

Restriction Endonucleases

TECHNICAL GUIDE





Cut Smarter *with* Restriction Enzymes *from* NEB®

Looking to bring CONVENIENCE to your workflow?

Simplify Reaction Setup and Double Digestion with CutSmart® Buffer

Over 205 restriction enzymes are 100% active in a single buffer, CutSmart Buffer, making it significantly easier to set up your double digest reactions. Since CutSmart Buffer includes BSA, there are fewer tubes and pipetting steps to worry about. Additionally, many DNA modifying enzymes are 100% active in CutSmart Buffer, eliminating the need for subsequent purification.

For more information, visit www.NEBCutSmart.com

Speed up Digestions with Time-Saver™ Qualified Restriction Enzymes

190 of our restriction enzymes are able to digest DNA in 5–15 minutes, and can safely be used overnight with no loss of sample. For added convenience and flexibility, most of these are supplied with our new CutSmart Buffer.

For more information, visit www.neb.com/timesaver

Keep it Simple with our RE-Mix® Restriction Enzyme Master Mixes

RE-Mix Restriction Enzyme Master Mixes are pre-mixed solutions that contain enzyme, buffer, BSA and loading dye. Just add your DNA and water; it's that simple! RE-Mix master mixes are Time-Saver qualified so you can trust your reaction to digest to completion in 15 minutes, or leave it to digest overnight, with no degradation of your final product.

For more information, visit www.NEBREMix.com

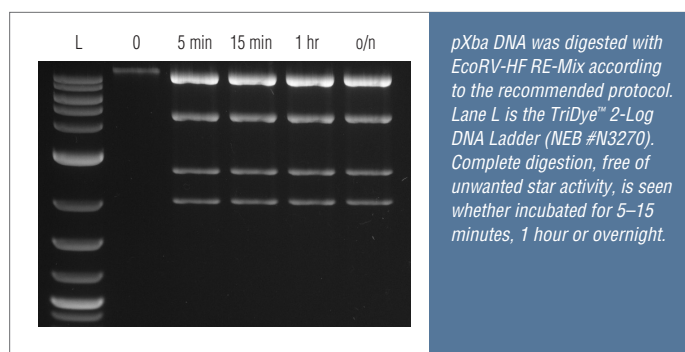
Bring Flexibility to your Workflow

NEB offers the largest selection of restriction enzymes commercially available. With an evergrowing list to choose from, currently at 280 enzymes – including traditional restriction enzymes, nicking endonucleases, homing endonucleases and methylation-sensitive enzymes for epigenetics studies – there is no need to look anywhere else.

Activity of DNA Modifying Enzymes in CutSmart Buffer

ENZYME	ACTIVITY IN CUTSMART	REQUIRED SUPPLEMENTS
Alkaline Phosphatase (CIP)	+++	
Antarctic Phosphatase	+++	Requires Zn ²⁺
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 Ca ²⁺
<i>E. coli</i> DNA Ligase	+++	Requires NAD
Endonuclease III (Nth), recombinant	+++	
Endonuclease VIII	+++	
Exonuclease III	+++	
GpC Methyltransferase (M. CviPI)	+	Requires DTT
McrBC	+++	
Micrococcal Nuclease	+++	
Nuclease BAL-31	+++	
phi29 DNA Polymerase	+++	
RecJ	+++	
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
T7 DNA Ligase	+++	Requires ATP + PEG
T7 DNA Polymerase (unmodified)	+++	
T7 Exonuclease	+++	
USER Enzyme, recombinant	+++	

+++ full functional activity ++ 50–100% functional activity + 0–50% functional activity



