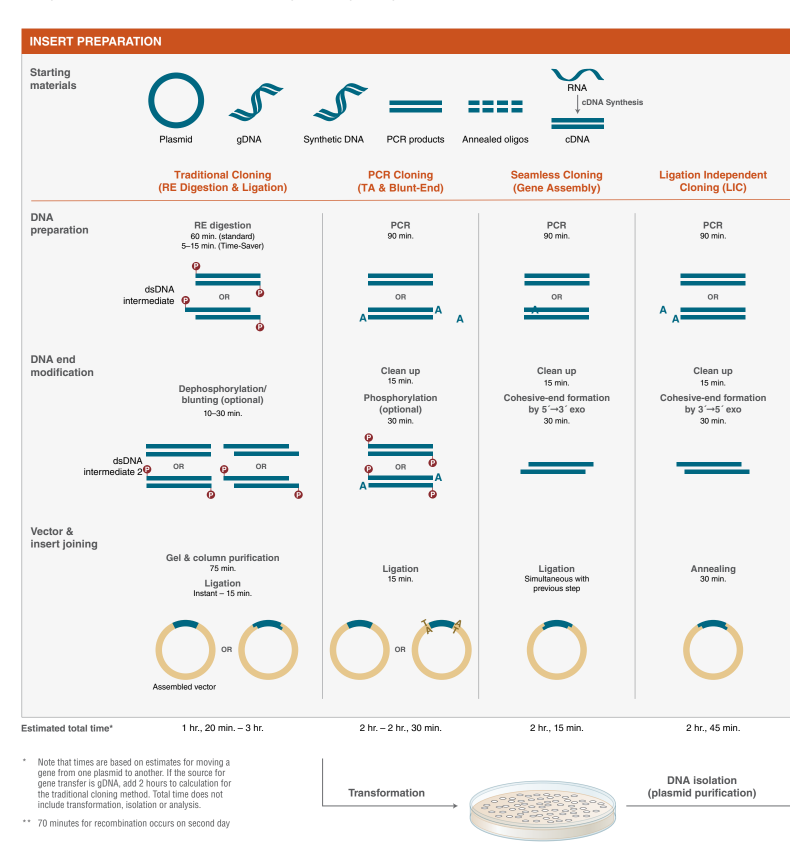
NEB Tools brings New England Biolabs' most popular web tools to your iPhone, iPad or Android devices.

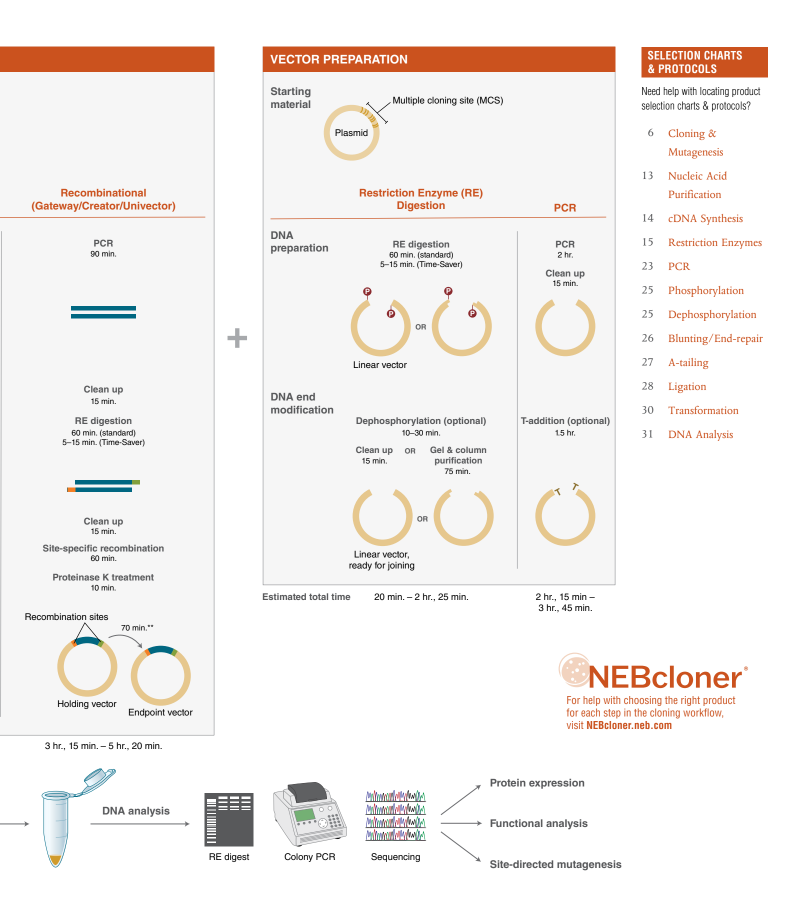
- Use Enzyme Finder to select a restriction enzyme by category or recognition sequence, or search by name to find information on any NEB enzyme. Sort your results so they make sense to you, then email them to your inbox or connect directly to www.neb.com.
- Use Double Digest Finder or NEBcloner to determine buffer and reaction conditions for experiments requiring two restriction enzymes.

When using either of these tools, look for rCutSmart[™], HF[®] and Time-Saver[™] enzymes for the ultimate in convenience. NEB Tools enables quick and easy access to the most requested restriction enzyme information, and allows you to plan your experiments from anywhere.

Cloning Workflow Comparison

The figure below compares the steps for the various cloning methodologies, along with the time needed for each step in the workflows.





DNA Assembly

DNA assembly refers to a molecular cloning method that physically links together multiple fragments of DNA, in an end-to-end fashion, to acheive a designed, higher-order assembly prior to joining to a vector. This process is the cornerstone of the synthetic biology field and allows the construction of novel biological systems and devices using defined components. These techniques are carried out *in vitro* and are typically enzymatically driven with the final constructions being maintained in microbial host cells.

To help select the best DNA assembly method for your needs, please refer to our Synthetic Biology/DNA Assembly Selection Chart below.

Synthetic Biology/DNA Assembly Selection Chart

| | NEBuilder HiFi DNA Assembly NEB #E2621 NEB #E5520 NEB #E2623 | NEB Gibson Assembly NEB #E2611 NEB #E5510 | NEBridge Golden Gate Assembly Kits (Bsal-HFv2/BsmBl-v2) NEB #E1601 NEB #E1602 NEBridge Ligase Master Mix NEB #M1100 | USER Enzyme NEB #M5505 Thermolabile USER II Enzyme NEB #M5508 |
|---|--|--|--|--|
| Properties | | | | |
| Removes 5' or 3' End Mismatches | *** | * | N/A | N/A |
| Assembles with High Fidelity at Junctions | *** | ** | *** | *** |
| Tolerates Repetitive Sequences at Ends | * | * | *** | *** |
| Generates Fully Ligated Product | *** | *** | *** | NR |
| Joins dsDNA with Single-stranded Oligo | *** | ** | NR | NR |
| Assembles Low Amounts of DNA with High Efficiency | *** | ** | ** | ** |
| Accommodates Flexible Overlap Lengths | *** | *** | * | ** |
| Applications | | | | |
| 2 Fragment Assembly (Simple cloning) | *** | *** | *** | *** |
| 3-6 Fragment Assembly (one pot) | *** | *** | *** | *** |
| 7-11 Fragment Assembly (one pot) | *** | ** | *** | *** |
| 12-50+ Fragment Assembly (one pot) (1) | * | * | *** | NR |
| Template Construction for In vitro Transcription | *** | *** | *** | *** |
| Synthetic Whole Genome Assembly | *** | * | *** | * |
| Multiple Site-directed Mutagenesis | *** | ** | ** | ** |
| Library Generation | *** | *** | *** | ** |
| Metabolic Pathway Engineering | *** | ** | *** | *** |
| TALENs | ** | ** | *** | ** |
| Short Hairpin RNA (shRNA) Cloning | *** | ** | * | * |
| gRNA Library Generation | *** | ** | * | * |
| Large Fragment (> 10 kb) Assembly | *** | *** | *** | ** |
| Small Fragment (< 100 bp) Assembly | *** | * | *** | *** |
| Use in Successive Rounds of Restriction Enzyme Assembly | *** | * | NR | * |

⁽¹⁾ Please visit **neb.com/GoldenGate** for more information

| KEY | |
|-----|---|
| *** | Optimal, recommended product for selected application |
| ** | Works well for selected application |
| * | Will perform selected application, but is not recommended |
| N/A | Not applicable to this application |
| NR | Not recommended |

Golden Gate Assembly

The efficient and seamless assembly of DNA fragments, commonly referred to as Golden Gate assembly (1,2) refers to multiple inserts being assembled into a vector backbone using only the sequential (3) or simultaneous (4) activities of a single Type IIS restriction enzyme and T4 DNA Ligase. Golden Gate has enabled single inserts, the cloning of inserts from diverse populations enabling library creation, and multi-module assemblies. NEB has made extraordinary improvements that touch every application of the Golden Gate technology.

Advances in Ligase Fidelity

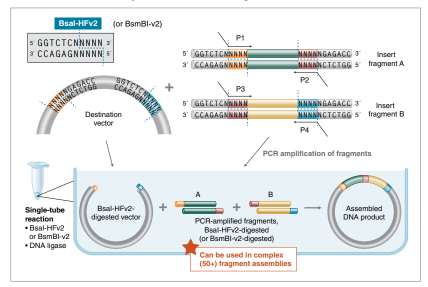
Research at NEB has led to increased understanding of ligase fidelity, including the development of a comprehensive method for profiling end-joining ligation fidelity to predict which overhangs have improved fidelity (5). This research led to Data-optimized Assembly Design (DAD) which allows careful choice of overhang sets and is especially important for achieving complex and high-fidelity Golden Gate Assemblies.

The NEBridge Ligase Fidelity tools utilize DAD to enable the design of new assemblies, including complex high fragment number assemblies, and the evaluation, modification, or expansion of existing assemblies, and more.

Type IIS Restriction Enzymes for Golden Gate Assembly

NEB offers more Type IIS (i.e., recognize asymmetric DNA sequences and cleave outside of their recognition sequence) restriction enzymes than any other supplier, many of which are used in Golden Gate Assembly. These enzymes, along with the ligase fidelity data, allows complex 50+ fragment assemblies with high efficiency, > 90% accuracy and low backgrounds.

Golden Gate Assembly Workflow for complex assemblies



In its simplest form, Golden Gate Assembly requires a Type IIS recognition site, in this case, Bsal-HFv2 (GGTCTC), or BsmBI-v2 (CGTCTC) added to both ends of a dsDNA fragment. After digestion, these sites are left behind, with each fragment bearing the designed 4-base overhangs that direct the assembly.

RECOMMENDED PRODUCTS

NEBridge Golden Gate Assembly Kits (Bsal-HFv2 or BsmBI-v2) (NEB #E1601, NEB #E1602)

- · Seamless cloning no scar remains following assembly
- · Includes destination plasmid with T7/SP6 promoters
- Ordered assembly of multiple fragments (2-50+) in a single reaction
- · Can also be used for cloning of single inserts and library preparations
- Efficient with regions with high GC content and areas of repeats
- · Compatible with a broad range of fragment sizes (< 100 bp to > 15 kb)

NEBridge Ligase Master Mix (NEB #M1100)

- · Optimized for efficient and accurate Golden Gate Assembly
- Convenient 3X Master Mix format
- · Use with NEB Type IIS restriction enzymes

Type IIS Enzymes used in Golden Gate

- Bbsl (NEB #R0539)
- BbsI-HF® (NEB #R3539)
- BspQI-HF[®] (NEB #R3712)
- BtgZI (NEB #R0703)
- Bsal-HF®v2 (NEB #R3733)
- Esp3I (NEB #R0734) - PaqCI® (NEB #R0745)
- BsmBI-v2 (NEB #R0739) - BspQI (NEB #R0712)
 - Sapl (NEB #R0569)

TOOLS & RESOURCES

Visit www.neb.com/GoldenGate to find:

- · Publications and protocols related to ligase fidelity and Golden Gate Assembly
- Access to NEBridge Golden Gate Assembly Tool, for help with designing your experiments. Also available at GoldenGate.neb.com
- Access to the NEBridge Ligase Fidelity Tools to facilitate the design of high-fidelity Golden Gate Assemblies. Also available at ligasefidelity.neb.com
- · View our webinar: Listen to DAD when constructing high-complexity Golden Gate Assembly targets
- View our MoClo Overhang Standards Usage Guidelines and our tutorial, Domestication and Golden Gate Assembly

- 1. Engler, C. et al. (2008) PLoS ONE, 3: e3647.
- 2. Engler, C. et al. (2009) PLoS ONE, 4: e5553.
- 3. Lee, J.H. et al. (1996) Genetic Analysis: Biomolecular Engineering, 13; 139-145.
- 4. Padgett, K.A. and Sorge, J.A. (1996) Gene, 168, 31-35. 5. Potapov, V. et. al. (2018) ACS Synth. Biol. DOI: 10.1021/acssynbio.8b00333.

Technical Tips for Optimizing Golden Gate Assembly Reactions

Looking to assemble multiple DNA fragments in a single reaction? Here are some tips to keep in mind when planning your Golden Gate Assembly experiment.

1. Check your sequences

Always check your assembly sequences for internal sites before choosing which Type IIS restriction endonuclease to use for your assembly. While single insert Golden Gate assembly has such high efficiencies of assembly that the desired product is obtainable regardless of the presence of an internal site, this is not true for assemblies with multiple inserts. Options include choosing a different Type IIS restriction enzyme to direct your assembly, or eliminating internal sites through domestication. Our tutorial video on Golden Gate Assembly Domestication provides a full description of the many options available for internal site issues. Note the use of a Type IIS restriction enzyme with a 7 base recognition site, such as PaqCI, is less likely to have internal sites present in any given sequence.

2. Orient your primers

When designing PCR primers to introduce Type IIS restriction enzyme sites, either for amplicon insert assembly or as an intermediate for pre-cloning the insert, remember that the recognition sites should always face inwards towards your DNA to be assembled. Consult the Golden Gate Assembly Kit manuals or videos for further information regarding the placement and orientation of the sites.

3. Choose the right plasmid

Consider switching to the pGGAselect Destination Plasmid for your Golden Gate assembly. This versatile new destination construct is included in all Golden Gate Assembly kits and can be used for BsaI-HFv2, BsmBI-v2 or BbsI directed assemblies. It also features T7 and SP6 promoter sequences flanking the assembly site, and has no internal BsaI, BsmBI or BbsI sites. The pGGAselect plasmid can also be transformed into any *E. coli* strain compatible with pUC19 for producing your own plasmid preparation if so desired.

4. Choose the right buffer

T4 DNA Ligase Buffer works best for Golden Gate Assembly with BsaI-HFv2, BsmBI-v2 and PaqCI. However, alternate buffers would be NEBuffer r1.1 for Bsa-HFv2, NEBuffer r2.1 for BsmBI-v2, or rCutSmart[™] Buffer for PaqCI, if these buffers are supplemented with 1 mM ATP and 5–10 mM DTT. NEB also offers NEBridge Ligase Master Mix that has been optimized for Golden Gate Assembly with our Type IIS restriction enzymes for Golden Gate.

5. Increase your complex assembly efficiency by increasing the Golden Gate cycling levels

T4 DNA Ligase, BsaI-HFv2, BsmBI-v2 and PaqCI are very stable and continue to be active during extended cycling protocols; an easy way to increase assembly efficiencies without sacrificing fidelity is to increase the total cycles from 30 to 45–65, even when using long (5-minute) segments for the temperature steps.

6. Make sure your plasmid prep is RNA-free

For pre-cloned inserts/modules, make sure your plasmid prep is free of RNA to avoid an overestimation of your plasmid concentrations.

7. Avoid primer dimers

For amplicon inserts/modules, make sure your PCR amplicon is a specific product and contains no primer dimers. Primer dimers, with Type IIS restriction endonuclease sites (introduced in the primers used for the PCR reactions), would be active in the assembly reaction and result in mis-assemblies.

8. Avoid PCR-induced errors

Do not over-cycle and use a proof reading high fidelity DNA polymerase, such as $Q5^{\ensuremath{\mathbb{R}}}$ DNA High-Fidelity Polymerase.

9. Decrease insert amount for complex assemblies

For complex assemblies involving >10 fragments, pre-cloned insert/ modules levels can be decreased from 75 to 50 ng each without significantly decreasing the efficiencies of assembly.

10. Carefully design EVERY insert's overhang

An assembly is only as good as its weakest junction. Research at NEB has led to an increased understanding of ligase fidelity, including the development of a comprehensive method for profiling end-joining ligation fidelity in order to predict which overhangs will result in improved accuracy. This ligase fidelity information can be paired with the appropriate Type IIS restriction enzyme chosen to direct your assembly to achieve high efficiencies and accurate complex assemblies. Please use the free NEBridge Golden Gate Assembly Tool to design primers for your Golden Gate Assembly reactions. Predict overhang fidelity or find optimal Golden Gate junctions for long sequences, refer to the NEBridge Ligase Fidelity Tools.

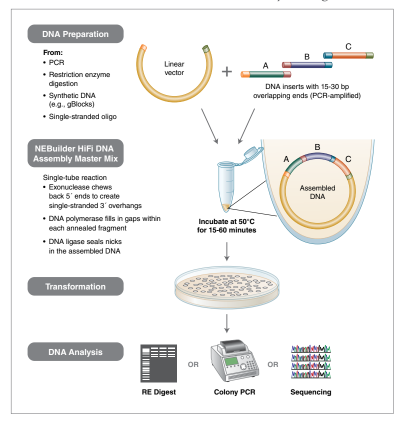
11. If using pre-cloned proven inserts that suddenly become problematic, check for a possible mutational event in your sequence.

Be aware that occasionally a pre-cloned insert/module can become corrupted by an error during propagation in *E. coli*, usually a frameshift due to slippage in a run of a single base (e.g., AAAA) by the *E. coli* DNA Polymerase. This should be suspected if previously functional assembly components suddenly become nonfunctional.

NEBuilder HiFi DNA Assembly

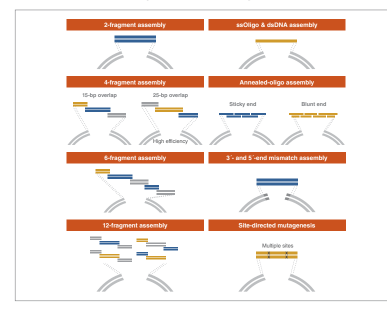
NEBuilder HiFi DNA Assembly enables virtually error-free joining of DNA fragments, even those with 5'- and 3'-end mismatches. Available with and without competent *E. coli*, this flexible kit enables simple and fast seamless cloning utilizing a new proprietary high-fidelity polymerase. Make NEBuilder HiFi DNA Assembly your first choice for DNA assembly and cloning.

Overview of the NEBuilder HiFi DNA Assembly cloning method



NEBuilder HiFi DNA Assembly

can be used for a variety of DNA assembly methods.



RECOMMENDED PRODUCTS

NEBuilder HiFi DNA Assembly Cloning Kit (NEB #E5520)

NEBuilder HiFi DNA Assembly Master Mix (NEB #E2621)

NEBuilder HiFi DNA Assembly Bundle for Large Fragments (NEB #E2623)

- · Simple and fast seamless cloning in as little as 15 minutes
- Use one system for both "standard-size" cloning and larger gene assembly products (up to 12 fragments and 20* kb)
- DNA can be used immediately for transformation or as template for PCR or RCA
- Adapts easily for multiple DNA manipulations, including site-directed mutagenesis
- Enjoy less screening/re-sequencing of constructs, with virtually error-free, high-fidelity assembly
- Use NEBuilder HiFi DNA Assembly in successive rounds of assembly, as it removes 5⁻ and 3⁻ end mismatches
- Bridge two ds-fragments with a synthetic ssDNA oligo for simple and fast construction (e.g., linker insertion or gRNA library)
- No PCR cleanup step required
- No licensing fee requirements from NEB for NEBuilder products
- NEBuilder HiFi DNA Assembly Cloning Kit includes the NEBuilder HiFi DNA Assembly Master Mix and NEB 5-alpha Competent *E. coli*
- NEBuilder HiFi DNA Assembly Bundle for Large Fragments includes the NEBuilder HiFi DNA Assembly Master Mix and NEB 10-beta Competent *E. coli* for assemblies larger than 15 kb.

TOOLS & RESOURCES

Visit NEBuilderHiFi.com to find:

- Online tutorials to help with assembly and primer design
- Application notes utilizing NEBuilder HiFi DNA Assembly
- Access to NEBuilder Assembly Tool, our online primer design tool
- Access to **NEBuilder Protocol Calculator**, our custom protocol generator
- Comparison to In-Fusion[®] Snap Assembly and GeneArt[®] Gibson Assembly[®]

Optimization Tips for NEBuilder HiFi DNA Assembly

Assembly Reaction

- PCR product purification is not necessary if the total volume of all PCR products is 20% or less of the assembly reaction volume. Higher volumes of unpurified PCR products may reduce the efficiency, so column purification of PCR products is highly recommended when performing assemblies of three or more PCR fragments or assembling longer fragments > 5 kb.
- Carefully follow guidelines as indicated in the protocol regarding total amount of DNA and ratios of insert:vector.
- Use the NEBuilder Protocol Calculator at **nebuildercalculator.neb.com** to help calculate the optimal amounts of input DNA sequences.

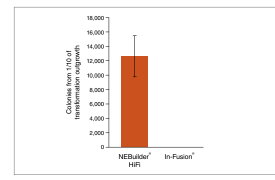
Primer Design

- For help with primer design, we recommend using NEBuilder Assembly Tool at nebuilder.neb.com.
- Vary overlap regions anywhere between 15–30 bp depending on the number of fragments being assembled.

Transformation

• The NEBuilder HiFi DNA Assembly Cloning Kit (NEB #E5520) includes NEB 5-alpha Competent *E. coli*. NEB recommends using the competent cells provided with the kit (NEB #C2987) because of their high efficiency. The components of the master mix may inhibit the functionality of competent cells from other companies if not diluted. The NEBuilder HiFi DNA Assembly Bundle for Large Fragments (NEB #E2623) includes NEB 10-beta Competent *E. coli* (NEB #C3019), ideal for assembling larger fragments (>15 kb).

NEBuilder HiFi DNA Assembly Cloning Kit can assemble a ssDNA oligo with a linearized vector.