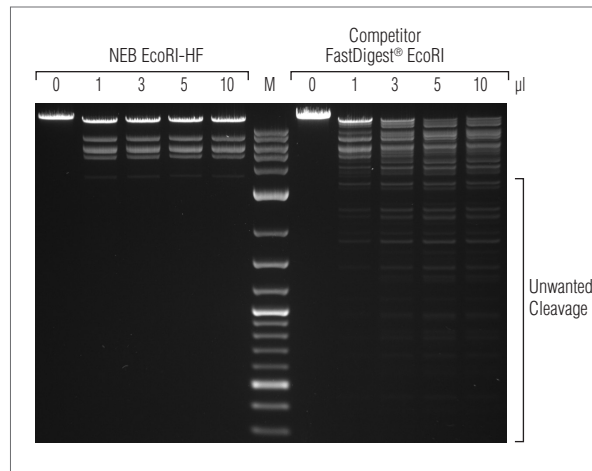


Looking to optimize PERFORMANCE in your reaction?

Choose a High-Fidelity (HF®) Restriction Enzyme

As part of our ongoing commitment to the advancement and improvement of enzymes for the cloning and manipulation of DNA, NEB has developed a line of High-Fidelity (HF) restriction enzymes. These engineered enzymes have the same specificity as the native enzyme, with the added benefit of reduced star activity, rapid digestion (5-15 minutes), and 100% activity in CutSmart Buffer. Enjoy the improved performance of NEB's engineered enzymes at the same price as the native enzymes!

For more information, visit www.neb.com/HF



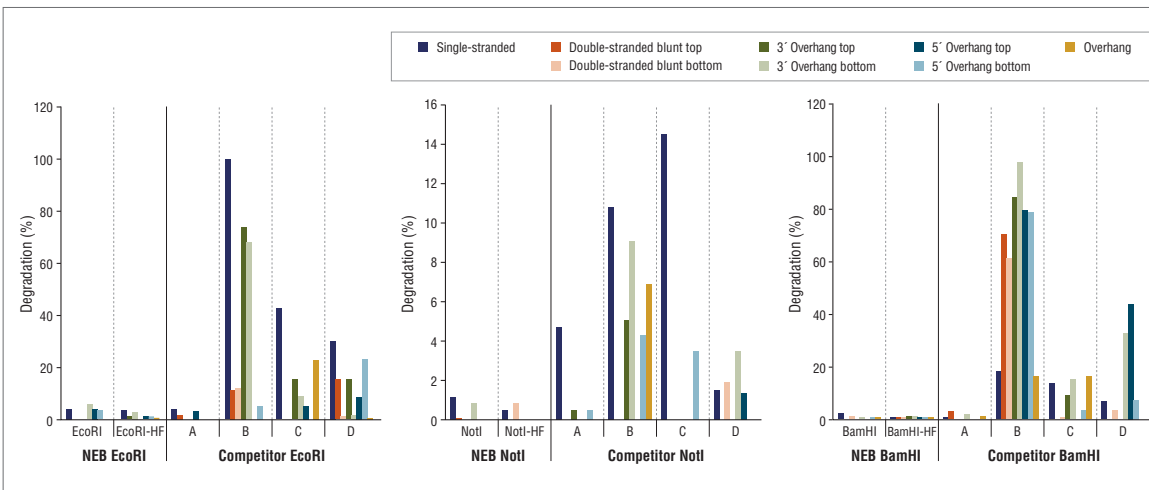
EcoRI-HF (NEB #R3101) shows no star activity in overnight digests, even when used at higher concentrations. 50 µl reactions were set up using 1 µg of Lambda DNA, the indicated amount of enzyme and the recommended reaction buffer. Reactions were incubated overnight at 37°C. Marker M is the 1 kb DNA Ladder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

Benefit from Industry-leading Quality

NEB's reputation as a leader in enzyme technologies stems from the quality and reliability of our restriction enzymes. All of our restriction enzymes undergo stringent quality control testing, ensuring the highest levels of purity and lot-to-lot consistency.