One pmol of a ssDNA oligo was assembled with a linearized vector (20 ng, CRISPR Nuclease OFP Reporter) by incubation at 50° C for 15 min. 2 µl of the assembled mix was transformed into to NEB 10-beta Competent E. coli (NEB #C3019). 9 out of 10 colonies selected show correct sequence, while no successful assembled constructs are found using In-Fusion HD.

Gibson Assembly and the Gibson Assembly Cloning Kit

Gibson Assembly enables multiple, overlapping DNA fragments to be joined in a singletube isothermal reaction, with no additional sequence added (scar-less). The Gibson Assembly Master Mix includes three different enzymatic activities that perform in a single buffer. The assembled, fully-sealed construct is then transformed into NEB 5-alpha competent *E. coli*. The entire protocol, from assembly to transformation, takes just under two hours. Visit **NEBGibson.com** to learn more.

Protocol: Assembly

Before use, thaw and vortex the master mix thoroughly and keep on ice.

1. Set up the following reaction on ice.

| | RECOMMENDED AMOUNT OF FRAGMENTS USED FOR ASSEMBLY | | |
|---|--|----------------------------|-------------------------------------|
| | 2–3 Fragment Assembly* | 4–6 Fragment Assembly** | NEBuilder Positive Control*** |
| Recommended DNA Molar Ratio | vector:insert= 1:2 | vector:insert= 1:1 | |
| Total Amount of Fragments | 0.03-0.2 pmol* X µl | 0.2-0.5 pmol** X μl | 10 µl |
| NEBuilder HiFi DNA Assembly Master Mix | 10 µl | 10 µl | 10 µl |
| Deionized H ₂ O | 10–X µl | 10–X µl | 0 |
| Total Volume | 20 µl**** | 20 µl**** | 20 µl |

- Optimized cloning efficiency is 50–100 ng of vector with 2-fold molar excess of each insert. Use 5-fold molar excess of any insert(s) less than 200 bp. Total volume of unpurified PCR tragments in the assembly reaction should not exceed 20%. To achieve optimal assembly efficiency, design 15-20 bp overlap regions between each fragment.
- ** To achieve optimal assembly efficiency, design 20-30 bp overlap regions between each fragment with equimolarity of all fragments (suggested: 0.05 pmol each).
- *** Control reagents are provided for 5 experiments.
- **** If greater numbers of fragments are assembled, increase the volume of the reaction, and use additional NEBuilder HiFi DNA Assembly Master Mix.
- Incubate samples in a thermocycler at 50°C for 15 minutes when 2 or 3 fragments are being assembled or 60 minutes (when 4–6 fragments are being assembled). Following incubation, store samples on ice or at –20°C for subsequent transformation.

Note: Extended incubation up to 60 minutes may help to improve assembly efficiency in some cases (for further details see FAQ section of product pages).

Protocol: Transformation with NEB 5-alpha cells

| | STANDARD PROTOCOL | |
|-------------------|--|--|
| DNA | 2 µl | |
| Competent E. coli | 50 μl | |
| Incubation | On ice for 30 minutes | |
| Heat Shock | Exactly 42°C for exactly 30 seconds | |
| Incubation | On ice for 5 minutes Add 950 µl room temperature SOC 37°C for 60 minutes, with shaking | |

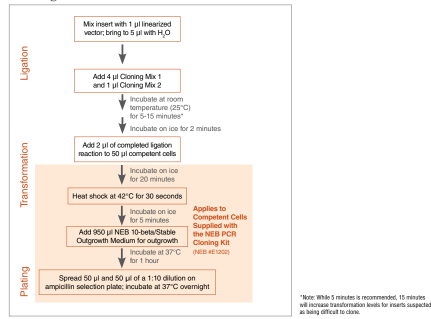


Cloning & Mutagenesis Kits

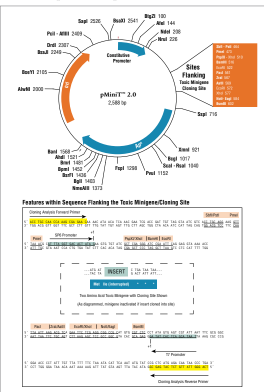
NEB PCR Cloning Kit

The NEB PCR Cloning Kit enables quick and simple cloning of all your PCR amplicons, regardless of the polymerase used. This kit utilizes a novel mechanism for background colony suppression - a toxic minigene is generated when the vector closes upon itself - and allows for direct cloning from your reaction, with no purification step.

Cloning Kit Protocol Overview



Vector Maps and Sequence



Map shown displays the construct formed if no insert is present. Unique restriction sites are shown in bold. Additional restriction sites that can be used for subcloning are also shown. Expanded box below shows location of sequencing primers, restriction sites for subcloning or linearization for in vitro transcription, RNA Polymerase promoter sequences and placement of insertion site within the toxic minigene.

Note: While 5 minutes is recommended, 15 minutes

RECOMMENDED PRODUCTS

NEB PCR Cloning Kit (NEB #E1202)

NEB PCR Cloning Kit

(without competent cells) (NEB #E1203)

- Easy cloning of all PCR products, including blunt and TA ends
- · Fast cloning experiments with 5-minute ligation step
- Simplified screening with low/no colony background and no blue/white selection required
- · Save time by eliminating purification steps
- Updated to allow for in vitro transcription with both SP6 and T7 promoters
- · Flanking restriction sites available for easy subcloning, including choice of two single digest options
- · Provided analysis primers allow for downstream colony PCR screening or sequencing
- · Ready-to-use kit components include 1 kb control amplicon, linearized cloning vector and optional single-use competent E. coli
- · Longer shelf life (12 months), as compared to some commercially available products
- Value pricing

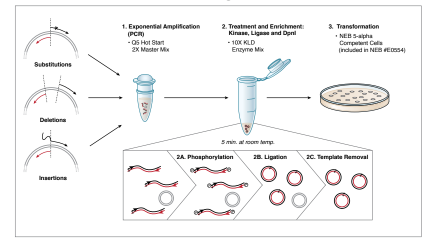
TIPS FOR OPTIMIZATION

- For first time use of the kit, prepare a positive control reaction containing 2 µl (30 ng) of the 1 kb amplicon cloning control included with the kit
- 3:1 insert:vector ratio is best, but ratios from 1:1 to 10:1 can also be utilized
- · Plate 50 µl or less of the 1 ml outgrowth. Plating too much of the outgrowth can increase background, and cause problems with colony PCR.
- Important to stop ligations: If you wish to store your ligations to allow transformations at a later time, make sure your freezer is cold enough (-20°C) to freeze the ligations.
- Do not incubate the transformation plates at room temperature. The slow growth rate of the cells at room temperature will increase the number of background colonies.
- · Follow the protocol, including the transformation protocol, carefully
- · Add the cloning mixes 1 and 2 (which can be mixed together for the day's experiment) to the reaction last

Q5 Site-Directed Mutagenesis Kit

The Q5 Site-Directed Mutagenesis Kit (with or without competent cells) enables rapid, site-specific mutagenesis of double-stranded plasmid DNA in less than 2 hours. The kit utilizes Q5 Hot Start High-Fidelity DNA Polymerase, along with custom mutagenic primers to create substitutions, deletions and insertions in a wide variety of plasmids. Transformation into high-efficiency NEB 5-alpha Competent *E. coli* cells ensures robust results with plasmids up to, at least, 14 kb in length.

Overview of Q5 Site-Directed Mutagenesis Kit



Protocol: Assembly

Before use, thaw and vortex the master mix thoroughly and keep on ice.

1. Exponential Amplification

| | 25 µl RXN | FINAL CONC. |
|---|-----------|-------------|
| Q5 Hot Start High-Fidelity 2X Master Mix | 12.5 µl | 1X |
| 10 µM Forward Primer | 1.25 µl | 0.5 µM |
| 10 µM Reverse Primer | 1.25 µl | 0.5 µM |
| Template DNA (1–25 ng/µl) | 1 µl | 1–25 ng |
| Nuclease-free water | 9.0 µl | |

2. KLD Reaction

| | VOLUME | FINAL CONC. |
|------------------------|--------|-------------|
| PCR Product | 1 µl | |
| 2X KLD Reaction Buffer | 5 µl | 1X |
| 10X KLD Enzyme Mix | 1 µl | 1X |
| Nuclease-free Water | 3 µl | |

Protocol: Transformation with NEB 5-alpha

| | STANDARD PROTOCOL |
|-------------------|--|
| KLD Mix | 5 µl |
| Competent E. coli | 50 µl |
| Incubation | On ice for 30 minutes |
| Heat Shock | Exactly 42°C for exactly 30 seconds |
| Incubation | On ice for 5 minutes. Add 950 μI room temperature SOC 37°C for 60 minutes, with shaking |

RECOMMENDED PRODUCTS

Q5 Site-Directed Mutagenesis Kit (NEB #E0554)

Q5 Site-Directed Mutagenesis Kit (Without Competent Cells) (NEB #E0552)

KLD Enzyme Mix (NEB #M0554)

- Generation of mutations, insertions or deletions in plasmid DNA
- Non-overlapping primer design ensures robust, exponential amplification and generates a high % of desired mutations from a wide range of templates
- Intramolecular ligation and transformation into NEB 5-alpha results in high colony yield
- Low error rate of Q5 High-Fidelity DNA Polymerase reduces screening time
- Use of standard primers eliminates need for phosphorylated or purified oligos

TIPS FOR OPTIMIZATION

- For optimal results, use NEBaseChanger at NEBaseChanger.neb.com to help design the primers for your SDM experiment
- No purification of your plasmid is necessary, either before or after the KLD reaction
- You can expect a high frequency of your desired mutation (> 90%)
- While the Q5 SDM Kit is supplied with high-efficiency, NEB competent *E. coli*, you can use your own chemically competent cells for cloning; results will vary, according to the quality and efficiency of the cells
- KLD Enzyme Mix (NEB #M0554) is available separately for customization
- For multi-site or combinatorial mutagenesis, we recommend NEBuilder[®] HiFi DNA Assembly products



Nucleic Acid Purification

The need for high quality, highly pure DNA and RNA is important for many molecular cloning workflows. These nucleic acids are being purified from a wide variety of sources, such as cells and tissues, enzymatic reactions (e.g., PCR, ligation, digestions), and agarose gel matrices. Purification methods are optimized for various starting materials to ensure excellent recovery, high purity, minimal processing time and compatibility with emerging techniques. The Monarch Nucleic Acid Purification product portfolio addresses the needs upstream and downstream of molecular cloning workflows, including products for isolation of DNA and RNA from biological samples, DNA and RNA cleanup, plasmid purification and gel extraction.

Monarch[®] DNA & RNA Purification Kits Designed with sustainability in mind

| _ | _ | |
|--|--|---|
| PRODUCT | APPLICATIONS | FEATURES |
| Monarch Spin Plasmid Miniprep Kit (NEB #T1110) | Purification of up to 20 µg of plasmid DNA from bacterial culture. | Elute in as little as 30 µl Prevent buffer retention and salt carryover with optimized column design Includes colored buffers to monitor completion of certain steps Reduced hands on time with faster protocols and less spin time |
| Monarch Spin PCR & DNA Cleanup Kit (NEB #T1130) | Purification and concentration of up to 5 μ g of DNA from enzymatic reactions. | Elute in as little as 5 μl Prevent buffer retention and salt carryover with optimized column design Purify oligos and other small DNA fragments with simple protocol modification |
| Monarch Spin DNA Gel Extraction Kit (NEB #T1120) | Purification of up to 5 µg of DNA from agarose gels. | Elute in as little as 5 µl Prevent buffer retention and salt carryover with optimized column design Fast, user-friendly protocol |
| Monarch Spin RNA Isolation Kit (Mini) (NEB #T2110) | Extraction and purification of up to 100 µg of total RNA from cells, tissues, bacteria, yeast, plants, insects, and other sample types. | Capture total RNA pool, including small RNAs <200 nt Proteinase K, and Monarch StabiLyse™ DNA/RNA Buffer for sample preservation Elute in as little as 6 μl Remove genomic DNA contamination with specialized columns and RNA-free DNase |
| Monarch Spin RNA Cleanup Kits (NEB #T2030, T2040, T2050) | Purification and concentration of RNA after: • Extraction (e.g., TRIzol®) • <i>in vitro</i> transcription • Enzymatic reactions • Gel electrophoresis | Highly pure RNA in as little as 5 minutes Available in 3 different binding capacities (10 µg, 50 µg and 500 µg) for flexibility in various applications Simple, user-friendly protocol Prevent buffer retention and salt carryover with optimized column design |
| Monarch Spin gDNA Extraction Kit (NEB #T3010) | Extraction and purification of genomic DNA from cells, blood, tissues and other sample types. | Optimized protocols and buffer chemistry for excellent yields from a wide variety of samples Purifies gDNA with a peak size > 50 kb Includes RNase A and Proteinase K Protocol also available for gDNA cleanup |
| Monarch HMW DNA Extraction Kit for Cells & Blood (NEB #T3050) and Tissue (NEB #T3060) | Extraction of high molecular weight (HMW) genomic DNA from a variety of samples including cells, blood, tissues, bacteria, yeast, insect, and more for long-read sequencing technologies, such as Oxford Nanopore Technologies® and PacBio®. | Isolate high molecular weight gDNA up to megabase range for long-read sequencing Extraction process taking only 30-60 minutes (cells & blood workflow) or 90 minutes (tissue workflow) compared to hours, days Compatible with a variety of sample types Utilizes unique, innovative glass bead technology and includes RNase A and Proteinase K and accessories for effective RNA removal and protein digestion |
| Monarch Mag Viral DNA/RNA Extraction Kit (NEB #T4010) | Viral nucleic acid purification from saliva, respiratory swab samples, wastewater, and more for qPCR, RT-qPCR, ddPCR, and NGS applications. | Utilizes magnetic beads for efficient and reliable viral DNA/RNA extraction Compatible with both high-throughput and manual workflows, and instrument platforms such as KingFisher® Flex, Agilent® Bravo®, and more Enables sensitive detection with included carrier RNA to enhance recovery of low amounts of viral nucleic acid |

TIPS FOR OPTIMIZATION

DNA PURIFICATION

- Ensure that the tip of the column doesn't contact with flow-through after washing: If in doubt, add a quick spin
- If working with DNA > 10 kb, heat the elution buffer to 50°C: Large DNA binds more tightly; heating helps to elute the DNA from the column

PLASMID MINIPREPS

- Don't use too many cells (culture should not exceed 15 0.D. units): Using the optimal amount of cells increases lysis efficiency and prevents clogging of the column
- Lyse cells completely: In order to release all plasmid DNA, all cells need to be lysed. Resuspend cells completely, and incubate for the recommended time.
- Don't vortex cells after lysis: Vortexing can cause shearing of host chromosomal DNA, resulting in gDNA contamination
- Allow the RNase to do its job: To prevent RNA contamination, do not skip or reduce the incubation with RNase (which is included in the neutralization buffer)
- Don't skip any washes: Proper washes ensure efficient removal of cell debris, endotoxins and salts

GEL EXTRACTION

- Use the smallest possible agarose plug: More agarose requires longer melting time and more dissolving buffer (introducing more salts which can co-elute with your sample
- Minimize exposure to UV light: UV exposure damages DNA. As long as the excision is done quickly, damage will be negligible.
- Melt the agarose completely: If the agarose is not completely melted, DNA remains trapped inside and cannot be extracted properly

GENOMIC DNA EXTRACTION

- Do not reload the same column: Over-exposure of the matrix to the lysed sample can dislodge the membrane
- Do not exceed recommended input amounts: Buffer volumes are optimized for recommended inputs. Exceeding these can result in inefficient lysis and can clog the column.
- Ensure samples are properly lysed: Samples should be disrupted and homogenized completely to release all DNA

RNA EXTRACTION & PURIFICATION

- Inactivate RNases after harvest: Nucleases in your sample will lead to degradation, so inactivating them is essential. Process samples quickly, or use preservation reagents, and always ensure nuclease-free working environments.
- Do not exceed recommended input amounts: Buffer volumes are optimized for recommended inputs. Exceeding these can result in inefficient lysis and can clog the column.
- Ensure samples are properly homogenized/ disrupted: Samples should be disrupted and homogenized completely to release all RNA

cDNA Synthesis

When RNA is used as starting material, a reverse transcriptase can be used to generate cDNA, which can then be used as template for any of the cloning methods listed previously. Depending on which workflow is being followed, the resulting DNA may require a clean-up step. This can be performed using a spin column or by gel extraction.

cDNA Synthesis Selection Chart

| 7 | | |
|---|---|--|
| cDNA SYNTHESIS | FEATURES | |
| KITS | | |
| LunaScript® RT SuperMix Kit (NEB #E3010/M3010) | Ideal for cDNA synthesis of shorter fragments Single tube supermix contains random hexamer and oligo-dT primers, dNTPs, Murine RNase Inhibitor, and Luna Reverse Transcriptase Visible blue tracking dye for easy reaction setup Fast 13-minute protocol | |
| LunaScript® RT Master Mix Kit (Primer-free) (NEB #E3025) | Ideal for first strand cDNA synthesis Compatible with random primers, oligo dT primers, and gene-specific primers for maximum cDNA synthesis flexibility 5X master mix contains dNTPs, Murine RNase Inhibitor and Luna Reverse Transcriptase Visible blue tracking dye for easy reaction setup Fast 13-minute protocol | |
| ProtoScript [®] II First Strand cDNA Synthesis Kit (NEB #E6560) | Generates cDNA at least 10 kb in length Contains ProtoScript II Reverse Transcriptase, an enzyme with increased thermostability and reduced RNase H activity Convenient 2-tube kit includes dNTPs, Oligo-dT primer and Random Primer Mix | |
| ProtoScript First Strand cDNA Synthesis Kit (NEB #E6300) | Generates cDNA at least 5 kb in length Contains M-MuLV Reverse Transcriptase Convenient 2-tube kit includes dNTPs, Oligo-dT primer and Random Primer Mix | |
| Template Switching RT Enzyme Mix (NEB #M0466) | Incorporates a universal adaptor sequence at the 3^{-'} end of cDNA during the RT reaction Enzyme mix and buffer are optimized for efficient template switching RT enzyme mix includes RNase Inhibitor High sensitivity for cDNA amplification – enables transcriptome analysis by RNA-seq from single cells or as low as 2 pg of human total RNA Robust and simple workflow for 5^{-'} Rapid Amplification of cDNA Ends (RACE) Retains the complete 5^{-'} end of transcripts for 2nd Strand cDNA Synthesis | |
| STANDALONE REAGENTS | | |
| ProtoScript II Reverse Transcriptase (NEB #M0368) An alternative to SuperScript® II | RNase H⁻ mutant of M-MuLV Reverse Transcriptase with increased thermostability and reduced RNase H activity Increased reaction temperatures (37–50°C) | |
| M-MuLV Reverse Transcriptase (NEB #M0253) | Robust reverse transcriptase for a variety of templates Standard reaction temperatures (37–45°C) | |
| Induro [®] Reverse Transcriptase (NEB #M0681) | Fast, processive, intron-encoded RT Ideal for long transcripts, direct RNA sequencing, cDNA synthesis from samples with strong secondary structure and/or with inhibitors | |
| WarmStart RTx Reverse Transcriptase (NEB #M0380) | Permits room temperature reaction setup Increased reaction temperatures (50–65°C) Optimized for RT-LAMP isothermal detection | |

TIPS FOR OPTIMIZATION

STARTING MATERIAL

- Intact RNA of high purity is essential for generating cDNA for cloning applications.
- Total RNA or mRNA can be used in the reverse transcription reaction. Total RNA is generally sufficient for cDNA synthesis reactions. However, if desired, mRNA can be easily obtained using a Magnetic mRNA Isolation Kit (NEB #S1550).
- The amount of RNA required for cDNA cloning depends on the abundance of the transcript-of-interest. In general, 1 ng to 1 µg total RNA or 0.1–100 ng mRNA is recommended.

PRODUCT SELECTION

 Streamline your reaction setup by using the ProtoScript II First Strand cDNA Synthesis Kit (NEB #E6560). This kit combines ProtoScript II Reverse Transcriptase (NEB #M0360), a thermostable M-MuLV (RNase H⁻) Reverse Transcriptase, and recombinant RNase Inhibitor in an enzyme Master Mix, along with a separate Reaction Mix containing dNTPs. Additionally, the kit contains two optimized reverse transcription primer mixes.

YIELD

- ProtoScript II Reverse Transcriptase is capable of generating cDNA of more than 10 kb up to 48°C. We recommend 42°C for routine reverse transcription.
- You can increase the yield of a long cDNA product by doubling the amount of enzyme and dNTPs.

ADDITIVES

 For most RT-PCR reactions, RNase H treatment is not required. But for some difficult amplicons or sensitive assays, add 2 units of *E. coli* RNase H to the reaction and incubate at 37°C for 20 minutes.

Protocol: cDNA Synthesis

| | DENATURATION PROTOCOL | |
|-------------------------------|--|--|
| Total RNA | 1—6 µI (up to 1 µg) | |
| d(T) ₂₃ VN (50 µM) | 2 µl | |
| Nuclease-free Water | to a total volume of 8 µl | |
| Incubation | 65°C for 5 minutes spin briefly and put on ice | |

| | SYNTHESIS PROTOCOL |
|---------------|-----------------------------------|
| Denatured RNA | 8 µl |
| Reaction Mix | 10 µl |
| Enzyme Mix | 2 µl |
| Incubation | 80°C for 5 minutes store at –20°C |

Restriction Enzyme Digestion

Restriction enzyme sites that are unique to both the insert and vector should be chosen. Unidirectional cloning is achieved using two different restriction enzymes, each with unique recognition sites at an end of the insert. Depending on the RE chosen, ends can be blunt or sticky (cohesive). Restriction enzyme digestion is generally used in traditional cloning.

Protocol: Restriction Enzyme Reactions

| | STANDARD PROTOCOL | TIME-SAVER® PROTOCOL |
|------------------------|-------------------|----------------------|
| DNA | up to 1 µg | up to 1 µg |
| 10X NEBuffer | 5 µl (1X) | 5 µl (1X) |
| Restriction Enzyme | 10 units* | 1 µl |
| Total Volume | 50 µl | 50 µl |
| Incubation Temperature | enzyme dependent | enzyme dependent |
| Incubation Time | 60 minutes | 5–15 minutes** |

*Sufficient to digest all types of DNAs.