HIGHLIGHTS

- Industry-leading product quality
- State-of-the-art production and purification
- Over 40 years of experience
- Stringent quality control testing
- Lot-to-lot consistency
- ISO 9001- and 13485-certified



Restriction Enzyme Competitor Study: Nuclease Contamination

EcoRI, NotI, and BamHI from multiple suppliers were tested in reactions containing a fluorescent labeled single stranded, double stranded blunt, 3' overhang or 5' overhang containing oligonucleotides. The percent degradation is determined by capillary electrophoresis and peak analysis. The resolution is at the single nucleotide level.

Visit NEBCutSmart.com for information on the smarter choice of restriction enzymes.



High Fidelity (HF) Enzymes

High-Fidelity (HF) restriction enzymes are engineered enzymes that have the same specificity as the native enzymes, are all active in CutSmart Buffer and have reduced star activity. Star activity, or off-target cleavage, is an intrinsic property of restriction enzymes. Most restriction enzymes will not exhibit star activity under recommended reaction conditions. However, for enzymes that have reported star activity, extra caution must be taken to set up reactions according to the recommended conditions to avoid unwanted cleavage. HF enzymes should be used in these cases.

In addition to reduced star activity, HF enzymes work optimally in CutSmart Buffer, which has the highest level of enzyme compatibility and will simplify double digest reactions. They are all Time-Saver qualified and digest substrate DNA in 5–15 minutes and are flexible enough to digest overnight. HF enzymes are supplied with our **purple gel loading dye**, which sharpens bands and eliminates UV shadow. Lastly, they are available at the same price as the native enzymes.

The following table indicates the number of units of HF enzyme that can be used compared to the native enzyme before any significant star activity is detected. The HF Factor refers to the X-fold increase in fidelity that is achieved by choosing an HF enzyme. This data clearly illustrates the flexibility that is offered by using an HF restriction enzyme.

PRODUCT NAME	PRODUCT NUMBER	BUFFER [†]	MAXIMUM UNITS WITH NO STAR ACTIVITY*	HF FACTOR
AgeI-HF	#R3552	CutSmart	≥ 250	≥ 8
AgeI	#R0552	1.1	32	
BamHI-HF	#R3136	CutSmart	≥ 4,000	≥ 125
BamHI	#R0136	3.1	32	
BmtI-HF	#R3658	CutSmart	1,000,000	31,250
BmtI	#R0658	3.1	32	
Bsal-HF	#R3535	CutSmart	≥ 8,000	≥ 250
Bsal	#R0535	CutSmart	32	
BsrGI-HF	#R3575	CutSmart	≥ 1,000	≥ 62
BsrGI	#R0575	2.1	16	
BstEII-HF	#R3162	CutSmart	≥ 2,000	≥ 125
BstEII	#R0162	3.1	16	
DrallI-HF	#R3510	CutSmart	≥ 2,000	≥ 1,000
DrallI**	N/A	3.1	2	
EagI-HF	#R3505	CutSmart	500	2
EagI	#R0505	3.1	250	
EcoRI-HF	#R3101	CutSmart	16,000	64
EcoRI	#R0101	U	250	
EcoRV-HF	#R3195	CutSmart	≥ 64,000	≥ 64
EcoRV	#R0195	3.1	1,000	
HindIII-HF	#R3104	CutSmart	≥ 500,000	≥ 2,000
HindIII	#R0104	2.1	250	
KpnI-HF	#R3142	CutSmart	≥ 1,000,000	≥ 62,500
KpnI	#R0142	1.1	16	
MfeI-HF	#R3589	CutSmart	≥ 500	≥ 16
MfeI	#R0589	CutSmart	32	
MluI-HF	#R3198	CutSmart	≥ 4,000	2
MluI	#R0198	3.1	≥ 2,000	
NcoI-HF	#R3193	CutSmart	≥ 64,000	≥ 530
NcoI	#R0193	3.1	120	