**Time-Saver qualified enzymes can also be incubated overnight with no star activity.

TIPS FOR OPTIMIZATION

ENZYME

- Keep on ice when not in the freezer
- · Should be the last component added to reaction
- Mix components by pipetting the reaction mixture up and down, or by "flicking" the reaction tube. Follow with a quick ("touch") spin-down in a microcentrifuge. Do not vortex the reaction.
- In general, we recommend 5 10 units of enzyme per μg DNA, and 10 – 20 units per μg of genomic DNA in a 1 hour digest
- Some restriction enzymes require more than one recognition site to cleave efficiently. These are designated with the "multi-site" icon 2*site. Please review recommendations on working with these enzymes at www.neb.com.

STAR ACTIVITY

- Unwanted cleavage that can occur when an enzyme is used under sub-optimal conditions, such as:
- Too much enzyme present
- Too long of an incubation time
- Using a non-recommended buffer
- Glycerol concentrations above 5%
- Star activity can be reduced by using a High-Fidelity (HF) enzyme, reducing the number of units, reducing incubation time, using a Time-Saver enzyme or increasing reaction volume

DNA

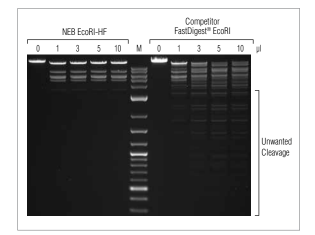
- Should be free of contaminants such as phenol, chloroform, alcohol, EDTA, detergents and salts. Spin column purification readily accomplishes this; extra washes during purification can also help.
- Methylation of DNA can affect digestion with certain enzymes.
 For more information about methylation visit www.neb.com/methylation.

BUFFER

- Use at a 1X concentration
- Recombinant Albumin is included in NEBuffer r1.1, r2.1, r3.1 and rCutSmart[™] Buffer. No additional albumin is needed.

REACTION VOLUME

- A 50 µl reaction volume is recommended for digestion of up to 1 µg of substrate. This helps maintain salt levels introduced by miniprepped DNA low enough that they don't affect enzyme activity.
- Enzyme volume should not exceed 10% of the total reaction volume to prevent star activity due to excess glycerol
- Additives in the restriction enzyme storage buffer (e.g., glycerol, salt), as well as contaminants found in the substrate solution (e.g., salt, EDTA, or alcohol), can be problematic in smaller reaction volumes



EcoRI-HF (NEB #R3101) shows no star activity in overnight digests, even when used at higher concentrations. 50 µl reactions were set up using 1 µg of Lambda DNA, the indicated amount of enzyme and the recommended reaction buffer. Reactions were incubated overnight at 37°C. Marker M is the 1 kb DNA Ladder (NEB# N3232).

| | RESTRICTION ENZYME* | DNA | 10X Nebuffer |
|------------------------|------------------------|--------|-----------------|
| 10 µl rxn […] | 1 unit | 0.1 µg | 1 µl |
| 25 µl rxn | 5 units | 0.5 µg | 2.5 µl |
| 50 µl rxn | 10 units | 1 µg | 5 µl |

- Restriction Enzymes can be diluted using the recommended diluent buffer when smaller amounts are needed
- ** 10 µl rxns should not be incubated for longer than 1 hour to avoid evaporation

INCUBATION TIME

- Incubation time for the Standard Protocol is 1 hour. Incubation for the Time-Saver Protocol is 5–15 minutes.
- Visit www.neb.com/timesaver for list of Time-Saver qualified enzymes
- It is possible, with many enzymes, to use fewer units and digest for up to 16 hours. For more information, visit www.neb.com.

STORAGE AND STABILITY

- Storage at -20°C is recommended for most restriction enzymes. For a few enzymes, storage at -80°C is recommended. Visit www.neb.com for storage information.
- 10X NEBuffers should be stored at -20°C
- The expiration date is found on the label
- Long term exposure to temperatures above -20°C should be minimized whenever possible

Performance Chart for Restriction Enzymes

New England Biolabs supplies > 210 restriction enzymes that are 100% active in rCutSmart Buffer. This results in increased efficiency, flexibility and ease-of-use, especially when performing double digests.

This performance chart summarizes the activity information of NEB restriction enzymes. To help select the best conditions for double digests, this chart shows the optimal (supplied) NEBuffer and approximate activity in the four standard NEBuffers for each enzyme. Note that Recombinant Albumin is included in all NEBuffers, and is not provided as a separate tube. In addition, this performance chart shows recommended reaction temperature, heat-inactivation temperature, recommended diluent buffer, methylation sensitivity, whether the enzyme is Time-Saver qualified, and whether the enzyme works better in a substrate with multiple sites.

Chart Legend

| U | Supplied with a unique reaction buffer that is different from the four standard NEBuffers. The compatibility with the four standard NEBuffers is indicated in the chart. | 2*site | Indicates that the restriction enzyme requires two or more sites for cleavage |
|-------------|--|--------|---|
| R }} | Recombinant | dcm | dcm methylation sensitivity |
| e | Engineered enzyme for maximum performance | dam | dam methylation sensitivity |
| 0 | Time-Saver qualified | CpG | CpG methylation sensitivity |

Activity Notes (see last column)

FOR STAR ACTIVITY

- 1. Star activity may result from extended digestion, high enzyme concentration or a glycerol concentration of > 5%
- 2. Star activity may result from extended digestion.
- 3. Star activity may result from a glycerol concentration of > 5%
- * May exhibit star activity in this buffer.
- + For added flexibility, NEB offers an isoschizomer or HF enzyme, supplied with rCutSmart Buffer.

FOR LIGATION AND RECUTTING

| а | Ligation | is less | than | 10% |
|----|----------|---------|--------|------|
| u. | Ligution | 10 1000 | tituti | 10/0 |

- b. Ligation is 25% 75%
- c. Recutting after ligation is < 5%
- d. Recutting after ligation is 50% 75%
- e. Ligation and recutting after ligation is not applicable since the enzyme is either a nicking enzyme, is affected by methylation, or the recognition sequence contains variable sequences.

| | | | Supplied | C | % Activity i | n NEBuffe | rs | Incub. Temp. | Inactiv. Temp. | | | Moth | vlation | |
|----|----------|---------|-----------|------|--------------|-----------|-----------|-----------------|-------------------|------|--------------------------------|------|----------|---------|
| | | Enzyme | NEBuffer | r1.1 | r2.1 | r3.1 | rCutSmart | (°C) | (°C) | Dil. | Unit Substrate | | sitivity | Notes |
| R% | 0 | AatII | rCutSmart | <10 | 50* | 50 | 100 | 37° | 80° | В | λ DNA | | CpG | |
| R% | 0 | Acc65I | r3.1 | 10 | 75* | 100 | 25 | 37° | 65° | А | pBC4 DNA | dcm | CpG | |
| R% | 0 | Accl | rCutSmart | 50 | 50 | 10 | 100 | 37° | 80° | А | λ DNA | | CpG | |
| R% | 0 | Acil | rCutSmart | <10 | 25 | 100 | 100 | 37° | 65° | А | λ DNA | | CpG | |
| R% | 0 | AcII | rCutSmart | <10 | <10 | <10 | 100 | 37° | No | В | λ DNA | | CpG | |
| R% | 0 | Acul | rCutSmart | 50 | 100 | 50 | 100 | 37° | 65° | В | λ DNA | | | 1, b, d |
| R% | | Afel | rCutSmart | 25 | 100 | 25 | 100 | 37° | 65° | В | pXba DNA | | CpG | |
| RX | 0 | AfIII | rCutSmart | 50 | 100 | 10 | 100 | 37° | 65° | А | ΦX174 RF I DNA | | | |
| RX | 0 | AfIIII | r3.1 | 10 | 50 | 100 | 50 | 37° | 80° | В | λ DNA | | | |
| RX | e | Agel-HF | rCutSmart | 100 | 50 | 10 | 100 | 37° | 65° | А | λDNA | | CpG | |
| R% | 0 | Ahdl | rCutSmart | 25 | 25 | 10 | 100 | 37° | 65° | А | λDNA | | CpG | а |
| R% | е | Alel-v2 | rCutSmart | <10 | <10 | <10 | 100 | 37° | 65° | В | λDNA | | CpG | |
| R% | 0 | Alul | rCutSmart | 25 | 100 | 50 | 100 | 37° | 80° | В | λDNA | | | b |
| RX | | Alwl | rCutSmart | 50 | 50 | 10 | 100 | 37° | No | А | λ DNA (dam-) | dam | | 1, b, d |
| R% | 0 | AlwNI | rCutSmart | 10 | 100 | 50 | 100 | 37° | 80° | А | λDNA | dcm | | |
| RX | 0 | Apal | rCutSmart | 25 | 25 | <10 | 100 | 37° | 65° | А | pXba DNA | dcm | CpG | |
| R% | 0 | ApaLI | rCutSmart | 100 | 100 | 10 | 100 | 37° | No | А | λ DNA (HindIII digest) | | CpG | |
| RX | 0 | ApeKI | r3.1 | 25 | 50 | 100 | 10 | 75° | No | В | λDNA | | CpG | |
| R% | e | Apol-HF | rCutSmart | 10 | 100 | 10 | 100 | 37° | 80° | В | λDNA | | | |
| RX | 0 | Ascl | rCutSmart | <10 | 10 | 10 | 100 | 37° | 80° | А | λDNA | | CpG | |
| R% | 0 | Asel | r3.1 | <10 | 50* | 100 | 10 | 37° | 65° | В | λDNA | | | 3 |
| RX | | AsiSI | rCutSmart | 100 | 100 | 25 | 100 | 37° | 80° | В | Xhol digested pXba | | CpG | 2, b |
| R% | 0 | Aval | rCutSmart | <10 | 100 | 25 | 100 | 37° | 80° | А | λDNA | | CpG | |
| R% | 0 | Avall | rCutSmart | 50 | 75 | 10 | 100 | 37° | 80° | А | λDNA | dcm | CpG | |
| R% | 0 | Avrll | rCutSmart | 100 | 50 | 50 | 100 | 37° | No | В | λ DNA (HindIII digest) | | | |
| R% | 0 | BaeGI | r3.1 | 75 | 75 | 100 | 25 | 37° | 80° | А | λDNA | | | |

a. Ligation is less than 10% b. Ligation is 25% – 75%

c. Recutting after ligation is < 5% d. Recutting after ligation is 50% - 75% e. Ligation and recutting after ligation is not applicable since the enzyme is either a nicking enzyme, is affected by methylation, or the recognition sequence contains variable sequences

| | | | | | | c | % Activity i | n NEBuffe | rs | Incub. | Inactiv. | | | | | |
|----------------|---|---|--------|-----------|----------------------|------|--------------|-----------|-----------|---------------|---------------|------|--------------------------------|-----|---------------------|---------|
| | | | | Enzyme | Supplied NEBuffer | r1.1 | r2.1 | r3.1 | rCutSmart | Temp. (°C) | Temp. (°C) | Dil. | Unit Substrate | | ylation sitivity | Notes |
| R% | 0 | | | Bael | rCutSmart | 50 | 100 | 50 | 100 | 37° | 65° | A | λDNA | | CpG | е |
| R% | 6 | | | BamHI | r3.1 | 75* | 100* | 100 | 100* | 37° | No | А | λDNA | | | 3 |
| R% | | е | | BamHI-HF | rCutSmart | 100 | 50 | 10 | 100 | 37° | No | А | λDNA | | | |
| R }} | | | | Banl | rCutSmart | 10 | 25 | <10 | 100 | 37° | 65° | А | λ DNA | dcm | CpG | 1 |
| R }} | | | | Banll | rCutSmart | 100 | 100 | 50 | 100 | 37° | 80° | А | λ DNA | | | 2 |
| RN | 6 | | | Bbsl | r2.1 | 100 | 100 | 25 | 75 | 37° | 65° | В | λ DNA | | | |
| RX | 0 | е | | BbsI-HF | rCutSmart | 10 | 10 | 10 | 100 | 37° | 65° | В | λ DNA | | | |
| Rn | | | | BbvCl | rCutSmart | 10 | 100 | 50 | 100 | 37° | No | В | λ DNA | | CpG | 1, a |
| Rn | 0 | | 2+site | Bbvl | rCutSmart | 100 | 100 | 25 | 100 | 37° | 65° | В | pBR322 DNA | | | 3 |
| R¦¦ | | | | Bccl | rCutSmart + DTT | 100 | 50 | 10 | 100 | 37° | 65° | А | pXba DNA | | | 3, b |
| R¦¦ | | | | BceAl | r3.1 | 100* | 100* | 100 | 100* | 37° | 65° | А | pBR322 DNA | | CpG | 1 |
| R {{ | | | 2+site | Bcgl | r3.1 | 10 | 75* | 100 | 50* | 37° | 65° | А | λ DNA | dam | CpG | е |
| R {{ | 0 | | | BciVI | rCutSmart | 100 | 25 | <10 | 100 | 37° | 80° | С | λ DNA | | | b |
| R % | 0 | | | Bcll | r3.1 | 50 | 100 | 100 | 75 | 37° | No | А | λ DNA (dam-) | dam | | |
| Rii | 0 | е | | Bcll-HF | rCutSmart | 100 | 100 | 10 | 100 | 37° | 65° | В | λ DNA (dam-) | dam | | |
| Rii | 0 | | | BcoDI | rCutSmart | 50 | 75 | 75 | 100 | 37° | No | В | λ DNA | | CpG | |
| Rii | | | | Bfal | rCutSmart | <10 | 10 | <10 | 100 | 37° | 80° | В | λ DNA | | | 2, b |
| Rii | 0 | | 2*site | BfuAl | r3.1 | <10 | 25 | 100 | 10 | 50° | 65° | В | λ DNA | | CpG | 3 |
| RN | 6 | | | Bgll | r3.1 | 10 | 25 | 100 | 10 | 37° | 65° | В | λ DNA | | CpG | |
| R\} | 6 | | | BgIII | r3.1 | 10 | 10 | 100 | <10 | 37° | No | А | λ DNA | | | |
| RN | 6 | | | Blpl | rCutSmart | 50 | 100 | 10 | 100 | 37° | No | А | λ DNA | | | d |
| RN | 6 | | | BmgBl | r3.1 | <10 | 10 | 100 | 10 | 37° | 65° | В | λ DNA | | CpG | 3, b, c |
| R\} | | | | Bmrl | r2.1 | 75 | 100 | 75 | 100* | 37° | 65° | В | λ DNA (HindIII digest) | | | b |
| R\} | 6 | e | | BmtI-HF | rCutSmart | 50 | 100 | 10 | 100 | 37° | 65° | В | pXba DNA | | | |
| RN | | | 2+site | Bpml | r3.1 | 75 | 100 | 100 | 100* | 37° | 65° | В | λ DNA | | | 2 |
| R\} | | | | Bpu10I | r3.1 | 10 | 25 | 100 | 25 | 37° | 80° | В | λ DNA | | | 3, b, c |
| RN | 6 | | | BpuEl | rCutSmart | 50* | 100 | 50* | 100 | 37° | 65° | В | λ DNA | | | d |
| Rn | 6 | | | BsaAl | rCutSmart | 100 | 100 | 100 | 100 | 37° | No | С | λ DNA | | CpG | |
| RX | | | | BsaBl | rCutSmart | 50 | 100 | 75 | 100 | 60° | 80° | В | λ DNA (dam-) | dam | CpG | 2 |
| Rii | 6 | | | BsaHI | rCutSmart | 50 | 100 | 100 | 100 | 37° | 80° | С | λ DNA | dcm | CpG | |
| Rii | 6 | е | | Bsal-HFv2 | rCutSmart | 100 | 100 | 100 | 100 | 37° | 80° | В | pXba DNA | dcm | CpG | |
| Rii | | | | BsaJI | rCutSmart | 50 | 100 | 100 | 100 | 60° | 80° | А | λ DNA | | | |
| Rii | 6 | | | BsaWI | rCutSmart | 10 | 100 | 50 | 100 | 60° | 80° | А | λ DNA | | | |
| | 6 | | | BsaXI | rCutSmart | 50* | 100* | 10 | 100 | 37° | No | С | λ DNA | | | е |
| Rii | 6 | | | BseRI | rCutSmart | 100 | 100 | 75 | 100 | 37° | 80° | А | λ DNA | | | d |
| RX | | | | BseYI | r3.1 | 10 | 50 | 100 | 50 | 37° | 80° | В | λ DNA | | CpG | d |
| Rľ | 6 | | 2+site | Bsgl | rCutSmart | 25 | 50 | 25 | 100 | 37° | 65° | В | λ DNA | | | d |
| R% | 6 | | | BsiEl | rCutSmart | 25 | 50 | <10 | 100 | 60° | No | А | λ DNA | | CpG | |
| R% | 6 | | | BsiHKAI | rCutSmart | 25 | 100 | 100 | 100 | 65° | No | А | λ DNA | | | |
| Rľ | 6 | | | BsiWI | r3.1 | 25 | 50* | 100 | 25 | 55° | 65° | В | ΦX174 DNA | | CpG | |
| RX | 0 | e | | BsiWI-HF | rCutSmart | 50 | 100 | 10 | 100 | 37° | No | В | ΦX174 DNA | | CpG | |
| R ₁ | 6 | | | BsII | rCutSmart | 50 | 75 | 100 | 100 | 55° | No | А | λ DNA | dcm | CpG | b |
| R % | 0 | | | BsmAl | rCutSmart | 50 | 100 | 100 | 100 | 55° | No | В | λ DNA | | CpG | |

1. Star activity may result from extended digestion, high enzyme concentration or a glycerol concentration of > 5%.

Star activity may result from extended digestion.
 Star activity may result from a glycerol concentration of > 5%.

* May exhibit star activity in this buffer. + NEB isoschizomer or HF enzyme supplied with rCutSmart Buffer.

| | | | | Striction Er | - | % Activity i | , | rs | Incub. | Inactiv. | | | | | |
|----------------|---|--------|------------|----------------------|------|--------------|------|-----------|---------------|---------------|------|--------------------------------|-----|---------------------|---------|
| | | | Enzyme | Supplied NEBuffer | r1.1 | r2.1 | r3.1 | rCutSmart | Temp. (°C) | Temp. (°C) | Dil. | Unit Substrate | | ylation sitivity | Notes |
| R?? | Ø | е | BsmBI-v2 | r3.1 | <10 | 50 | | 25 | 55° | 80° | В | λ DNA | | CpG | |
| R% | | | BsmFl | rCutSmart | 25 | 50 | 50 | 100 | 65° | 80° | А | pBR322 DNA | dcm | CpG | 1 |
| R _{ | 0 | | Bsml | rCutSmart | 25 | 100 | <10 | 100 | 65° | 80° | А | λDNA | | | |
| R _* | Ø | | BsoBl | rCutSmart | 25 | 100 | 100 | 100 | 37° | 80° | А | λDNA | | | |
| R?? | 0 | | Bsp1286I | rCutSmart | 25 | 25 | 25 | 100 | 37° | 65° | А | λDNA | | | 3 |
| R% | 6 | | BspCNI | rCutSmart | 100 | 75 | 10 | 100 | 37° | 80° | А | λDNA | | | b |
| R?? | | | BspDI | rCutSmart | 25 | 75 | 50 | 100 | 37° | 80° | А | λDNA | dam | CpG | |
| RX | Ø | | BspEl | r3.1 | <10 | 10 | 100 | <10 | 37° | 80° | В | λ DNA (dam-) | dam | CpG | |
| R {{ | 9 | | BspHI | rCutSmart | 10 | 50 | 25 | 100 | 37° | 80° | А | λ DNA | | | |
| RX | | 2*site | BspMI | r3.1 | 10 | 50* | 100 | 10 | 37° | 65° | В | λDNA | | | |
| R _{ | 0 | | BspQI | r3.1 | 100* | 100* | | 100* | 50° | 80° | В | λDNA | | | 3 |
| R _{ | 6 | e | BspQI-HF | rCutSmart | 25 | 100 | 50 | 100 | 37° | 80° | В | λ DNA | | | |
| RX | 9 | | BsrBl | rCutSmart | 50 | 100 | 100 | 100 | 37° | 80° | А | λ DNA | | CpG | d |
| RX | Ø | | BsrDI | r2.1 | 10 | 100 | 75 | 25 | 37° | 80° | А | λDNA | | | 3, d |
| R? | 0 | е | BsrFI-v2 | rCutSmart | 25 | 25 | 0 | 100 | 37° | No | С | pBR322 DNA | | CpG | |
| R% | 6 | e | BsrGI-HF | rCutSmart | 10 | 100 | 100 | 100 | 37° | 80° | А | λ DNA | | | |
| | 6 | | Bsrl | r3.1 | <10 | 50 | 100 | 10 | 65° | 80° | В | ΦX174 DNA | | | b |
| R% | 6 | | BssHII | rCutSmart | 100 | 100 | 100 | 100 | 50° | 65° | В | λ DNA | | CpG | |
| R _{ | 0 | е | BssSI-v2 | rCutSmart | 10 | 25 | <10 | 100 | 37° | No | В | λDNA | | | |
| R _{ | | | BstAPI | rCutSmart | 50 | 100 | 25 | 100 | 60° | 80° | А | λ DNA | | CpG | b |
| R _{ | 0 | | BstBl | rCutSmart | 75 | 100 | 10 | 100 | 65° | No | А | λDNA | | CpG | |
| R _{ | 6 | e | BstEII-HF | rCutSmart | <10 | 10 | <10 | 100 | 37° | No | А | λ DNA | | | |
| R _{ | 0 | | BstNI | r3.1 | 10 | 100 | 100 | 75 | 60° | No | А | λDNA | | | а |
| Rľ | Ø | | BstUI | rCutSmart | 50 | 100 | 25 | 100 | 60° | No | А | λ DNA | | CpG | b |
| RX | 9 | | BstXI | r3.1 | <10 | 50 | 100 | 25 | 37° | 80° | В | λ DNA | dcm | | 3 |
| Rľ | Ø | | BstYI | rCutSmart | 25 | 100 | 75 | 100 | 60° | No | А | λ DNA | | | |
| R {{ | 9 | e | BstZ17I-HF | rCutSmart | 100 | 100 | 10 | 100 | 37° | No | А | λ DNA | | CpG | |
| R _* | Ø | | Bsu36I | rCutSmart | 25 | 100 | 100 | 100 | 37° | 80° | С | λ DNA (HindIII digest) | | | b |
| R % | 6 | | Btgl | rCutSmart | 50 | 100 | 100 | 100 | 37° | 80° | В | pBR322 DNA | | | |
| R% | | | BtgZl | rCutSmart | 10 | 25 | <10 | 100 | 60° | 80° | А | λDNA | | CpG | 3, b, d |
| R _* | 0 | | BtsCI | rCutSmart | 10 | 100 | 25 | 100 | 50° | 80° | В | λ DNA | | | |
| R% | 6 | e | BtsI-v2 | rCutSmart | 100 | 100 | 25 | 100 | 37° | No | А | λ DNA | | | 1 |
| R% | | e | BtsIMutI | rCutSmart | 100 | 50 | 10 | 100 | 55° | 80° | А | pUC19 DNA | | | b |
| | Ø | | Cac8I | rCutSmart | 50 | 75 | 100 | 100 | 37° | 65° | В | λDNA | | CpG | b |
| R % | Ø | | Clal | rCutSmart | 10 | 50 | 50 | 100 | 37° | 65° | А | λ DNA (dam-) | dam | CpG | |
| R% | Ø | 2*site | CspCI | rCutSmart | 10 | 100 | 10 | 100 | 37° | 65° | А | λDNA | | | е |
| R% | | | CviKI-1 | rCutSmart | 25 | 100 | 100 | 100 | 37° | No | А | pBR322 DNA | | | 1, b |
| R % | 0 | | CviQI | r3.1 | 75 | 100* | 100 | 75* | 25° | No | С | λDNA | | | b |
| R /{ | Ø | | Ddel | rCutSmart | 75 | 100 | 100 | 100 | 37° | 65° | В | λDNA | | | |
| RX | Ø | | Dpnl | rCutSmart | 100 | 100 | 75 | 100 | 37° | 80° | В | pBR322 DNA (dam methylated) | | CpG | b |
| R?? | 6 | | DpnII | U | 25 | 25 | 100* | 25 | 37° | 65° | В | λ DNA (dam-) | dam | | |
| R# | Ø | | Dral | rCutSmart | 75 | 75 | 50 | 100 | 37° | 65° | А | λDNA | | | |
| | | | | | | | | | | | | | | | |

a. Ligation is less than 10% b. Ligation is 25% – 75% c. Recutting after ligation is < 5% d. Recutting after ligation is 50% – 75% e. Ligation and recutting after ligation is not applicable since the enzyme is either a nicking enzyme, is affected by methylation, or the recognition sequence contains variable sequences.

| | | | | 0 | | % Activity i | n NEBuffe | ers | Incub. | Inactiv. | | | | | |
|-------------|---|--------|------------|----------------------|------|--------------|-----------|-----------|---------------|---------------|------|--------------------------------|-----|---------------------|-------|
| | | | Enzyme | Supplied NEBuffer | r1.1 | r2.1 | r3.1 | rCutSmart | Temp. (°C) | Temp. (°C) | Dil. | Unit Substrate | | ylation sitivity | Notes |
| RX | Ø | e | DrallI-HF | rCutSmart | <10 | 50 | 10 | 100 | 37° | No | В | λDNA | | CpG | b |
| R}{ | 6 | 1 | Drdl | rCutSmart | 25 | 50 | 10 | 100 | 37° | 65° | А | pUC19 DNA | | CpG | 3 |
| ?]] | | | Eael | rCutSmart | 10 | 50 | <10 | 100 | 37° | 65° | А | λ DNA | dcm | CpG | b |
| R11 | 6 | e | Eagl-HF | rCutSmart | 25 | 100 | 100 | 100 | 37° | 65° | В | pXba DNA | | CpG | |
| R]{ | 6 | 1 | Earl | rCutSmart | 50 | 10 | <10 | 100 | 37° | 65° | В | λ DNA | | CpG | b, d |
| R/{ | | | Ecil | rCutSmart | 100 | 50 | 50 | 100 | 37° | 65° | А | λ DNA | | CpG | 2 |
| Rii | Ø | 1 | Eco53kl | rCutSmart | 100 | 100 | <10 | 100 | 37° | 65° | А | pXba DNA | | CpG | 3, b |
| Rľ | Ø | 1 | EcoNI | rCutSmart | 50 | 100 | 75 | 100 | 37° | 65° | А | λ DNA | | | b |
| RII | 6 | 1 | Eco0109I | rCutSmart | 50 | 100 | 50 | 100 | 37° | 65° | А | λ DNA (HindIII digest) | dcm | | 3 |
| RN | Ø | 2*site | EcoP15I | r3.1 + ATP | 75 | 100 | 100 | 100 | 37° | 65° | А | pUC19 DNA | | | е |
| Rii | Ø | 1 | EcoRI | U | 25 | 100* | 50 | 50* | 37° | 65° | С | λ DNA | | CpG | |
| RN | Ø | е | EcoRI-HF | rCutSmart | 10 | 100 | <10 | 100 | 37° | 65° | С | λ DNA | | CpG | |
| Rłł | Ø | 1 | EcoRV | r3.1 | 10 | 50 | | 10 | 37° | 80° | А | λ DNA | | CpG | |
| RX | 6 | e | EcoRV-HF | rCutSmart | 25 | 100 | 100 | 100 | 37° | 65° | В | λ DNA | | CpG | |
| Rłł | 6 | 1 | Esp3I | rCutSmart | 100 | 100 | <10 | 100 | 37° | 65° | В | λ DNA | | CpG | |
| R/{ | | | Fatl | r2.1 | 10 | 100 | 50 | 50 | 55° | 80° | А | pUC19 DNA | | | |
| ?]] | | | Faul | rCutSmart | 100 | 50 | 10 | 100 | 55° | 65° | А | λ DNA | | CpG | 3, b, |
| ? }} | Ø | 1 | Fnu4HI | rCutSmart | <10 | <10 | <10 | 100 | 37° | No | А | λ DNA | | CpG | а |
| R]{ | | 2*site | Fokl | rCutSmart | 100 | 100 | 75 | 100 | 37° | 65° | А | λ DNA | dcm | CpG | 3, b, |
| R11 | 6 | 1 | Fsel | rCutSmart | 100 | 75 | <10 | 100 | 37° | 65° | В | pBC4 DNA | dcm | CpG | |
| ? }} | Ø | 1 | Fspl | rCutSmart | 10 | 100 | 10 | 100 | 37° | No | С | λ DNA | | CpG | b |
| ? }} | Ø | 1 | Haell | rCutSmart | 25 | 100 | 10 | 100 | 37° | 80° | А | λ DNA | | CpG | |
| ?]] | Ø | 1 | HaeIII | rCutSmart | 50 | 100 | 25 | 100 | 37° | 80° | А | λ DNA | | | |
| R}{ | | | Hgal | r1.1 | 100 | 100 | 25 | 100* | 37° | 65° | А | ΦX174 DNA | | CpG | 1 |
| ? }} | 6 | 1 | Hhal | rCutSmart | 25 | 100 | 100 | 100 | 37° | 65° | А | λ DNA | | CpG | |
| Rii | Ø | 1 | HinP1I | rCutSmart | 100 | 100 | 100 | 100 | 37° | 65° | А | λ DNA | | CpG | |
| Rii | Ø | 1 | Hincll | rCutSmart | 25 | 100 | 100 | 100 | 37° | 65° | В | λ DNA | | CpG | |
| RN | | | HindIII | r2.1 | 25 | 100 | 50 | 50 | 37° | 80° | В | λ DNA | | | 2 |
| Rii | Ø | e | HindIII-HF | rCutSmart | 10 | 100 | 10 | 100 | 37° | 80° | В | λ DNA | | | |
| ? }} | Ø | 1 | Hinfl | rCutSmart | 50 | 100 | 100 | 100 | 37° | 80° | А | λ DNA | | CpG | |
| R }} | | | Hpal | rCutSmart | <10 | 75* | 25 | 100 | 37° | No | А | λ DNA | | CpG | 1 |
| Rii | Ø | 1 | Hpall | rCutSmart | 100 | 50 | <10 | 100 | 37° | 80° | А | λ DNA | | CpG | |
| ?]] | Ø | 1 | Hphl | rCutSmart | 50 | 50 | <10 | 100 | 37° | 65° | В | λ DNA | dcm | | 1, b, |
| ? }} | 6 | 1 | Hpy166II | rCutSmart | 100 | 100 | 50 | 100 | 37° | 65° | С | pBR322 DNA | | CpG | |
| R]{ | | | Hpy188I | rCutSmart | 25 | 100 | 50 | 100 | 37° | 65° | А | pBR322 DNA | dam | | 1, b |
| R/(| | | Hpy188III | rCutSmart | 100 | 100 | 10 | 100 | 37° | 65° | В | pUC19 DNA | dam | CpG | 3, b |
| ? }} | | | Hpy99I | rCutSmart | 50 | 10 | <10 | 100 | 37° | 65° | А | λ DNA | | CpG | |
| ? }} | Ø | 1 | HpyAV | rCutSmart | 100 | 100 | 25 | 100 | 37° | 65° | | λ DNA | | CpG | 3, b, |
| ? }} | | | HpyCH4III | rCutSmart | 100 | 25 | <10 | 100 | 37° | 65° | А | λDNA | | | b |
| R11 | Ø | 1 | HpyCH4IV | rCutSmart | 100 | 50 | 25 | 100 | 37° | 65° | А | pUC19 DNA | | CpG | |
| R?? | 6 | | HpyCH4V | rCutSmart | 50 | 50 | 25 | 100 | 37° | 65° | А | λDNA | | | |

1. Star activity may result from extended digestion, high enzyme concentration or a glycerol concentration of > 5%.

Star activity may result from extended digestion.
 Star activity may result from a glycerol concentration of > 5%.

* May exhibit star activity in this buffer. + NEB isoschizomer or HF enzyme supplied with rCutSmart Buffer.

| | | | | 0 | 0 | % Activity i | n NEBuffe | rs | Incub. | Inactiv. | | | | | |
|-------------|-------------------|--------|----------|----------------------------|------|--------------|-----------|-----------|---------------|---------------|------|--|-----|---------------------|-------|
| | | | Enzyme | Supplied NEBuffer | r1.1 | r2.1 | r3.1 | rCutSmart | Temp. (°C) | Temp. (°C) | Dil. | Unit Substrate | | ylation sitivity | Notes |
| Rii | | | I-Ceul | rCutSmart | 10 | 10 | 10 | 100 | 37° | 65° | В | pBHS Scal-linearized Control Plasmid | | | |
| RX | | | I-Scel | rCutSmart | 10 | 50 | 25 | 100 | 37° | 65° | В | pGPS2 Notl-linearized Control Plasmid | | | |
| R¦{ | | | Kasl | rCutSmart | 50 | 100 | 50 | 100 | 37° | 65° | В | pBR322 DNA | | CpG | 3 |
| Rłł | e | | Kpnl-HF | rCutSmart | 100 | 25 | <10 | 100 | 37° | No | А | pXba DNA | | | |
| Rłł | 0 | | Mbol | rCutSmart | 75 | 100 | 100 | 100 | 37° | 65° | А | λ DNA (dam-) | dam | CpG | |
| Rii | Ø | 2*site | Mboll | rCutSmart | 100* | 100 | 50 | 100 | 37° | 65° | С | λ DNA (dam-) | dam | | b |
| R}} | e | | Mfel-HF | rCutSmart | 75 | 25 | <10 | 100 | 37° | No | А | λ DNA | | | |
| R }} | Ø | | MluCl | rCutSmart | 100 | 10 | 10 | 100 | 37° | No | А | λ DNA | | | |
| R }} | e | | Mlul-HF | rCutSmart | 25 | 100 | 100 | 100 | 37° | No | А | λ DNA | | CpG | |
| ? }} | 0 | | Mlyl | rCutSmart | 50 | 50 | 10 | 100 | 37° | 65° | А | λ DNA | | | b, d |
| R }} | 0 | 2*site | Mmel | rCutSmart | 50 | 100 | 50 | 100 | 37° | 65° | В | ΦX174 RF I DNA | | CpG | b, c |
| RN | 0 | | Mnll | rCutSmart | 75 | 100 | 50 | 100 | 37° | 65° | В | λ DNA | | | b |
| RN | | | Mscl | rCutSmart | 25 | 100 | 100 | 100 | 37° | 80° | С | λ DNA | dcm | | |
| R11 | 0 | | Msel | rCutSmart | 75 | 100 | 75 | 100 | 37° | 65° | А | λ DNA | | | |
| R11 | 0 | | MsII | rCutSmart | 50 | 50 | <10 | 100 | 37° | 80° | А | λ DNA | | | |
| RR | 0 | | MspA1I | rCutSmart | 10 | 50 | 10 | 100 | 37° | 65° | В | λ DNA | | CpG | |
| Rii | 0 | | Mspl | rCutSmart | 75 | 100 | 50 | 100 | 37° | No | А | λ DNA | | | |
| R11 | | | MspJI | rCutSmart + Enz. Activ. | <10 | <10 | <10 | 100 | 37° | 65° | В | pBR322 (dcm+) DNA | | | 1, e |
| R}} | Ø | | Mwol | rCutSmart | <10 | 100 | 100 | 100 | 60° | No | В | λ DNA | | CpG | |
| R# | | 2+site | Nael | rCutSmart | 25 | 25 | <10 | 100 | 37° | No | А | pXba DNA | | CpG | b |
| R{{ | | 2+site | Narl | rCutSmart | 100 | 100 | 10 | 100 | 37° | 65° | А | pXba DNA | | CpG | |
| R# | | | Nb.BbvCI | rCutSmart | 25 | 100 | 100 | 100 | 37° | 80° | А | supercoiled plasmid DNA | | | е |
| R11 | | | Nb.Bsml | r3.1 | <10 | 50 | | 10 | 65° | 80° | A | supercoiled plasmid pBR322 DNA | | | е |
| R# | | | Nb.BsrDI | rCutSmart | 25 | 100 | 100 | 100 | 65° | 80° | А | supercoiled pUC19 DNA | | | е |
| R{{ | | | Nb.BssSI | r3.1 | 10 | 100 | 100 | 25 | 37° | No | В | supercoiled pUC19 DNA | | | е |
| RN | | | Nb.Btsl | rCutSmart | 75 | 100 | 75 | 100 | 37° | 80° | A | supercoiled pUC101 DNA (dam-/dcm-) | | | е |
| R}} | Ø | | Ncil | rCutSmart | 100 | 25 | 10 | 100 | 37° | No | А | λ DNA | | CpG | b |
| RN | 0 | | Ncol | r3.1 | 100 | 100 | 100 | 100 | 37° | 80° | А | λ DNA | | | |
| Rii | 6 e | | Ncol-HF | rCutSmart | 50 | 100 | 10 | 100 | 37° | 80° | В | λ DNA | | | |
| RN | 0 | | Ndel | rCutSmart | 75 | 100 | 100 | 100 | 37° | 65° | А | λ DNA | | | |
| R}} | 0 | 2*site | NgoMIV | rCutSmart | 100 | 50 | 10 | 100 | 37° | No | А | pXba DNA | | CpG | 1 |
| R# | 6 <i>e</i> | | Nhel-HF | rCutSmart | 100 | 25 | 10 | 100 | 37° | 80° | С | λ DNA (HindIII digest) | | CpG | |
| R}} | 0 | | NIaIII | rCutSmart | <10 | <10 | <10 | 100 | 37° | 65° | В | ΦX174 RF I DNA | | | |
| R{{ | | | NIalV | rCutSmart | 10 | 10 | 10 | 100 | 37° | 65° | В | pBR322 DNA | dcm | CpG | |
| R{{ | | 2*site | NmeAIII | rCutSmart | 10 | 10 | <10 | 100 | 37° | 65° | В | ΦX174 RF I DNA | | | С |
| | 0 | | Notl | r3.1 | <10 | 50 | 100 | 25 | 37° | 65° | С | pBC4 DNA | | CpG | |
| | 9 e | | NotI-HF | rCutSmart | 25 | 100 | 25 | 100 | 37° | 65° | А | pBC4 DNA | | CpG | |
| | 9 e | | Nrul-HF | rCutSmart | 0 | 25 | 50 | 100 | 37° | No | А | λDNA | dam | CpG | |
| | 0 | | Nsil | r3.1 | 10 | 75 | 100 | 25 | 37° | 65° | В | λDNA | | | |

a. Ligation is less than 10% b. Ligation is 25% – 75% c. Recutting after ligation is < 5% d. Recutting after ligation is 50% – 75% e. Ligation and recutting after ligation is not applicable since the enzyme is either a nicking enzyme, is affected by methylation, or the recognition sequence contains variable sequences.

| | | | | | | % Activity i | in NEBuffe | rs | Incub. | Inactiv. | | | | | |
|-------------|---|------------|-----------|------------------------------|------|--------------|------------|-----------|---------------|---------------|------|--|----------------|-----|-------|
| | | | Enzyme | Supplied NEBuffer | r1.1 | r2.1 | r3.1 | rCutSmart | Temp. (°C) | Temp. (°C) | Dil. | Unit Substrate | Methy Sensi | | Notes |
| RX | Ø |) e | Nsil-HF | rCutSmart | <10 | 20 | <10 | 100 | 37° | 80° | В | λDNA | | | |
| Rii | Ø | | Nspl | rCutSmart | 100 | 100 | <10 | 100 | 37° | 65° | А | λDNA | | | |
| Rn | | | Nt.Alwl | rCutSmart | 10 | 100 | 100 | 100 | 37° | 80° | А | pUC101 DNA (dam-/dcm-) | dam | | е |
| RN | | | Nt.BbvCl | rCutSmart | 50 | 100 | 10 | 100 | 37° | 80° | А | supercoiled plasmid DNA | | CpG | е |
| RX | | | Nt.BsmAl | rCutSmart | 100 | 50 | 10 | 100 | 37° | 65° | А | supercoiled plasmid DNA | | CpG | е |
| Rn | | | Nt.BspQI | r3.1 | <10 | 25 | 100 | 10 | 50° | 80° | В | supercoiled pUC19 DNA | | | е |
| RX | | | Nt.BstNBI | r3.1 | 0 | 10 | 100 | 10 | 55° | 80° | А | T7 DNA | | | е |
| RX | | | Nt.CviPII | rCutSmart | 10 | 100 | 25 | 100 | 37° | 65° | А | pUC19 DNA | | CpG | е |
| RX | | | PI-Pspl | U + rAlbumin | 10 | 10 | 10 | 10 | 65° | No | В | pAKR7 XmnI-linearized Control Plasmid | | | |
| RX | | | PI-Scel | U + rAlbumin | 10 | 10 | 10 | 10 | 37° | 65° | В | pBSvdeX XmnI-linearized Control Plasmid | | | |
| RX | Ø |] | Pacl | rCutSmart | 100 | 75 | 10 | 100 | 37° | 65° | A | pNEB193 DNA | | | |
| RX | Ø |] | PaeR7I | rCutSmart | 25 | 100 | 10 | 100 | 37° | No | А | λ DNA (HindIII digest) | | CpG | |
| RX | | 2*site | PaqCI® | rCutSmart PaqCI Activator | 10 | 100 | 10 | 100 | 37° | 65° | В | λ DNA | | CpG | 1 |
| RX | | | Pcil | r3.1 | 50 | 75 | 100 | 50* | 37° | 80° | В | pXba DNA | | | |
| RX | Ø |] | PfIFI | rCutSmart | 25 | 100 | 25 | 100 | 37° | 65° | А | pBC4 DNA | | | b |
| Rłł | Ø | 3 | PfIMI | r3.1 | 0 | 100 | 100 | 50 | 37° | 65° | А | λDNA | dcm | | 3, b, |
| Rłł | | 2+site | Plel | rCutSmart | 25 | 50 | 25 | 100 | 37° | 65° | А | λDNA | | CpG | b, d |
| RN | | 2+site | PluTl | rCutSmart | 100 | 25 | <10 | 100 | 37° | 65° | А | pXba DNA | | CpG | b |
| Rłł | 0 | 3 | Pmel | rCutSmart | <10 | 50 | 10 | 100 | 37° | 65° | А | λDNA | | CpG | |
| RN | Ø | 3 | Pmll | rCutSmart | 100 | 50 | <10 | 100 | 37° | 65° | А | λ DNA (HindIII digest) DNA | | CpG | |
| R# | 0 | 3 | PpuMI | rCutSmart | <10 | <10 | <10 | 100 | 37° | No | В | λ DNA (HindIII digest) | dcm | | |
| RN | Ø | 3 | PshAl | rCutSmart | 25 | 50 | 10 | 100 | 37° | 65° | А | λDNA | | CpG | |
| ?]} | 0 |) <i>e</i> | Psil-v2 | rCutSmart | 25 | 50 | 10 | 100 | 37° | 65° | В | λDNA | | | 3 |
| Rłł | | | PspGI | rCutSmart | 25 | 100 | 50 | 100 | 75° | No | А | T7 DNA | dcm | | 3 |
| Rłł | | | PspOMI | rCutSmart | 10 | 10 | <10 | 100 | 37° | 65° | В | pXba DNA | dcm | CpG | |
| RX | | | PspXI | rCutSmart | <10 | 100 | 25 | 100 | 37° | No | В | λ DNA (HindIII digest) | | CpG | |
| RX | 0 |] | Pstl | r3.1 | 75 | 75 | 100 | 50* | 37° | 80° | С | λDNA | | | |
| RX | 6 |) <i>e</i> | PstI-HF | rCutSmart | 10 | 75 | 50 | 100 | 37° | No | С | λDNA | | | |
| Rii | Ø |) e | Pvul-HF | rCutSmart | 25 | 100 | 100 | 100 | 37° | No | В | pXba DNA | | CpG | |
| RN | Ø |] | Pvull | r3.1 | 50 | 100 | 100 | 100* | 37° | No | В | λDNA | | | |
| Rii | Ø |) e | Pvull-HF | rCutSmart | <10 | <10 | <10 | 100 | 37° | No | В | λDNA | | | |
| RX | 0 |] | Rsal | rCutSmart | 25 | 50 | <10 | 100 | 37° | No | А | λDNA | | CpG | |
| Rłł | | 2*site | RsrII | rCutSmart | 25 | 75 | 10 | 100 | 37° | 65° | С | λDNA | | CpG | |
| RN | Ø |) <i>e</i> | SacI-HF | rCutSmart | 10 | 50 | <10 | 100 | 37° | 65° | А | λ DNA (HindIII digest) | | CpG | |
| Rłł | 0 | 2+site | SacII | rCutSmart | 10 | 100 | 10 | 100 | 37° | 65° | А | pXba DNA | | CpG | |
| RX | Ø | 3 | Sall | r3.1 | <10 | <10 | 100 | <10 | 37° | 65° | А | λ DNA (HindIII digest) | | CpG | |
| Rłł | e |) <i>e</i> | Sall-HF | rCutSmart | 10 | 100 | 100 | 100 | 37° | 65° | А | λ DNA (HindIII digest) | | CpG | |
| RX | Ø | 3 | Sapl | rCutSmart | 75 | 50 | <10 | 100 | 37° | 65° | В | λDNA | | | |
| RN | | | Sau3AI | r1.1 | 100 | 50 | 10 | 100 | 37° | 65° | А | λDNA | | CpG | b |
| RN | | | Sau96I | rCutSmart | 50 | 100 | 100 | 100 | 37° | 65° | А | λDNA | dcm | CpG | |
| RN | 0 | e | SbfI-HF | rCutSmart | 50 | 25 | <10 | 100 | 37° | 80° | В | λ DNA | | | |

1. Star activity may result from extended digestion, high enzyme concentration or a glycerol concentration of > 5%.

Star activity may result from extended digestion.
 Star activity may result from a glycerol concentration of > 5%.

* May exhibit star activity in this buffer. + NEB isoschizomer or HF enzyme supplied with rCutSmart Buffer.

| | | | | | % Activity i | in NEBuffe | rs | Incub. | Inactiv. | | | | | |
|-------------|------------|------------------------|----------------------|------|--------------|------------|-----------|---------------|---------------|------|--------------------------------------|-----|--------------------|---------|
| | | Enzyme | Supplied NEBuffer | r1.1 | r2.1 | r3.1 | rCutSmart | Temp. (°C) | Temp. (°C) | Dil. | Unit Substrate | | ylation itivity | Notes |
| R % | в е | Scal-HF | rCutSmart | 100 | 100 | 10 | 100 | 37° | 80° | В | λDNA | | | |
| R }} | | ScrFI | rCutSmart | 100 | 100 | 100 | 100 | 37° | 65° | С | λDNA | dcm | CpG | 2, а |
| RX | | SexAl | rCutSmart | 100 | 75 | 50 | 100 | 37° | 65° | А | pBC4 DNA (dcm-) | dcm | | 3, b, d |
| R {{ | | SfaNI | r3.1 | <10 | 75 | 100 | 25 | 37° | 65° | В | ΦX174 RF I DNA | | CpG | 3, b |
| R % | | SfcI | rCutSmart | 75 | 50 | 25 | 100 | 37° | 65° | В | λDNA | | | 3 |
| R % | C 2*si | e Sfil | rCutSmart | 25 | 100 | 50 | 100 | 50° | No | С | pXba DNA | dcm | CpG | |
| R % | 0 | Sfol | rCutSmart | 50 | 100 | 100 | 100 | 37° | No | В | λ DNA (HindIII digest) | dcm | CpG | |
| R % | 2*si | e SgrAl | rCutSmart | 100 | 100 | 10 | 100 | 37° | 65° | А | λ DNA | | CpG | 1 |
| R % | 0 | Smal | rCutSmart | <10 | <10 | <10 | 100 | 37° | 65° | В | λ DNA (HindIII digest) | | CpG | b |
| R % | | Smll | rCutSmart | 25 | 75 | 25 | 100 | 55° | No | А | λ DNA | | | b |
| R % | | SnaBl | rCutSmart | 50* | 50 | 10 | 100 | 37° | 80° | А | T7 DNA | | CpG | 1 |
| R {{ | 9 e | Spel-HF | rCutSmart | 25 | 50 | 10 | 100 | 37° | 80° | С | pXba-Xbal DNA | | | |
| R {{ | | Sphl | r2.1 | 100 | 100 | 50 | 100 | 37° | 65° | В | λ DNA | | | 2 |
| RX | 9 e | SphI-HF | rCutSmart | 50 | 25 | 10 | 100 | 37° | 65° | В | λDNA | | | |
| R {{ | 0 | Srfl | rCutSmart | 10 | 50 | 0 | 100 | 37° | 65° | В | pNEB193-SrfI DNA | | CpG | |
| R {{ | 9 e | SspI-HF | rCutSmart | 25 | 100 | <10 | 100 | 37° | 65° | В | λ DNA | | | |
| R {{ | 0 | Stul | rCutSmart | 50 | 100 | 50 | 100 | 37° | No | А | λ DNA | dcm | | |
| R % | 6 | StyD4I | rCutSmart | 10 | 100 | 100 | 100 | 37° | 65° | В | λ DNA | dcm | CpG | |
| R % | 🚱 e | Styl-HF | rCutSmart | 25 | 100 | 25 | 100 | 37° | 65° | А | λ DNA | | | |
| Rii | Ø | Swal | r3.1 | 10 | 10 | 100 | 10 | 25° | 65° | В | pXba DNA | | | b, d |
| R }} | e | TaqI-v2 | rCutSmart | 50 | 100 | 50 | 100 | 65° | No | В | λDNA | dam | | |
| R }} | Ø | Tfil | rCutSmart | 50 | 100 | 100 | 100 | 65° | No | С | λDNA | | CpG | |
| R % | 0 | Tsel | rCutSmart | 75 | 100 | 100 | 100 | 65° | No | В | λ DNA | | CpG | 3 |
| R % | | Tsp45I | rCutSmart | 100 | 50 | <10 | 100 | 65° | No | А | λ DNA | | | |
| | 0 | TspMI | rCutSmart | 50* | 75* | 50* | 100 | 75° | No | В | pBC4 DNA | | CpG | d |
| R % | 6 | TspRI | rCutSmart | 25 | 50 | 25 | 100 | 65° | No | В | λ DNA | | | |
| R % | 0 | Tth111I | rCutSmart | 25 | 100 | 25 | 100 | 65° | No | В | pBC4 DNA | | | b |
| RX | | WarmStart Nt.BstNBI | r3.1 | 0 | 10 | 100 | 25 | 55° | 80° | A | T7 DNA | | | |
| RX | 0 | Xbal | rCutSmart | <10 | 100 | 75 | 100 | 37° | 65° | А | λ DNA (dam-/Hind III digest) | dam | | |
| R % | | XcmI | r2.1 | 10 | 100 | 25 | 100* | 37° | 65° | С | λDNA | | | 2 |
| RX | Ø | Xhol | rCutSmart | 75 | 100 | 100 | 100 | 37° | 65° | А | λ DNA (HindIII digest) | | CpG | b |
| R }} | G | Xmal | rCutSmart | 25 | 50 | <10 | 100 | 37° | 65° | А | pXba DNA | | CpG | 3 |
| R {{ | G | Xmnl | rCutSmart | 50 | 75 | <10 | 100 | 37° | 65° | А | λ DNA | | | b |
| R¦ł | | Zral | rCutSmart | 100 | 25 | 10 | 100 | 37° | 80° | В | λ DNA | | CpG | |

a. Ligation is less than 10% b. Ligation is 25% – 75%

c. Recutting after ligation is < 5% d. Recutting after ligation is 50% – 75% e. Ligation and recutting after ligation is not applicable since the enzyme is either a nicking enzyme, is affected by methylation, or the recognition sequence contains variable sequences.

Activity of DNA Modifying Enzymes in rCutSmart Buffer

A selection of DNA modifying enzymes were assayed in rCutSmart Buffer, in lieu of their supplied buffers. Functional activity was compared to the activity in its supplied buffer, plus required supplements. Reactions were set up according to the recommended reaction conditions, with rCutSmart Buffer replacing the supplied buffer.

| Enzyme | Activity in rCutSmart | Required Supplements |
|--|-----------------------|---------------------------|
| Antarctic Phosphatase | +++ | Requires Zn ²⁺ |
| Authenticase® | ++ | |
| Bst DNA Polymerase | +++ | |
| CpG Methyltransferase (M.Sssl) | +++ | |
| DNA Polymerase I | +++ | |
| DNA Polymerase I, Large (Klenow) Fragment | +++ | |
| DNA Polymerase Klenow Exo- | +++ | |
| DNase I (RNase-free) | +++ | Requires Ca2+ |
| DNase I-XT | +++ | Requires Ca ²⁺ |
| Duplex DNase | + | |
| E. coli DNA Ligase | +++ | Requires NAD |
| Endonuclease III (Nth), recombinant | +++ | |
| Endonuclease VIII | +++ | |
| Exonuclease I | +++ | |
| Exonuclease III | +++ | |
| Exonuclease VII | +++ | |
| Exonuclease V (Rec BCD) | +++ | Requires ATP |
| Fpg | +++ | noquilos / m |
| GpC Methyltransferase (M.CviPI) | + | Requires DTT |
| Hi-T4 [™] DNA Ligase | ++++ | Requires ATP |
| Lambda Exonuclease | +++ | nequiles All |
| Micrococcal Nuclease | | Requires Ca ²⁺ |
| Mismatch Endonuclease I | +++ | nequires da- |
| | ++ | Dequiree DTT |
| phi29 DNA Polymerase | +++ | Requires DTT |
| Quick CIP | +++ | |
| RecJ _f | +++ | D ATD |
| Salt-T4® DNA Ligase | + | Requires ATP |
| Shrimp Alkaline Phosphatase (rSAP) | +++ | |
| T3 DNA Ligase | +++ | Requires ATP + PEG |
| T4 DNA Ligase | +++ | Requires ATP |
| T4 DNA Polymerase | +++ | |
| T4 Phage β -glucosyltransferase (T4-BGT) | +++ | |
| T4 Polynucleotide Kinase | +++ | Requires ATP + DTT |
| T4 PNK (3´ phosphatase minus) | + + + | Requires ATP + DTT |
| T5 Exonuclease | + + + | |
| T7 DNA Ligase | + + + | Requires ATP + PEG |
| T7 DNA Polymerase (unmodified) | +++ | |
| T7 Exonuclease | + + + | |
| Thermolabile Exonuclease I | + + + | |
| Thermolabile USER II Enzyme | + + + | |
| Thermolabile USER III Enzyme | ++ | |
| Thermostable Endonuclease Q | +++ | |
| Thermostable OGG | +++ | |
| USER Enzyme, recombinant | +++ | |
| WarmStart [®] Afu UDG | + | |

RECOMMENDED PRODUCT

Gel Loading Dye, Purple (6X) (NEB #B7024) NEB provides Gel Loading Dye, Purple (6X) with most of our restriction enzymes.

- No UV Shadow, allowing for publication grade images
- Contains Ficoll[®] for brighter, tighter bands
- · Contains SDS for improved band sharpness
- Contains EDTA to stop enzymatic reactions
- Compatible with agarose and non-denaturing polyacrylamide gels
- Contains EDTA to stop enzymatic reactions
- Our Purple Gel Loading Dye sharpens bands and eliminates the UV shadow seen with other dyes. Available also without SDS (NEB #B7025).



COMPARISON OF DYE FRONTS

+++ full functional activity

++ 50–100% functional activity

+ 0-50% functional activity

PCR/Amplification

Amplification can be performed to generate a blunt insert, or to have a 1-base overhang, depending on the polymerase used. Additionally, primers can be used to incorporate RE recognition sites. After amplification, the insert can be used directly or cloned into a holding vector, or RE digestion can be performed to generate cohesive ends. Amplification is often the first step for PCR cloning, seamless cloning, ligation independent cloning and recombinational cloning.

PCR Polymerase Selection Chart for Cloning

For over 50 years, New England Biolabs, Inc. has been a world leader in the discovery and production of reagents for the life science industry. NEB offers a wide range of DNA polymerases, and through our commitment to research, ensures the development of innovative and high quality tools for PCR and related applications. The following table simplifies the selection of a polymerase that best suits your cloning experiment.

| | | DARD Cr | | IDELITY Cr | | IALTY Cr |
|---------------------------|---|--------------------------------------|----------------------|---|--|-----------------|
| | | | Highest | Fidelity | Long Amplicons | dU Tolerance |
| Num | One <i>Taq®/</i> One <i>Taq</i> Hot Start | <i>Taq </i> Hot Start <i>Taq</i> | Q5®/Q5 Hot Start | Phusion ^{™(1)} / Phusion ⁽¹⁾ Flex | LongAmp®/ LongAmp Hot Start <i>Taq</i> | Q5U® |
| PROPERTIES | | | | | | |
| Fidelity vs. Taq | 2X | 1X | ~280X ⁽³⁾ | > 50X | 2X | ND |
| Amplicon Size | < 6 kb | ≤ 5 kb | ≤ 20 kb | ≤ 20 kb | ≤ 30 kb | app-specific |
| Extension Time | 1 kb/min | 1 kb/min | 6 kb/min | 4 kb/min | 1.2 kb/min | 2 kb/min |
| Resulting Ends | 3´ A/Blunt | 3´ A | Blunt | Blunt | 3´ A/Blunt | Blunt |
| 3´→ 5´ exo | Yes | No | Yes | Yes | Yes | Yes |
| 5´→ 3´ exo | Yes | Yes | No | No | Yes | No |
| Units/50 µl Reaction | 1.25 | 1.25 | 1.0 | 1.0 | 5.0 | 1.0 |
| Annealing Temperature | Tm⁻5 | Tm⁻5 | Tm*3 | Tm*3 | Tm⁻5 | Tm*3 |
| APPLICATIONS | | | | | | |
| Routine PCR | * | • | • | • | ٠ | |
| Colony PCR | * | • | | | | |
| Enhanced Fidelity | • | | * | • | • | |
| High Fidelity | | | * | • | | |
| High Yield | * | • | * | • | | |
| Fast | | | * | • | | |
| Long Amplicon | | | * | • | * | |
| GC-rich Targets | * | | * | | • | |
| AT-rich Targets | * | • | * | • | • | * |
| High Throughput | • | • | • | • | | * |
| Multiplex PCR | • | ★(2) | • | • | | |
| DNA Labeling | | * | | | | |
| Site-directed Mutagenesis | | | * | • | | |
| Carryover Prevention | | | | | | * |
| USER® Cloning | | | | | | * |
| FORMATS | | | | | | |
| Hot Start Available | ٠ | • | • | • | ٠ | • |
| Kit | | • | • | • | • | • |
| Master Mix Available | • | • | • | • | • | • |
| Direct Gel Loading | ٠ | • | | | | |

(1) Phusion DNA Polymerase was developed by Finnzymes Oy, now a part of Thermo Fisher Scientific. This product is manufactured by New England Biolabs, Inc. under agreement with, and under the performance specifications of Thermo Fisher Scientific.

(2) Use Multiplex PCR 5X Master Mix.

(3) Due to the very low frequency of misincorporation events being measured, the error rate of high-fidelity enzymes like Q5 is challenging to measure in a statistically significant manner. We continue to investigate improved assays to characterize Q5's very low error rate to ensure that we present the most robust accurate fidelity data possible (Popatov, V. and Ong, J.L. (2017) PLoS One, 12(1):e0169774. doi 10.1371/journal. pone. 0169774).

GETTING STARTED

 When choosing a polymerase for PCR, we recommend starting with One *Taq* or Q5 DNA Polymerases (highlighted to the left in orange). Both offer robust amplification and can be used on a wide range of templates (routine, AT- and GC-rich). Q5 provides the benefit of maximum fidelity, and is also available in a formulation specifically optimized for next generation sequencing.

TOOLS & RESOURCES

Visit NEBPCRPolymerases.com to find:

- The full list of polymerases available
- FAQs & troubleshooting guides
- · Interactive tools to help with experimental design
- Online tutorials for setting up PCR reactions



LEARN HOW TO AMPLIFY GC-RICH DNA



* indicates recommended choice for application

For additional help with choosing the right polymerase for your PCR, we recommend using our PCR Selector at **PCRSelector.neb.com.**

Protocol: High-Fidelity PCR with Q5

| | 25 µI Reaction | 50 µI Reaction | FINAL Concentration |
|--|-------------------|-------------------|------------------------|
| 5X Q5 Reaction Buffer* | 5 µl | 10 µl | 1X |
| 10 mM dNTPs | 0.5 µl | 1 µl | 200 µM |
| 10 μ M primers (forward and reverse) | 1.25 µl | 2.5 µl | 0.5 µM |
| Template DNA | variable | variable | < 1 µg |
| Nuclease-free water | to 25 µl | to 50 µl | |
| Q5 High-Fidelity DNA Polymerase** | 0.25 µl | 0.5 µl | 0.02 units/50 µl rxn |

* Q5 High GC Enhancer can be used for difficult amplicons.

** For amplicons > 6 kb, up to 2 units/50 µl rxn can be added.

Protocol: Routine PCR with One Taq

| | 25 µI Reaction | 50 µI Reaction | FINAL Concentration |
|--------------------------------------|-------------------|-------------------|------------------------|
| One Taq Standard 5X Reaction Buffer* | 5 µl | 10 µl | 1X |
| 10 mM dNTPs | 0.5 µl | 1 µl | 200 µM |
| 10 µM primers (forward and reverse) | 0.5 µl | 1 µl | 0.2 μM |
| Template DNA | variable | variable | < 1 µg |
| Nuclease-free water | to 25 µl | to 50 µl | |
| One <i>Taq</i> DNA Polymerase** | 0.125 µl | 0.25 µl | 1.25 units/50 µl rxn |

* If reaction buffer is 5X, volume should be doubled.

** Amount of polymerase added will depend on polymerase used. Refer to neb.com for more information.

TIPS FOR OPTIMIZATION

When switching from a *Taq* product to a high-fidelity polymerase, remember to use:

- Higher annealing temps –
 check TmCalculator.neb.com
- Higher denaturation temps particularly beneficial for difficult templates
- · Higher primer concentrations
- Shorter cycling protocols

DNA TEMPLATE

- Use high-quality, purified DNA templates whenever possible. Refer to specific product information for amplification from unpurified DNA (i.e., colony or direct PCR).
- For low-complexity templates (i.e., plasmid, lambda, BAC DNA), use 1 pg–10 ng of DNA per 50 µl reaction
- For higher complexity templates (i.e., genomic DNA), use 1 ng-1 µg of DNA per 50 µl reaction
- Higher DNA concentrations tend to decrease amplicon specificity, particularly for high numbers of cycles

PRIMERS

- Primers should typically be 20–40 nucleotides in length, with 40–60% GC content
- Primer Tm values should be determined with NEB's Tm Calculator (TmCalculator.neb.com)
- Primer pairs should have Tm values that are within 5°C

- Avoid secondary structure (i.e., hairpins) within each primer and potential dimerization between the primers
- Higher than recommended primer concentrations
 may decrease specificity
- When engineering restriction sites onto the end of primers, 6 nucleotides should be added 5^{-'} to the site

ENZYME CONCENTRATION

- Optimal concentration is specific to each polymerase
- Master mix formulations already contain optimal enzyme concentrations for most applications

MAGNESIUM CONCENTRATION

- Most PCR buffers provided by NEB already contain sufficient levels of Mg⁺⁺ at 1X concentrations
- Excess Mg⁺⁺ may lead to spurious amplification; insufficient Mg⁺⁺ concentrations may cause reaction failure

DEOXYNUCLEOTIDES

- Ideal dNTP concentration is typically 200 µM each
- The presence of uracil in the primer, template, or deoxynucleotide mix will cause reaction failure when using archaeal PCR polymerases. Use One *Taq*, *Taq* or Q5U DNA Polymerases for these applications.

STARTING REACTIONS

- Unless using a hot start enzyme, assemble all reaction components on ice
- Add the polymerase last, whenever possible

| | CYCLES | TEMP. | TIME |
|--------------------------|--------|----------|----------------------|
| Initial denaturation: | 1 | 98°C | 30 seconds |
| Denaturation | | 98°C | 5–10 seconds |
| Annealing | 30 | 50-72°C* | 10–30 seconds |
| Extension | | 72°C | 20–30 seconds per kb |
| Final extension: | 1 | 72°C | 2 minutes |
| Hold: | 1 | 4–10°C | |

Tm values should be determined using the NEB Tm calculator (TmCalculator.neb.com) Please note that Q5 and Phusion[®] annealing temperature recommendations are unique.

| | CYCLES | TEMP. | ТІМЕ |
|--------------------------|--------|----------|-----------------|
| Initial denaturation: | 1 | 94°C | 30 seconds |
| Denaturation | | 94°C | 15–30 seconds |
| Annealing | 30 | 45-68°C* | 15–60 seconds |
| Extension | | 68°C | 1 minute per kb |
| Final extension: | 1 | 68°C | 5 minutes |
| Hold: | 1 | 4–10°C | |

* Tm values should be determined using the NEB Tm calculator (TmCalculator.neb.com).

 Transfer reactions to a thermocycler that has been pre-heated to the denaturation temperature. Preheating the thermocycler is not necessary when using a hot start enzyme (e.g., Q5 Hot Start or One Tag Hot Start).

DENATURATION

- Avoid longer or higher temperature incubations unless required due to high GC content of the template
- NEB's aptamer-based hot start enzymes do not require additional denaturation steps to activate the enzymes

ANNEALING

- Primer Tm values should be determined using the NEB Tm Calculator (**TmCalculator.neb.com**)
- Non-specific product formation can often be avoided by optimizing the annealing temperature or by switching to a hot start enzyme (e.g., Q5 Hot Start High-Fidelity DNA Polymerase or One *Taq* Hot Start DNA Polymerase)

EXTENSION

- Extension rates are specific to each PCR polymerase. In general, extension rates range from 15–60 s/kb.
- Longer than recommended extension times can result in higher error rates, spurious banding patterns and/or reduction of amplicon yields

Common DNA End Modifications

Modification of the termini of double-stranded DNA is often necessary to prepare the molecule for cloning. DNA ligases require a 5['] monophosphate on the donor end, and the acceptor end requires a 3['] hydroxyl group. Additionally, the sequences to be joined need to be compatible, either a blunt end being joined to another blunt end, or a cohesive end with a complementary overhang to another cohesive end. End modifications are performed to improve the efficiency of the cloning process, and ensure the ends to be joined are compatible.

Phosphorylation

Vectors and inserts digested by restriction enzymes contain the necessary terminal modifications (5[°] phosphate and 3[°] hydroxyl), while ends created by PCR may not. Typical amplification by PCR does not use phosphorylated primers. In this case, the 5[°] ends of the amplicon are non-phosphorylated and need to be treated by a kinase, such as T4 Polynucleotide Kinase (NEB #M0201), to introduce the 5[°] phosphate. Alternatively, primers for PCR can be ordered with 5[°] phosphate to avoid the need to separately phosphorylate the PCR product with a kinase.

Protocol: Phosphorylation with T4 Polynucleotide Kinase

| | STANDARD PROTOCOL |
|---------------------------------------|---------------------------------|
| DNA | 1-2 µg |
| 10X Polynucleotide Kinase Buffer | 5 µl |
| 10 mM Adenosine 5´-Triphosphate (ATP) | 5 µl (1 mM final concentration) |
| T4 Polynucleotide Kinase (PNK) | 1 µl (10 units) |
| Nuclease-free water | to 50 μl |
| Incubation | 37°C, 30 minutes |

Dephosphorylation

Dephosphorylation is a common step in traditional cloning to ensure the vector does not re-circularize during ligation. If a vector is linearized by a single restriction enzyme or has been cut with two enzymes with compatible ends, use of a phosphatase to remove the 5['] phosphate reduces the occurrence of vector re-closure by intramolecular ligation and thereby reduces the background during subsequent transformation. If the vector is dephosphorylated, it is essential to ensure the insert contains a 5['] phosphate to allow ligation to proceed. Each double-strand break requires that one intact phosphodiester bond be created before transformation (and *in vivo* repair).

Phosphatase Selection Chart

| | Recombinant Shrimp Alkaline Phosphatase (rSAP) (NEB #M0371) | Antarctic Phosphatase (AP) (NEB #M0289) | Quick CIP (NEB #M0525) |
|-------------------------------|---|---|----------------------------------|
| FEATURES | | | |
| 100% heat inactivation | 5 minutes/65°C | 2 minutes/80°C | 2 minutes/80°C |
| High specific activity | • | | • |
| Improved stability | • | | • |
| Works directly in NEB buffers | • | • | • |
| Requires additive | | (Zn²⁺) | |
| Quick Protocol | | | ٠ |

Protocol: Dephosphorylation using Quick CIP

| | STANDARD PROTOCOL |
|----------------------|---------------------|
| DNA | 1 pmol of ends |
| 10X rCutSmart Buffer | 2 µl |
| Quick CIP | 1μl |
| Nuclease-free water | to 20 μl |
| Incubation | 37°C for 10 minutes |
| Heat Inactivation | 80°C for 2 minutes |

TIPS FOR OPTIMIZATION

ENZYME

- T4 Polynucleotide Kinase (NEB #M0201) and T4 DNA Ligase (NEB #M0202) can be used together in the T4 DNA Ligase Buffer
- T4 Polynucleotide Kinase is inhibited by high levels of salt (50% inhibition by 150 mM NaCl), phosphate (50% inhibition by 7 mM phosphate) and ammonium ions (75% inhibited by 7 mM (NH_a)₂SO_a)
- If using T4 Polynucleotide Kinase and working with 5'-recessed ends, heat the reaction mixture for 10 min at 70°C, chill rapidly on ice before adding the ATP (or Ligase Buffer containing ATP) and enzyme, then incubate at 37°C

ADDITIVES

• The addition of PEG 8000 (up to 5%) can improve results



TIPS FOR OPTIMIZATION

ENZYME

- When dephosphorylating a fragment following a restriction enzyme digest, a DNA clean up step is required if the restriction enzyme(s) used is NOT heat inactivatable.
 We recommend the Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130).
- NEB recommends Quick CIP as a starting point for dephosphorylation
- When working with the Quick CIP (NEB #M0525), rSAP (NEB #M0371) or AP (NEB #M0289), which are heat-inactivatable enzymes, a DNA clean-up step after dephosphorylation is not necessary prior to the ligation step.

ADDITIVES

• AP requires the presence of Zn²⁺ in the reaction, so don't forget to supplement the reaction with 1X Antarctic Phosphatase Reaction Buffer when using other NEBuffers

Blunting/End-repair

Blunting is a process by which the single-stranded overhang created by a restriction digest is either "filled in", by adding nucleotides on the complementary strand using the overhang as a template for polymerization, or by "chewing back" the overhang, using an exonuclease activity. Vectors and inserts are often "blunted" to allow non-compatible ends to be joined. Sequence information is lost or distorted by doing this and a detailed understanding of the modification should be considered before performing this procedure. Often, as long as the sequence being altered is not part of the translated region or a critical regulatory element, the consequence of creating blunt ends is negligible. Blunting a region of translated coding sequence, however, usually creates a shift in the reading frame. DNA polymerases, such as the Klenow Fragment of DNA Polymerase I and T4 DNA Polymerase, included in our Quick Blunting Kit (NEB #E1202), are often used to fill in $(5' \rightarrow 3')$ and chew back $(3' \rightarrow 5')$. Removal of a 5' overhang can be accomplished with a nuclease, such as Mung Bean Nuclease (NEB #M0250).

Blunting Selection Chart

| | T4 DNA Polymerase* (NEB #M0203) | DNA Polymerase I, Large (Klenow) Fragment (NEB #M0210) | Quick Blunting Kit (NEB #E1201) | Mung Bean Nuclease (NEB #M0250) |
|-------------------------------------|---|---|---------------------------------------|---------------------------------------|
| APPLICATION | | | | |
| Fill in of 5 [°] overhangs | • | • | • | |
| Removal of 3' overhangs | • | • | • | • |
| Removal of 5' overhangs | | | | • |

* T4 DNA Polymerase has a strong 3' \rightarrow 5' exo activity.

Protocol: Blunting using the Quick Blunting Kit

| | STANDARD PROTOCOL |
|---------------------|--|
| DNA | up to 5 μg |
| 10X Blunting Buffer | 2.5 μl |
| 1 mM dNTP Mix | 2.5 μl |
| Blunt Enzyme Mix | 1 µl |
| Nuclease-free water | to 25 µl |
| Incubation | room temperature; 15 min for RE-digested DNA; 30 min for sheared/nebulized DNA or PCR products* |
| Heat Inactivation | 70°C, 10 minutes |

 * PCR generated DNA must be purified before blunting by using a purification kit (NEB #T1130), phenol extraction/ethanol precipitation, or gel extraction (NEB #T1120).

TIPS FOR OPTIMIZATION

ENZYME

- Make sure that you choose the correct enzyme to blunt your fragment. The Quick Blunting Kit (NEB #E1201), T4 DNA Polymerase (NEB #M0203) and DNA Polymerase I, Large (Klenow) Fragment (NEB #M0210) will fill 5' overhangs and degrade 3' overhangs. Mung Bean Nuclease (NEB #M0250) degrades 5' overhangs.
- T4 DNA Polymerase and DNA Polymerase I, Large (Klenow) Fragment are active in all NEBuffers. Please remember to add dNTPs.

CLEAN-UP

- When trying to blunt a fragment after a restriction enzyme digestion, if the restriction enzyme(s) used are heat inactivable, then a clean up step prior to blunting is not needed. Alternatively, if the restriction enzyme(s) used are not heat inactivable, a DNA clean up step is recommended prior to blunting.
- When trying to blunt a fragment amplified by PCR, a DNA clean up step (e.g., Monarch Spin PCR & DNA Cleanup Kit (5 μg), NEB #T1130) is necessary prior to the blunting step to remove the nucleotides and polymerase
- When trying to dephosphorylate a fragment after the blunting step, you will need to add a DNA clean up step (e.g., Monarch Spin PCR & DNA Cleanup Kit (5 µg), NEB #T1130) after the blunting and before the addition of the phosphatase

TEMPERATURE

 When trying to blunt a fragment with Mung Bean Nuclease, the recommended temperature of incubation is room temperature, since higher temperatures may cause sufficient breathing of the dsDNA ends that the enzyme may degrade some of the dsDNA sequence. The number of units to be used and time of incubation may be determined empirically to obtain best results.

HEAT INACTIVATION

 Mung Bean nuclease reactions should not be heat inactivated. Although Mung Bean Nuclease can be inactivated by heat, this is not recommended because the DNA begins to "breathe" before the Mung Bean Nuclease is inactivated and undesirable degradation occurs at breathing sections. Purify DNA by phenol/chloroform extraction and ethanol precipitation or spin column purification (NEB #T1130).



The DNA blunting tutorial will teach you how to identify what type of overhang you have, as well as which enzyme will blunt that end, and how.

A-tailing

Tailing is an enzymatic method to add a non-templated nucleotide to the 3['] end of a blunt, double-stranded DNA molecule. Tailing is typically done to prepare a T-vector for use in TA cloning or to A-tail a PCR product produced by a high-fidelity polymerase (not *Taq* DNA Polymerase) for use in TA cloning. TA cloning is a rapid method of cloning PCR products that utilizes stabilization of the single-base extension (adenosine) produced by *Taq* DNA Polymerase by the complementary T (thymidine) of the T-vector prior to ligation and transformation. This technique does not utilize restriction enzymes and PCR products can be used directly without modification. Additionally, PCR primers do not need to be designed with restriction sites, making the process less complicated. One drawback is that the method is non-directional; the insert can go into the vector in both orientations.

A-tailing Selection Chart

| FEATURES | Klenow Fragment (3´→5´ exo⁻) (NEB #M0212) | Taq DNA Polymerase |
|----------------------|---|--------------------|
| Reaction temperature | 37°C | 75°C |
| Heat inactivated | 75°C, 20 minutes | No |
| Nucleotide cofactor | dATP | dATP |

Protocol: A-tailing with Klenow Fragment $(3 \rightarrow 5 \text{ exo})$

| | STANDARD PROTOCOL |
|---|-----------------------|
| Purified, blunt DNA | 1—5 µg* |
| NEBuffer 2 (10X) | 5 µl |
| dATP (1 mM) | 0.5 µl (0.1 mM final) |
| Klenow Fragment (3´→5´ exo-) (NEB #M0212) | 3 µI |
| H ₂ 0 | to 50 μl |
| Incubation | 37°C, 30 minutes |

* If starting with blunt-ended DNA that has been prepared by PCR or end polishing, DNA must be purified to remove the blunting enzymes.

TIPS FOR OPTIMIZATION

If the fragment to be tailed has been amplified with a high-fidelity polymerase, the DNA needs to be purified prior to the tailing reaction. For this we recommend the Monarch Spin PCR & DNA Cleanup Kit (5 µg) (NEB #T1130). Otherwise, any high-fidelity polymerase present in the reaction will be able to remove any non-templated nucleotides added to the end of the fragments.

Vector and Insert Joining

DNA Ligation

Ligation of DNA is a critical step in many modern molecular biology workflows. The sealing of nicks between adjacent residues of a single-strand break on a double-strand substrate and the joining of double-strand breaks are enzymatically catalyzed by DNA ligases. The formation of a phosphodiester bond between the 3['] hydroxyl and 5['] phosphate of adjacent DNA residues proceeds in three steps: Initially, the ligase is self-adenylated by reaction with free ATP. Next, the adenyl group is transferred to the 5['] phosphorylated end of the "donor" strand. Lastly, the formation of the phosphodiester bond proceeds after reaction of the adenylated donor end with the adjacent 3['] hydroxyl acceptor and the release of AMP. In living organisms, DNA ligates are essential enzymes with critical roles in DNA replication and repair. In the lab, DNA ligation is performed for both cloning and non-cloning applications.

Molecular cloning is a method to prepare a recombinant DNA molecule, an extrachromosomal circular DNA that can replicate autonomously within a microbial host. DNA ligation is commonly used in molecular cloning projects to physically join a DNA vector to a sequence of interest ("insert"). The ends of the DNA fragments can be blunt or cohesive and at least one must contain a monophosphate group on its 5^{\prime} ends. Following the mechanism described above, the covalent bonds are formed and a closed circular molecule is created that is capable of transforming a host bacterial strain. The recombinant plasmid maintained in the host is then available for amplification prior to downstream applications such as DNA sequencing, protein expression, or gene expression/functional analysis.

Recently, NEB has published research on T4 DNA Ligase fidelity. This information enables improved DNA assembly methods (such as Golden Gate). Please visit www.neb. com/GoldenGate to try our free Ligase Fidelity Tools and for more information.

DNA Ligase Selection Chart for Cloning

| DNA APPLICATIONS | Instant Sticky-end Ligase Master Mix (NEB #M0370) | Blunt/TA Ligase Master Mix (NEB #M0367) | ElectroLigase® (NEB #M0369) | T4 DNA Ligase (NEB #M0202) | Quick Ligation Kit (NEB #M2200) | T3 DNA Ligase (NEB #M0317) | T7 DNA Ligase (NEB #M0318) | HiFi <i>Taq</i> DNA Ligase (NEB #M0647) | Salt-T4® DNA Ligase (NEB #M0467) | Hi-T4 ^w DNA Ligase (NEB #M2622) | NEBridge Ligase Master Mix (NEB #M1100) |
|---|--|--|--------------------------------|--------------------------------------|------------------------------------|-------------------------------|-------------------------------|--|-------------------------------------|---|--|
| Ligation of sticky ends | ••• | •• | •• | •• | ••• | •• | •• | • | •• | •• | • |
| Ligation of blunt ends | • | ••• | •• | •• | ••• | •• | | | •• | •• | • |
| T/A cloning | • | ••• | •• | •• | •• | • | • | | •• | •• | |
| Electroporation | | | ••• | •• | | | | | | •• | |
| Ligation of sticky ends only | | | | | | | ••• | | | | |
| Repair of nicks in dsDNA | •• | •• | •• | ••• | •• | •• | •• | •• | ••• | ••• | |
| High complexity library cloning | •• | •• | •• | ••• | •• | | | | | | |
| FEATURES | - | - | - | - | - | - | - | - | - | - | |
| Salt tolerance (> 2X that of T4 DNA Ligase) | | | | | | ~ | | | ~ | | |
| Ligation in 15 min. or less | 1 | 1 | | 1 | 1 | 1 | 1 | 1 | 1 | 1 | |
| Master Mix Formulation | 1 | 1 | | | | | | | | | 1 |
| Thermostable | | | | | | | | 1 | | | |
| Thermotolerant | | | | | | | | | | 1 | |
| Recombinant | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |

GETTING STARTED

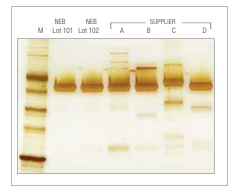
For traditional cloning, follow the ligation guidelines specified by the ligase supplier. If they suggest a 3:1 molar ratio of insert to vector, try this first for the best result. Using a 3:1 mass ratio is not the same thing (unless the insert and vector have the same mass). To calculate how much of your insert and vector to add, use **NEBioCalculator** at **NEBioCalculator.neb.com**. Ligation usually proceeds very quickly and, unless your cloning project requires the generation of a high-complexity library that benefits from the absolute capture of every possible ligation product, long incubation times are not necessary.

TOOLS & RESOURCES

Visit NEBStickTogether.com to find:

- The full list of DNA ligases available
- FAQs
- Videos about ligation and help with setting up ligation reactions

Experience extreme purity with NEB's T4 DNA Ligase



Equivalent amounts of protein were loaded and silver stained using SilverXpress[™]. Marker M is NEB's Broad Range Protein Marker (NEB #P7702).



Protocol: Ligation

| | Quick Ligation Kit (NEB #M2200) | T4 DNA Ligase (NEB #M0202) | Instant Sticky-end Master Mix (NEB #M0370) | Blunt/TA Master Mix (NEB #M0367) |
|---------------------|---|--------------------------------------|--|--|
| Format | Kit | Enzyme | Master Mix | Master Mix |
| Vector (4 kb) | 50 ng | 50 ng | 50 ng | 50 ng |
| Insert (1 kb) | 37.5 ng | 37.5 ng | 37.5 ng | 37.5 ng |
| Buffer | 2X Quick Ligation Buffer | T4 DNA Ligase Reaction Buffer | 5 µl (Master Mix) | 5 µl (Master Mix) |
| Ligase | 1 µl | 1 µl | N/A | N/A |
| Nuclease-free water | to 20 μΙ | to 20 µl | to 10 µI | to 10 μΙ |
| Incubation | 25°C, 5 minutes | 25°C, 2 hrs; 16°C, overnight* | N/A, instant ligation | 25°C, 15 minutes |

* For sticky-end ligation, the incubation time can be shortened to 25°C for 10 minutes.



For more information on the mechanisms of ligation and tips for optimization, view our videos at **NEBStickTogether.com**

TIPS FOR OPTIMIZATION

REACTION BUFFERS

- T4 DNA Ligase Buffer (NEB #B0202) should be thawed on the bench or in the palm of your hand, and not at 37°C (to prevent breakdown of ATP)
- Once thawed, T4 DNA Ligase Buffer should be placed on ice
- Ligations can also be performed in any of the four standard restriction endonuclease NEBuffers or in T4 Polynucleotide Kinase Buffer (NEB #B0201) supplemented with 1 mM ATP
- When supplementing with ATP, use ribo-ATP (NEB #P0756). Deoxyribo-ATP will inhibit ligation.
- Before ligation, completely inactivate the restriction enzyme by heat inactivation, spin column (e.g., Monarch Spin PCR & DNA Cleanup Kit (5 µg), NEB #T1130) or Phenol/ EtOH purification

DNA

• Either heat inactivate (AP, rSAP, Quick CIP) or remove phosphatase (rSAP) before ligation

- Keep total DNA concentration between 5-10 µg/ml
- Vector:Insert molar ratios between 1:1 and 1:10 are optimal for single insertions
- For cloning more than one insert, we recommend the NEBuilder HiFi DNA Assembly Master Mix (NEB #E2621) or Cloning Kit (NEB #E5520)
- If you are unsure of your DNA concentration, perform multiple ligations with varying ratios

LIGASE

- For cohesive-end ligations, standard T4 DNA Ligase. Instant Sticky-end Ligase Master Mix or the Quick Ligation Kit are recommended.
- For blunt and single-base overhangs the Blunt/TA Ligase Master Mix is recommended
- For ligations that are compatible with electroporation, Electroligase is recommended
- Standard T4 DNA Ligase can be heat inactivated at 65°C for 20 minutes
- Do not heat inactivate the Quick Ligation Kit or the ligase master mixes

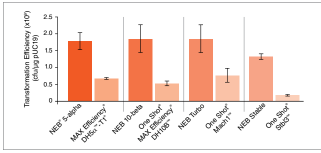
TRANSFORMATION

- Add between 1–5 µl of ligation mixture to competent cells for transformation
- Extended ligation with PEG causes a drop off in transformation efficiency
- Electroporation is recommended for larger constructs (> 10,000 bp). Dialyze samples or use a spin column first if you have used the Quick Ligation Kit or ligase master mixes.
- For ligations that are compatible with electroporation, Electroligase is recommended

Transformation

Transformation is the process by which an organism acquires exogenous DNA. Transformation can occur in two ways: natural transformation and artificial transformation. Natural transformation describes the uptake and incorporation of naked DNA from the cell's natural environment. Artificial transformation encompasses a wide array of methods for inducing uptake of exogenous DNA. In cloning protocols, artificial transformation is used to introduce recombinant DNA into host bacteria. The most common method of artificial transformation of bacteria involves use of divalent cations (e.g., calcium chloride) to increase the permeability of the bacterium's membrane, making them chemically competent, and thereby increasing the likelihood of DNA acquisition. Another artificial method of transformation is electroporation, in which cells are shocked with an electric current, to create holes in the bacterial membrane. With a newly-compromised cell membrane, the transforming DNA is free to pass into the cytosol of the bacterium. Regardless of which method of transformation is used, outgrowth of bacteria following transformation allows repair of the bacterial surface and selection of recombinant cells if the newly acquired DNA conveys antibiotic resistance to the transformed cells.

Benefit from High Transformation Efficiencies



The transformation efficiencies were compared using manufacturers' recommended protocols. Values shown are the average of triplicate experiments.

Competent Cell Selection Chart

| | NEB 5-alpha Competent <i>E. coli</i> (NEB #C2987) | NEB Turbo Competent <i>E. coli</i> (NEB #C2984) | NEB 5-alpha F [*] I ^q Competent E. coli (NEB #C2992) | NEB 10-beta Competent <i>E. coli</i> (NEB #C3019) | dam ⁻ /dcm ⁻ Competent <i>E. coli</i> (NEB #C2925) | NEB Stable Competent <i>E. coli</i> (NEB #C3040) |
|--------------------------------------|--|--|---|--|---|---|
| FEATURES | | | | | _ | |
| Versatile | • | | | | | • |
| Fast growth (< 8 hours) | | • | | | | |
| Toxic gene cloning | | | • | | | • |
| Large plasmid/BAC cloning | | | | • | | |
| Dam/Dcm-free plasmid growth | | | | | ٠ | |
| Retroviral/lentiviral vector cloning | | | | | | • |
| <i>recA</i> - | • | | • | • | | • |
| FORMATS | | | | | | |
| Chemically competent | • | • | • | • | ٠ | • |
| Electrocompetent | | | | • | | |
| Subcloning | • | | | | | |
| 96-well format | • | | | • | | |
| 384-well format | • | | | | | |
| 12 x 8-tube strips | • | | | | | |

TIPS FOR OPTIMIZATION

THAWING

- · Cells are best thawed on ice
- DNA should be added as soon as the last trace of ice in the tube disappears
- Cells can be thawed by hand, but warming above 0°C decreases efficiency

DNA

- Up to 10 μl of DNA from a ligation mix can be used with only a 2-fold loss of efficiency

INCUBATION & HEAT SHOCK

- Incubate on ice for 30 minutes. Expect a 2-fold loss in transformation efficiency (TE) for every 10 minutes this step is shortened.
- Both temperature and time are specific to the transformation volume and vessel. Typically, 30 seconds at 42°C is recommended, except when using BL21 (NEB #C2530) which requires exactly 10 seconds.

OUTGROWTH

- Outgrowth at 37°C for 1 hour is best for cell recovery and for expression of antibiotic resistance. Expect a 2-fold loss in TE for every 15 minutes this step is shortened.
- SOC gives 2-fold higher TE than LB medium
- Incubation with shaking or rotation results in 2-fold higher TE

PLATING

- Selection plates can be used warm or cold, wet or dry with no significant effects on TE
- Warm, dry plates are easier to spread and allow for the most rapid colony formation

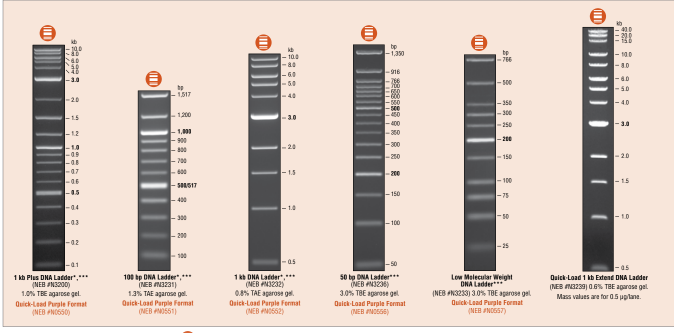
DNA CONTAMINANTS TO AVOID

| CONTAMINANT | REMOVAL METHOD |
|---------------------------|---|
| Detergents | Ethanol precipitate |
| Phenol | Extract with chloroform and ethanol precipitate |
| Ethanol or Isopropanol | Dry pellet before resuspending |
| PEG | Column purify (e.g., Monarch Spin PCR & DNA Cleanup Kit (5 µg)) or phenol/chloroform extract and ethanol precipitate |



DNA Markers and Ladders

Agarose-gel electrophoresis is the standard method used for separation, identification and purification of DNA fragments. DNA is visualized on a gel after soaking or pre-casting the gel with a visualization dye, such as Ethidium Bromide, which is a DNA intercalating agent that fluoresces under UV illumination. DNA markers and ladders are composed of DNA fragments of known sizes and masses which are used as a reference to determine the size and relative mass of the DNA of interest. Bands are visible under UV illumination or under blue light illumination, depending on the visualization dye used. DNA markers and DNA samples have to be combined with loading dyes to give them density in the wells and to track the migration on the gel; some of NEB's ladders come pre-mixed with loading dye for convenience.



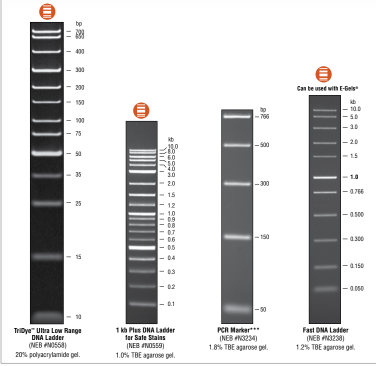
Quick-Load and Quick-Load Purple DNA Ladders

* Available in Quick-Load[®] and TriDye[™] formats

Ready-to-Load **

*** Free Loading Dye included





- Sharp, crisp bands
- Excellent quality and value
- Convenient 1 kb Plus DNA Ladder available in a variety of formats, including one specifically optimized for safe stains (e.g., GelRed and SYBR[®] Safe)
- TriDye Ultra Low Range DNA Ladder ranges as low as 10 bp and is suitable for polyacrylamide gels
- Use our selection tool to help choose a ladder at www.neb.com/ DNAmarkersandladders



Traditional Cloning Quick Guide

Preparation of insert and vectors

Insert from a plasmid source

• Digest plasmid with the appropriate restriction enzymes to produce a DNA fragment that can be cloned directly into a vector. Unidirectional cloning is achieved with restriction enzymes that produce non-compatible ends.

Insert from a PCR product

- Design primers with appropriate restriction sites to clone unidirectionally into a vector
- Addition of 6 bases upstream of the restriction site is sufficient for digestion with most enzymes
- If fidelity is a concern, choose a proofreading polymerase such as Q5 High-Fidelity DNA Polymerase (NEB #M0491)
- Visit www.NEBPCRPolymerases.com for additional guidelines for PCR optimization
- Purify PCR product by running the DNA on an agarose gel and excising the band or by using a spin column (e.g., Monarch Spin DNA Gel Extraction Kit, NEB #T1120, Monarch Spin PCR & DNA Cleanup Kit (5 μg), NEB #T1130)
- Digest with the appropriate restriction enzyme

Standard Restriction Enzyme Protocol

| DNA | 1 µg |
|------------------------|---|
| 10X NEBuffer | 5 μl (1X) |
| Restriction Enzyme | 10 units is sufficient, generally 1 μ l is used |
| Nuclease-free Water | to 50 μl |
| Incubation Time | 1 hour* |
| Incubation Temperature | Enzyme dependent |

* Can be decreased by using a Time-Saver qualified enzyme.

Time-Saver Restriction Enzyme Protocol

| DNA | 1 µg |
|------------------------|------------------|
| 10X NEBuffer | 5 µl (1X) |
| Restriction Enzyme | 1 µl |
| Nuclease-free Water | to 50 µl |
| Incubation Time | 5–15 minutes* |
| Incubation Temperature | Enzyme dependent |

* Time-Saver qualified enzymes can also be incubated overnight with no star activity.

Insert from annealed oligos

- Annealed oligos can be used to introduce a fragment (e.g., promoter, polylinker, etc.)
- Anneal two complementary oligos that leave protruding 5' or 3' overhangs for ligation into a vector cut with appropriate enzymes
- Non-phosphorylated oligos can be phosphorylated using T4 Polynucleotide Kinase (NEB #M0201)

Typical Annealing Reaction

| Oligo 1 | 20 µM Final concentration |
|---------------------|---|
| Oligo 2 | 20 µM Final concentration |
| 10X NEBuffer r2.1 | 5 μl |
| Nuclease-free Water | to 50 μl |
| Incubation | 95°C for 5 minutes, cool slowly to room temp. |

Vector

• Digest vector with appropriate restriction enzymes. Enzymes that leave non-compatible ends are ideal as they prevent vector self-ligation.

Dephosphorylation

- Dephosphorylation is sometimes necessary to prevent self-ligation. NEB offers four products for dephosphorylation of DNA:
- Quick CIP (NEB #M0525), Shrimp Alkaline Phosphatase (rSAP) (NEB #M0371) and Antarctic Phosphatase (AP) (NEB #M0289) are heat-inactivatable phosphatases. They work in all NEBuffers, but AP requires supplementation with Zn²⁺.

| Dephosphorylation of 5' ends of DNA using Quick CIP | | |
|---|---------------------|--|
| DNA | 1 pmol of DNA ends | |
| 10X rCutSmart Buffer | 2 µl | |
| Quick CIP | 1 µl | |
| Nuclease-free Water | to 20 µl | |
| Incubation | 37°C for 10 minutes | |

80°C for 2 minutes

Note: Scale larger reaction volumes proportionally.

Blunting

Heat Inactivation

- In some instances, the ends of the insert or vector require blunting
- PCR with a proofreading polymerase will leave a predominantly blunt end
- T4 DNA Polymerase (NEB #M0203) or Klenow (NEB #M0210) will fill in a 5' overhang and chew back a 3' overhang
- The Quick Blunting Kit (NEB #E1201) is optimized to blunt and phosphorylate DNA ends for cloning in less than 30 minutes
- Analyze agarose gels with longwave UV (365 nM) to minimize UV exposure that may cause DNA damage

Blunting with the Quick Blunting Kit

| DNA | Up to 5 µg |
|---------------------|---|
| Blunting Buffer | 2.5 µl |
| dNTP Mix (1 mM) | 2.5 µl |
| Blunt Enzyme Mix | 1 µl |
| Nuclease-free Water | to 25 μl |
| Incubation | room temperature; 15 min for RE-digested DNA; 30 min for sheared/nebulized DNA or PCR products* |
| Heat Inactivation | 70°C for 10 minutes |

* PCR-generated DNA must be purified before blunting using a purification kit (NEB #T1130), phenol extraction/ethanol precipitation or gel extraction (NEB #T1120).



Getting started with Molecular Cloning:

Simple tips to improve your cloning efficiency.

Phosphorylation

- For ligation to occur, at least one of the DNA ends (insert or vector) should contain a 5' phosphate
- Primers are usually supplied non-phosphorylated; therefore, the PCR product will not contain a 5' phosphate
- Digestion of DNA with a restriction enzyme will always produce a 5' phosphate
- A DNA fragment can be phosphorylated by incubation with T4 Polynucleotide Kinase (NEB #M0201). T4 PNK can be inactivated at 65°C for 20 minutes.

Phosphorylation with T4 PNK

| DNA (20 mer) | up to 300 pmol of 5´ termini |
|---------------------|------------------------------|
| 10X T4 PNK Buffer | 5 µl |
| 10 mM ATP | 5 µl (1 mM final conc.) |
| T4 PNK | 1 µl (10 units) |
| Nuclease-free Water | to 50 μl |
| Incubation | 37°C for 30 minutes |

Purification of Vector and Insert

- Purify the vector and insert by either running the DNA on an agarose gel and excising the appropriate bands or by using a spin column, such as Monarch Spin DNA Gel Extraction Kit or Monarch Spin PCR & DNA Cleanup Kit (5 μ g) (NEB #T1120 or T1130)
- DNA can also be purified using β-Agarase I (NEB #M0392) with low melt agarose or an appropriate spin column or resin
- Analyze agarose gels with longwave UV (365 nM) to minimize UV exposure that may cause DNA damage

Ligation of Vector and Insert

- Use a molar ratio of 1:3 vector to insert. Use NEBioCalculator to calculate molar ratios.
- If using T4 DNA Ligase (NEB #M0202) or the Quick Ligation Kit (NEB #M2200), thaw and resuspend the Ligase Buffer at room temp. If using Ligase Master Mixes, no thawing is necessary.
- The Quick Ligation Kit (NEB #M2200) is optimized for ligation of both sticky and blunt ends
- Instant Sticky-end Ligase Master Mix (NEB #M0370) is optimized for instant ligation of sticky/cohesive ends
- Blunt/TA Ligase Master Mix (NEB #M0367) is optimized for ligation of blunt or single base overhangs, which are the more challenging type of ends for T4 DNA Ligase
- Following ligation, chill on ice and transform
- DO NOT heat inactivate when using the Quick Ligation Buffer or Ligase Master Mixes, as this will inhibit transformation
- Electroligase[®] (NEB #M0369) is optimized for ligation of both sticky and blunt ends and is compatible with electroporation (i.e., no cleanup step required)
- Improved Golden Gate Assembly can be achieved by selecting high fidelity overhangs [Potapov, V. et al (2018) ACS Synth. Biol. 7(11),

2665–2674].

Ligation with the Quick Ligation Kit

| Vector DNA (4 kb) | 50 ng |
|--------------------------|--------------------------------|
| Insert DNA (1 kb) | 37.5 ng |
| 2X Quick Ligation Buffer | 10 µl |
| Quick T4 DNA Ligase | 1µl |
| Nuclease-free Water | 20 µl (mix well) |
| Incubation | Room temperature for 5 minutes |

Ligation with Instant Sticky-end Ligase Master Mix

| Vector DNA (3 kb) | 50 ng |
|---------------------|----------|
| Insert DNA (1 kb) | 50 ng |
| Master Mix | 5 μΙ |
| Nuclease-free Water | to 10 μl |
| Incubation | None |

Ligation with Blunt/TA Ligase Master Mix

| Vector DNA (3 kb) | 50 ng |
|---------------------|---------------------------------|
| Insert DNA (1 kb) | 50 ng |
| Master Mix | 5 μΙ |
| Nuclease-free Water | to 10 µl |
| Incubation | Room temperature for 15 minutes |

Transformation

- If recombination is a concern, then use the *recA⁻* strains NEB 5-alpha Competent *E. coli* (NEB #C2987), or NEB-10 beta Competent *E. coli* (NEB #C3019) or NEB Stable Competent *E. coli* (NEB #C3040)
- NEB-10 beta Competent E. coli works well for constructs larger than 5 kb
- NEB Stable Competent *E. coli* (NEB #C3040) can be used for constructs with repetitive sequences such as lentiviral constructs
- If electroporation is required, use NEB 10-beta Electrocompetent *E. coli* (NEB #C3020)
- Use pre-warmed selection plates
- Perform several 10-fold serial dilutions in SOC or NEB 10-beta/Stable Outgrowth Medium for plating

Transformation with NEB 5-alpha Competent E. coli

| DNA | 1–5 μl containing 1 pg–100 ng of plasmid DNA |
|-------------------|--|
| Competent E. coli | 50 μl |
| Incubation | On ice for 30 minutes |
| Heat Shock | Exactly 42°C for exactly 30 seconds |
| Incubation | On ice for 5 minutes Add 950 µl room temperature SOC 37°C for 60 minutes, with shaking |

Troubleshooting Guide for Cloning

We strongly recommend running the following controls during transformations. These controls may help troubleshoot which step(s) in the cloning workflow has failed.

- 1 Transform 100 pg 1 ng of uncut vector to check cell viability, calculate transformation efficiency and verify the antibiotic resistance of the plasmid.
- 2 Transform the cut vector to determine the amount of background due to undigested plasmid. The number of colonies in this control should be < 1% of the number of colonies in the uncut plasmid control transformation (from control #1).
- 3 Transform a vector only ligation reaction. The ends of the vector should not be able to re-ligate because either they are incompatible (e.g., digested with two restriction enzymes that do not generate compatible ends) or the 5' phosphate group has been removed in a dephosphorylation reaction (e.g., blunt ends treated with rSAP). This control transformation should yield the same number of colonies as control #2.
- 4 Digest vector DNA with a single restriction enzyme, re-ligate and transform. The ends of the vector DNA should be compatible and easily joined during the ligation reaction, resulting in approximately the same number of colonies as control #1.

The cloning workflow often benefits from an accurate quantitation of the amount of DNAs that are being worked with. We recommend quantification of DNAs whenever possible.

| PROBLEM | CAUSE | SOLUTION |
|----------------------------|--|---|
| | Cells are not viable | • Transform an uncut plasmid (e.g., pUC19) and calculate the transformation efficiency of the competent cells. If the transformation efficiency is low (<10 ⁴) re-make the competent cells or consider using commercially available high efficiency competent cells. |
| | Incorrect antibiotic or antibiotic concentration | Confirm antibiotic and antibiotic concentration |
| | DNA fragment of interest is toxic to the cells | Incubate plates at lower temperature (25–30°C) Transformation may need to be carried out using a strain that exerts tighter transcriptional control over the DNA fragment of interest [e.g., NEB-5-alpha F' I^o Competent <i>E. coli</i> (NEB #C2992)] |
| | If using chemically competent cells, the wrong heat-shock protocol was used | Follow the manufacturer's specific transformation protocol (Note: going above the recommended temperature during the heat shock can result in competent cell death) |
| | If using electrocompetent cells, PEG is present in the ligation mix | Clean up DNA by drop dialysis prior to transformation with Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130) Try NEB's ElectroLigase (NEB #M0369) |
| | If using electrocompetent cells, arcing was observed or no voltage was registered | Clean up the DNA prior to the ligation step Tap the cuvette to get rid of any trapped air bubbles Be sure to follow the manufacturer's specified electroporation parameters |
| | Construct is too large | Select a competent cell strain that can be transformed efficiently with large DNA constructs [≥ 10 kb, we recommend trying NEB 10-beta Competent <i>E. coli</i> (NEB #C3019)] or NEB Stable Competent <i>E. coli</i> (NEB #C3040) For very large constructs (> 10 kb), consider using electroporation |
| | Construct may be susceptible to recombination | • Select a <i>recA</i> - strain such as NEB 5-alpha (NEB #C2987) or NEB 10-beta Competent <i>E. coli</i> (NEB #C3019) or NEB Stable Competent <i>E. coli</i> (NEB #C3040) |
| Few or no transformants | The insert comes directly from mammalian or plant DNA and contains methylated cytosines, which are degraded by many <i>E. coli</i> strains | • Use a strain that is deficient in McrA, McrBC and Mrr, such as NEB 10-beta Competent <i>E. coli</i> |
| | Too much ligation mixture was used | \bullet Use < 5 μl of the ligation reaction for the transformation |
| | Inefficient ligation | Make sure that at least one fragment being ligated contains a 5' phosphate moiety Vary the molar ratio of vector to insert from 1:1 to 1:10. Use NEBioCalculator to calculate molar ratios. Purify the DNA to remove contaminants such as salt and EDTA with Monarch Spin PCR & DNA Cleanup Kit (5 µg) (NEB #T1130) ATP will degrade after multiple freeze-thaws; repeat the ligation with fresh buffer Heat inactivate or remove the phosphatase prior to ligation Ligation of single base-pair overhangs (most difficult) may benefit from being carried out with Blunt/TA Master Mix (NEB #M0367), Quick Ligation Kit (NEB #M2200) or concentrated T4 DNA Ligase (NEB #M0202) |
| | Inefficient phosphorylation | Purify the DNA prior to phosphorylation with Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130). Excess salt, phosphate or ammonium ions may inhibit the kinase. If the ends are blunt or 5' recessed, heat the substrate/buffer mixture for 10 minutes at 70°C. Rapidly chill on ice before adding the ATP and enzyme, then incubate at 37°C. ATP was not added. Supplement the reaction with 1 mM ATP, as it is required by T4 Polynucleotide Kinase (NEB #M0201) Alternatively, use 1X T4 DNA Ligase Buffer (contains 1 mM ATP) instead of the 1X T4 PNK Buffer |
| | Inefficient blunting | Heat inactivate or remove the restriction enzymes prior to blunting Clean up the PCR fragment prior to blunting with Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130) Sonicated gDNA should be blunted for at least 30 minutes Do not use > 1 unit of enzyme/μg of DNA Do not incubate for > 15 minutes Do not incubate at temperatures > 12°C (for T4 DNA Polymerase, NEB #M0203) or > 24°C (for Klenow, NEB #M0210) Make sure to add a sufficient amount of dNTPs to the reaction (33 μM each dNTP for DNA Polymerase I, Large (Klenow) Fragment, NEB #M0210 and 100 μM each dNTP for T4 DNA Polymerase, NEB #M0203). When using Mung Bean Nuclease (NEB #M0250), incubate the reaction at room temperature. Do not use > 1 unit of enzyme/μg DNA or incubate the reaction > 30 minutes. |

Troubleshooting Guide for Cloning (cont.)

| PROBLEM | CAUSE | SOLUTION | |
|--|---|---|--|
| | Inefficient A-Tailing | Clean up the PCR prior to A-tailing. NEB recommends the Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130). High-fidelity enzymes will remove any non-templated nucleotides. | |
| Few or no transformants (cont.) | | Check the methylation sensitivity of the enzyme(s) to determine if the enzyme is blocked by methylation of the recognition sequence Use the recommended buffer supplied with the restriction enzyme Clean up the DNA to remove any contaminants that may inhibit the enzyme. NEB recommends the Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130). When digesting a PCR fragment, make sure to have at least 6 nucleotides between the recognition site and the end of the DNA molecule | |
| Colonies don't | Antibiotic level used was too low | Increase the antibiotic level on plates to the recommended amount Use fresh plates with fresh antibiotics | |
| contain a plasmid | Satellite colonies were selected | Choose large, well-established colonies for analysis | |
| | Recombination of the plasmid has occurred | • Use a recA ⁻ strain such NEB 5-alpha, or NEB 10-beta Competent E. coli, or NEB Stable Competent E. coli | |
| | Incorrect PCR amplicon was used during cloning | Optimize the PCR conditions Gel purify the correct PCR fragment. NEB recommends the Monarch Spin DNA Gel Extraction Kit (NEB #T1120). | |
| Colonies contain the wrong construct | Internal recognition site was present | • Use NEBcutter to analyze insert sequence for presence of an internal recognition site | |
| wrong construct | DNA fragment of interest is toxic to the cells | Incubate plates at lower temperature (25–30°C) Transformation may need to be carried out using a strain that exerts tighter transcriptional control of the DNA fragment of interest (e.g., NEB 5-alpha F' / Competent E. coli) | |
| | Mutations are present in the sequence | Use a high-fidelity polymerase (e.g., Q5 High-Fidelity DNA Polymerase, NEB #M0491) Re-run sequencing reactions | |
| | Inefficient dephosphorylation | Heat inactivate or remove the restriction enzymes prior to dephosphorylation | |
| | Kinase is present/active | • Heat inactivate the kinase after the phosphorylation step. Active kinase will re-phosphorylate the dephosphorylated vector. | |
| Too much background | Restriction enzyme(s) didn't cleave completely | Check the methylation sensitivity of the restriction enzyme(s) to be sure it is not inhibited by methylation of the recognition sequence Use the recommended buffer supplied with the restriction enzyme Clean up the DNA to remove contaminants (e.g., too much salt). NEB recommends the Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130). | |
| | Antibiotic level is too low | Confirm the correct antibiotic concentration | |
| Ran the ligation on a gel and saw no ligated product | Inefficient ligation | Make sure at least one DNA fragment being ligated contains a 5' phosphate Vary the molar ratios of vector to insert from 1:1 to 1:10. Use NEBioCalculator to calculate molar ratios. Purify the DNA to remove contaminants such as salt and EDTA. NEB recommends the Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130). ATP will degrade after multiple freeze-thaws; repeat the ligation with fresh buffer Heat inactivate or remove the phosphatase prior to ligation Ligation of single base-pair overhangs (most difficult) may benefit from being carried out with Blunt/TA Master Mix, Quick Ligation Kit or concentrated T4 DNA Ligase | |
| The ligated DNA ran as a smear on an agarose gel | The ligase is bound to the substrate DNA | • Treat the ligation reaction with Proteinase K (NEB #P8107) prior to running on a gel | |
| The digested DNA | The restriction enzyme(s) is bound to the substrate DNA | Lower the number of units Add SDS (0.1–0.5%) to the loading buffer to dissociate the enzyme from the DNA or use Gel Loading Dye, Purple (6X) (NEB #B7024) | |
| ran as a smear on an agarose gel | Nuclease contamination | Use fresh, clean running buffer Use a fresh agarose gel Clean up the DNA. NEB recommends the Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130). | |
| | Cleavage is blocked by methylation | DNA isolated from a bacterial source may be blocked by Dam and Dcm methylation DNA isolated from eukaryotic source may be blocked by CpG methylation Check the methylation sensitivity of the enzyme(s) to determine if the enzyme is blocked by methylation of the recognition sequence If the enzyme is inhibited by Dam or Dcm methylation, grow the plasmid in a <i>dam-/dcm-</i> strain (NEB #C2925) | |
| | Salt inhibition | Enzymes that have low activity in salt-containing buffers (NEBuffer r3.1) may be salt sensitive, so clean up the DNA prior to digestion. NEB recommends the Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130). DNA purification procedures that use spin columns can result in high salt levels, which inhibit enzyme activity. Monarch Spin kits (NEB #T1110, #T1120, #T1130) use columns that have been designed to minimize salt carry over into the eluted DNA, so using them can minimize this issue. To prevent this, DNA solution should be no more than 25% of total reaction volume. | |
| | Inhibition by PCR components | • Clean up the PCR fragment prior to restriction digest. NEB recommends the Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130). | |
| Incomplete restriction | Using the wrong buffer | Use the recommended buffer supplied with the restriction enzyme | |
| enzyme digestion | Too few units of enzyme used | • Use at least 5–10 units of enzyme per µg of DNA | |
| | Incubation time was too short | Increase the incubation time | |
| | Digesting supercoiled DNA | • Some enzymes have a lower activity on supercoiled DNA. Increase the number of enzyme units in the reaction. | |
| | Presence of slow sites | • Some enzymes can exhibit slower cleavage towards specific sites. Increase the incubation time, 1–2 hours is typically sufficient. | |
| | Two sites required | • Some enzymes require the presence of two recognition sites to cut efficiently. For more information, visit the table "Restriction Enzymes Requir- ing Multi-sites" on neb.com. | |
| | DNA is contaminated with an inhibitor | Assay substrate DNA in the presence of a control DNA. Control DNA will not cleave if there is an inhibitor present. Miniprep DNA is particularly susceptible to contaminants. NEB recommends the Monarch Spin PCR & DNA Cleanup Kit (5 µg) (NEB #T1130). Clean DNA with a spin column with Monarch Spin PCR & DNA Cleanup Kit (5 µg) (NEB #T1130), resin or drop dialysis, or increase volume to dilute contaminant | |

| PROBLEM | CAUSE | SOLUTION | |
|--|---|--|--|
| If larger bands than expected are seen in the gel, this may indicate binding of the enzyme(s) to the substrate Star activity | | Lower the number of units in the reaction Add SDS (0.1–0.5%) to the loading buffer to dissociate the enzyme from the substrate or add Gel Loading Dye, Purple (6X) (NEB #B7024) | |
| | | Use the recommended buffer supplied with the restriction enzyme Decrease the number of enzyme units in the reaction Make sure the amount of enzyme added does not exceed 10% of the total reaction volume. This ensures that the total glycerol concentration does not exceed 5% v/v Decrease the incubation time. Using the minimum reaction time required for complete digestion will help prevent star activity. Try using a High-Fidelity (HF) restriction enzyme. HF enzymes have been engineered for reduced star activity. | |
| | Partial restriction enzyme digest | Enzymes that have low activity in salt-containing buffers (e.g., NEBuffer r3.1) may be salt sensitive. Make sure to clean up the DNA prior to digestion. NEB recommends the Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130). DNA purification procedures that use spin columns can result in high salt levels, which inhibit enzyme activity. Monarch Spin kits (NEB #T1110, #T1120, #T1130) use columns that have been designed to minimize salt carry over into the eluted DNA, so using them can minimize this issue. To prevent this, DNA solution should be no more than 25% of total reaction volume Clean-up the PCR fragment prior to restriction digest. NEB recommends the Monarch Spin PCR & DNA Cleanup Kit (5 μg) (NEB #T1130). Use the recommended buffer supplied with the restriction enzyme Use at least 3–5 units of enzyme per μg of DNA Digest the DNA for 1–2 hours | |
| Used the wrong primer sequence | | Double check the primer sequence | |
| | Incorrect annealing temperature | • Use the NEB Tm calculator to determine the correct annealing temperature (www.neb.com/TmCalculator) | |
| | Incorrect extension temperature | • Each polymerase type has a different extension temperature requirement. Follow the manufacturer's recommendations. | |
| No PCR fragment | Too few units of polymerase | Use the recommended number of polymerase units based on the reaction volume | |
| amplified | Incorrect primer concentration | • Each polymerase has a different primer concentration requirement. Make sure to follow the manufacturer's recommendations. | |
| | Mg ²⁺ levels in the reaction are not optimal | • Titrate the Mg ²⁺ levels to optimize the amplification reaction. Follow the manufacturer's recommendations. | |
| | Difficult template | • With difficult templates, try different polymerases and/or buffer combinations | |
| The PCR reaction is a smear on a gel | If bands are larger than expected it may indicate binding of the enzyme(s) to the DNA | • Add SDS (0.1–0.5%) to the loading buffer to dissociate the enzyme from the DNA | |
| | Annealing temperature is too low | Use the NEB Tm calculator to determine the annealing temperature of the primers | |
| Extra bands in | Mg ²⁺ levels in the reaction are not optimal | • Titrate the Mg ²⁺ levels to optimize the amplification reaction. Make sure to follow the manufacturer's recommendations. | |
| PCR reaction | Additional priming sites are present | • Double check the primer sequence and confirm it does not bind elsewhere in the DNA template | |
| | Formation of primer dimers | • Primer sequence may not be optimal. Additional primers may need to be tested in the reaction. | |
| | Incorrect polymerase choice | • Try different polymerases and/or buffer combinations | |

Selected Products for PCR & Mutagenesis

| | | 0175 |
|---|------------------|------------------------------|
| PRODUCT | NEB # | SIZE |
| HIGH-FIDELITY DNA POLYMERASES | | |
| Q5 High-Fidelity DNA Polymerase | M0491S/L | 100/500 units |
| Q5 Hot Start High-Fidelity DNA Polymerase | M0493S/L | 100/500 units |
| Q5 High-Fidelity 2X Master Mix | M0492S/L | 100/500 reactions |
| Q5 Hot Start High-Fidelity 2X Master Mix | M0494S/L/X | 100/500/500 reactions |
| Q5 High-Fidelity PCR Kit | E0555S/L | 50/200 reactions |
| Q5U Hot Start High-Fidelity DNA Polymerase | M0515S/L | 100/500 units |
| Phusion High-Fidelity PCR Master Mix with HF Buffer | M0531S/L | 100/500 reactions |
| Phusion High-Fidelity PCR Master Mix with GC Buffer | M0532S/L | 100/500 reactions |
| Phusion Hot Start Flex 2X Master Mix | M0536S/L | 100/500 reactions |
| Phusion High-Fidelity PCR Kit | E0553S/L | 50/200 reactions |
| Phusion High-Fidelity DNA Polymerase | M0530S/L | 100/500 units |
| Phusion Hot Start Flex High-Fidelity DNA Polymerase | M0535S/L | 100/500 units |
| DNA POLYMERASES | | |
| One Taq DNA Polymerase | M0480S/L/X | 200/1,000/5,000 units |
| One Taq Quick-Load DNA Polymerase | M0509S/L/X | 100/500/2,500 units |
| One Taq Hot Start DNA Polymerase | M0481S/L/X | 200/1,000/5,000 units |
| One Taq 2X Master Mix with Standard Buffer | M0482S/L | 100/500 reactions |
| One <i>Taq</i> Quick-Load 2X Master Mix with Standard Buffer | M0486S/L | 100/500 reactions |
| One Taq Hot Start 2X Master Mix with Standard Buffer | M0484S/L | 100/500 reactions |
| One <i>Taq</i> Hot Start 2X Master Mix with GC Buffer | M0485S/L | 100/500 reactions |
| One <i>Taq</i> Hot Start Quick-Load 2X Master Mix with Standard Buffer | M0488S/L | 100/500 reactions |
| One <i>Taq</i> Hot Start Quick-Load 2X Master Mix with GC Buffer | M0489S/L | 100/500 reactions |
| <i>Taq</i> DNA Polymerase with ThermoPol [™] Buffer | M0267S/L/X/E | 400/2,000/4,000/20,000 units |
| <i>Taq</i> DNA Polymerase with Standard <i>Taq</i> Buffer | M0273S/L/X | 400/2,000/4,000 units |
| <i>Taq</i> DNA Polymerase with Standard <i>Taq</i> (Mg-free) Buffer | M0320S/L | 400/2,000 units |
| Taq PCR Kit | E5000S | 200 reactions |
| Quick-Load Taq 2X Master Mix | M0271L | 500 reactions |
| Taq 2X Master Mix | M0270L | 500 reactions |
| Taq 5X Master Mix | M0285L | 500 reactions |
| Multiplex PCR 5X Master Mix | M0284S | 100 reactions |
| Hot Start Taq DNA Polymerase | M0495S/L | 200/1,000 units |
| Hot Start Taq 2X Master Mix | M0496S/L | 100/500 reactions |
| Vent DNA Polymerase | M0254S/L | 200/1,000 units |
| Vent (exo ⁻) DNA Polymerase | M0257S/L | 200/1,000 units |
| Deep Vent DNA Polymerase | M0258S/L | 200/1,000 units |
| Deep Vent (exo-) DNA Polymerase | M0259S/L | 200/1,000 units |
| LongAmp Taq DNA Polymerase | M0323S/L | 500/2,500 units |
| LongAmp Hot Start <i>Taq</i> DNA Polymerase | M0534S/L | 500/2,500 units |
| LongAmp Taq 2X Master Mix | M0287S/L | 100/500 reactions |
| LongAmp Hot Start Taq 2X Master Mix | M0533S/L | 100/500 reactions |
| LongAmp Taq PCR Kit | E5200S | 100 reactions |
| PCR CLONING & MUTAGENESIS | | |
| NEB PCR Cloning Kit | E1202S | 20 reactions |
| NEB PCR Cloning Kit (Without Competent Cells) | E1203S | 20 reactions |
| Q5 Site-Directed Mutagenesis Kit | E0554S | 10 reactions |
| Q5 Site-Directed Mutagenesis Kit | E05529 | 10 reactions |
| (Without Competent Cells) KLD Enzyme Mix | E0552S M0554S | 10 reactions 25 reactions |
| | 0100010 | 201000000 |

Products for cDNA Synthesis

| PRODUCT | NEB # | SIZE |
|--|------------|---------------------------|
| Deoxynucleotide (dNTP) Solution Set | N0446S | 25 µmol of each |
| Deoxynucleotide (dNTP) Solution Mix | N0447S/L | 8/40 µmol of each |
| AMV Reverse Transcriptase | M0277S/L | 200/1,000 units |
| LunaScript RT SuperMix Kit | E3010S/L | 25/100 reactions |
| LunaScript RT Master Mix Kit (Primer-free) | E3025S/L | 25/100 reactions |
| LunaScript RT SuperMix | M3010L/X/E | 100/500/2,500 reactions |
| M-MuLV Reverse Transcriptase | M0253S/L | 10,000/50,000 units |
| ProtoScript II First Strand cDNA Synthesis Kit | E6560S/L | 30/150 reactions |
| ProtoScript First Strand cDNA Synthesis Kit | E6300S/L | 30/150 reactions |
| Template Switching RT Enzyme Mix | M0466S/L | 20/100 reactions |
| ProtoScript II Reverse Transcriptase | M0368S/L/X | 4,000/10,000/40,000 units |
| WarmStart® RTx Reverse Transcriptase | M0380S/L | 50/250 reactions |

Products for Restriction Digestion

| PRODUCT | NEB # | SIZE |
|---------------------------------------|--------------|---------------------|
| HIGH-FIDELITY (HF®) RESTRICTION ENZYM | IES | |
| Agel-HF | R3552S/L | 300/1,500 units |
| Apol-HF | R3566S/L | 1,000/5,000 units |
| BamHI-HF | R3136S/L/T/M | 10,000/50,000 units |
| BbsI-HF | R3539S/L | 300/1,500 units |
| BcII-HF | R3160S/L | 3,000/15,000 units |
| BmtI-HF | R3658S/L | 300/1,500 units |
| Bsal-HFv2 | R3733S/L | 1,000/5,000 units |
| BsiWI-HF | R3553S/L | 300/1,500 units |
| BspQI-HF | R3712S/L | 500/2,500 units |
| BsrGI-HF | R3575S/L | 1,000/5,000 units |
| BstEII-HF | R3162S/L/M | 2,000/10,000 units |
| BstZ171-HF | R3594S/L | 1,000/5,000 units |
| DrallI-HF | R3510S/L | 1,000/5,000 units |
| Eagl-HF | R3505S/L/M | 500/2,500 units |
| EcoRI-HF | R3101S/L/T/M | 10,000/50,000 units |
| EcoRV-HF | R3195S/L/T/M | 4,000/20,000 units |
| HindIII-HF | R3104S/L/T/M | 10,000/50,000 units |
| KpnI-HF | R3142S/L/M | 4,000/20,000 units |
| Mfel-HF | R3589S/L | 500/2,500 units |
| Mlul-HF | R3198S/L | 1,000/5,000 units |
| Ncol-HF | R3193S/L/M | 1,000/5,000 units |
| Nhel-HF | R3131S/L/M | 1,000/5,000 units |
| NotI-HF | R3189S/L/M | 500/2,500 units |
| Nrul-HF | R3192S/L | 1,000/5,000 units |
| Nsil-HF | R3127S/L | 1,000/5,000 units |
| PstI-HF | R3140S/L/T/M | 10,000/50,000 units |
| Pvul-HF | R3150S/L | 500/2,500 units |
| Pvull-HF | R3151S/L/M | 5,000/25,000 units |
| SacI-HF | R3156S/L/M | 2,000/10,000 units |
| Sall-HF | R3138S/L/T/M | 2,000/10,000 units |
| SbfI-HF | R3642S/L | 500/2,500 units |
| Scal-HF | R3122S/L/M | 1,000/5,000 units |
| SpeI-HF | R3133S/L/M | 500/2,500 units |
| SphI-HF | R3182S/L/M | 500/2,500 units |
| SspI-HF | R3132S/L/M | 1,000/5,000 units |
| Styl-HF | R3500S/L | 3,000/15,000 units |
| FEATURED GEL LOADING DYE | | |
| Gel Loading Dye, Purple (6X) | B7024S | 4 ml |
| Gel Loading Dye, Purple (6X), no SDS | B7025S | 4 ml |

For the full list of restriction enzymes available, visit **www.neb.com**.

Products for End Modification

| PRODUCT | NEB # | SIZE |
|---|------------|-----------------------|
| Quick CIP | M0525S/L | 1,000/5,000 units |
| Shrimp Alkaline Phosphatase (Recombinant) | M0371S/L | 500/2,500 units |
| Antarctic Phosphatase | M0289S/L | 1,000/5,000 units |
| T4 DNA Polymerase | M0203S/L | 150/750 units |
| DNA Polymerase I, Large (Klenow) Fragment | M0210S/L/M | 200/1,000/1,000 units |
| Quick Blunting Kit | E1201S/L | 20/100 reactions |
| Mung Bean Nuclease | M0250S/L | 1,000/5,000 units |
| T4 Polynucleotide Kinase | M0201S/L | 500/2,500 units |
| Klenow Fragment (3 ^{\sim} \rightarrow 5 ^{\prime} exo ⁻) | M0212S/L/M | 200/1,000/1,000 units |
| β-Agarase I | M0392S/L | 100/500 units |

Products for Ligation

| PRODUCT | NEB # | SIZE |
|--------------------------------------|--------------|-----------------------|
| Blunt/TA Ligase Master Mix | M0367S/L | 50/250 reactions |
| Instant Sticky-End Ligase Master Mix | M0370S/L | 50/250 reactions |
| NEBridge Ligase Master Mix | M1100S/L | 50/250 reactions |
| ElectroLigase | M0369S | 50 reactions |
| T4 DNA Ligase | M0202S/L/T/M | 20,000/100,000 units |
| Salt-T4 DNA Ligase | M0467S/L | 20,000/100,000 units |
| Hi-T4 DNA Ligase | M2622S/L | 20,000/100,000 units |
| Quick Ligation Kit | M2200S/L | 30/150 reactions |
| T3 DNA Ligase | M0317S/L | 100,000/750,000 units |
| T7 DNA Ligase | M0318S/L | 100,000/750,000 units |
| Taq DNA Ligase | M0208S/L | 2,000/10,000 units |

Products for Transformation

| PRODUCT | NEB # | SIZE |
|---|----------------|---|
| dam-/dcm- Competent E. coli | C2925H/I | 20 x 0.05 ml/tube/ 6 x 0.2 ml ml/tube |
| NEB 5-alpha Competent <i>E. coli</i> (High Efficiency) | C2987H/I/P/R/U | 20 x 0.05 ml/tube/ 6 x 0.2 ml/tube/ 1 x 96 well plate/ 1 x 384 well plate/ 12 x 8 tube strips |
| NEB 5-alpha Competent <i>E. coli</i> (Subcloning Efficiency) | C2988J | 6 x 0.4 ml/tube |
| NEB 5-alpha F ^r / ^a Competent <i>E. coli</i> (High Efficiency) | C2992H/I | 20 x 0.05/6 x 0.2 ml |
| NEB 10-beta Competent <i>E. coli</i> (High Efficiency) | C3019H/I/P | 20 x 0.05 ml/tube/ 6 x 0.2 ml ml/tube 1 x 96 well plate |
| NEB 10-beta Electrocompetent E. coli | C3020K | 6 x 0.1 ml/tube |
| NEB Turbo Competent <i>E. coli</i> (High Efficiency) | C2984H/I | 20 x 0.05 ml/tube/ 6 x 0.2 ml/tube |
| NEB Stable Competent <i>E. coli</i> (High Efficiency) | C3040H/I | 20 x 0.05 ml/tube/ 6 x 0.2 ml/tube |
| NEB Cloning Competent E. coli Sampler | C1010S | 8 tubes |

For the full list of competent cells available, visit **www.neb.com**.

Products for Nucleic Acid Purification

| PRODUCT | NEB # | SIZE |
|---|------------|--------------------|
| Monarch Spin Plasmid Miniprep Kit | T1110S/L | 50/250 preps |
| Monarch Spin DNA Gel Extraction Kit | T1120S/L | 50/250 preps |
| Monarch Spin PCR & DNA Cleanup Kit (5 µg) | T1130S/L | 50/250 preps |
| Monarch Spin gDNA Extraction Kit | T3010S/L | 50/150 preps |
| Monarch Spin RNA Isolation Kit (Mini) | T2110S | 50 preps |
| Monarch Spin RNA Cleanup Kit (10 µg) | T2030S/L | 10/100 preps |
| Monarch Spin RNA Cleanup Kit (50 µg) | T2040S/L | 10/100 preps |
| Monarch Spin RNA Cleanup Kit (500 µg) | T2050S/L | 10/100 preps |
| Monarch HMW DNA Extraction Kit for Cells & Blood | T3050S/L | 5/50 preps |
| Monarch HMW DNA Extraction Kit for Tissue | T3060S/L | 5/50 preps |
| Monarch Mag Viral DNA/RNA Extraction Kit | T4010S/L/X | 100/600/1800 preps |

Columns and buffers also available separately.

Products for DNA Analysis

| PRODUCT | NEB # | SIZE |
|---|----------|---------------------|
| 1 kb DNA Ladder | N3232S/L | 200/1,000 gel lanes |
| TriDye 1 kb DNA Ladder | N3272S | 125 gel lanes |
| Quick-Load 1 kb DNA Ladder | N0468S/L | 125/375 gel lanes |
| 100 bp DNA Ladder | N3231S/L | 100/500 gel lanes |
| TriDye 100 bp DNA Ladder | N3271S | 125 gel lanes |
| Quick-Load 100 bp DNA Ladder | N0467S/L | 125/375 gel lanes |
| 1 kb Plus DNA Ladder | N3200S/L | 200/1,000 gel lanes |
| 1 kb Plus DNA Ladder for Safe Stains | N0559S | 50 μg/ml |
| TriDye 1 kb Plus DNA Ladder | N3270S | 250 gel lanes |
| Quick-Load 1 kb Plus DNA Ladder | N0469S | 250 gel lanes |
| Quick-Load Purple 1 kb Plus DNA Ladder | N0550S/L | 250/750 gel lanes |
| TriDye Ultra Low Range DNA Ladder | N0558S | 100 µg/ml |
| 50 bp DNA Ladder | N3236S/L | 200/1,000 gel lanes |
| Quick-Load Purple 50 bp DNA Ladder | N0556S | 250 gel lanes |
| Quick-Load 1 kb Extend DNA Ladder | N3239S | 125 gel lanes |
| Quick-Load Purple 1 kb DNA Ladder | N0552S/L | 125/375 gel lanes |
| Quick-Load Purple 100 bp DNA Ladder | N0551S/L | 125/375 gel lanes |
| Low Molecular Weight DNA Ladder | N3233S/L | 100/500 gel lanes |
| Quick-Load Purple Low Molecular Weight DNA Ladder | N0557S | 125 gel lanes |
| Fast DNA Ladder | N3238S | 200 gel lanes |
| PCR Marker | N3234S/L | 100/500 gel lanes |

Conventional and PFG markers are also available, visit www.neb.com/DNAMarkersandLadders.

Products for Seamless Cloning

| PRODUCT | NEB # | SIZE |
|---|--------------|----------------------|
| NEBuilder HiFi DNA Assembly Cloning Kit | E5520S | 10 reactions |
| NEBuilder HiFi DNA Assembly Master Mix | E2621S/L | 10/50 reactions |
| NEBuilder HiFi DNA Assembly Bundle for Large Fragments | E2623S | 20 reactions |
| Gibson Assembly Cloning Kit | E5510S | 10 reactions |
| Gibson Assembly Master Mix | E2611S/L | 10/50 reactions |
| NEBridge Golden Gate Assembly Kit (Bsal-HFv2) | E1601S/L | 20/100 reactions |
| NEBridge Golden Gate Assembly Kit (BsmBI-v2) | E1602S/L | 20/100 reactions |
| NEBridge Ligase Master Mix | M1100S/L | 50/250 reactions |
| Bbsl | R0539S/L | 300/1,500 units |
| BbsI-HF | R3539S/L | 300/1,500 units |
| Bsal-HFv2 | R3733S/L | 1,000/5,000 units |
| BsmBI-v2 | R0739S/L | 200/1,000 units |
| BspQI | R0712S/L | 500/2,500 units |
| BspQI-HF | R3712S/L | 500/2,500 units |
| BtgZI | R0703S/L | 100/500 units |
| Esp3I | R0734S/L | 300/1,500 units |
| PaqCI | R0745S/L | 200/1,000 units |
| Sapl | R0569S/L | 250/1,250 units |
| T4 DNA Polymerase | M0203S/L | 150/750 units |
| Taq DNA Ligase | M0208S/L | 2,000/10,000 units |
| T4 DNA Ligase | M0202S/L/T/M | 20,000/100,000 units |
| T5 Exonuclease | M0363S/L | 1,000/5,000 units |
| Exonuclease V (RecBCD) | M0345S/L | 1,000/5,000 units |
| USER Enzyme | M5505S/L | 50/250 units |
| Thermolabile USER Enzyme II | M5508S/L | 50/250 units |

Products for Recombinational Cloning

| PRODUCT | NEB # | SIZE |
|-----------------|------------|--------------|
| Cre Recombinase | M0298S/L/M | 50/250 units |



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