PRODUCT NAME	PRODUCT NUMBER	BUFFER [†]	MAXIMUM UNITS WITH NO STAR ACTIVITY*	HF FACTOR
NheI-HF	#R3131	CutSmart	≥ 32,000	≥ 266
NheI	#R0131	2.1	120	
NotI-HF	#R3189	CutSmart	≥ 64,000	≥ 16
NotI	#R0189	3.1	4,000	
NruI-HF	#R3192	CutSmart	≥ 32,000	64
NruI	#R0192	3.1	≥ 500	
NsiI-HF	#R3127	CutSmart	≥ 8,000	2
NsiI	#R0127	3.1	≥ 4,000	
PstI-HF	#R3140	CutSmart	4,000	33
PstI	#R0140	3.1	120	
PvuI-HF	#R3150	CutSmart	≥ 16,000	≥ 32
PvuI	#R0150	3.1	500	
PvuII-HF	#R3151	CutSmart	500	32
PvuII	#R0151	2.1	16	
SacI-HF	#R3156	CutSmart	≥ 32,000	≥ 266
SacI	#R0156	1.1	120	
Sall-HF	#R3138	CutSmart	≥ 32,000	≥ 8,000
Sall	#R0138	3.1	4	
SbfI-HF	#R3642	CutSmart	250	32
SbfI	#R0642	CutSmart	8	
Scal-HF	#R3122	CutSmart	250	62
Scal**	#R0122	3.1	4	
SpeI-HF	#R3133	CutSmart	≥ 8,000	≥ 16
SpeI	#R0133	CutSmart	500	
SphI-HF	#R3182	CutSmart	8,000	250
SphI	#R0182	2.1	32	
SspI-HF	#R3132	CutSmart	500	16
SspI	#R0132	U	32	
StyI-HF	#R3500	CutSmart	4,000	125
StyI	#R0500	3.1	32	

TOOLS & RESOURCES

Visit NEBRestrictionEnzymes.com to find:

- The full list of HF restriction enzymes available
- Online tutorials on how to avoid star activity and setting up digests using the Time-Saver protocol



WHAT IS A HIGH-FIDELITY ENZYME?

[†] Wild type enzymes were tested in supplied buffer for comparisons.

* Wei, H. et al (2008) Nucleic Acids Research 36, e50.

** No longer available.



Avoiding Star Activity

Tips for preventing unwanted cleavage in restriction enzyme digests

Under non-standard reaction conditions, some restriction enzymes are capable of cleaving sequences which are similar but not identical to their defined recognition sequence. This altered specificity has been termed “star activity”. It has been suggested that star activity is a general property of restriction endonucleases (1) and that any restriction endonuclease will cleave noncanonical sites under certain extreme conditions, some of which are listed below. Although the propensity for star activity varies, the vast majority of enzymes from New England Biolabs will not exhibit star activity when used under recommended conditions in their supplied NEBuffers. If an enzyme has been reported to exhibit star activity, it will be indicated in the product entry found in the catalog, on the supplied card and on our website.

CONDITIONS THAT CONTRIBUTE TO STAR ACTIVITY	STEPS THAT CAN BE TAKEN TO INHIBIT STAR ACTIVITY
High glycerol concentration (> 5% v/v)	Restriction enzymes are stored in 50% glycerol, therefore the amount of enzyme added should not exceed 10% of the total reaction volume. Use the standard 50 µl reaction volume to reduce evaporation during incubation.
High concentration of enzyme/µg of DNA ratio (varies with each enzyme, usually 100 units/µg)	Use the fewest units possible to achieve digestion. This avoids overdigestion and reduces the final glycerol concentration in the reaction.
Non-optimal buffer	Whenever possible, set up reactions in the recommended buffer. Buffers with differing ionic strengths and pHs may contribute to star activity.
Prolonged reaction time	Use the minimum reaction time required for complete digestion. Prolonged incubation may result in increased star activity, as well as evaporation.
Presence of organic solvents [DMSO, ethanol (4), ethylene glycol, dimethylacetamide, dimethylformamide, sulphalane (5)]	Make sure the reaction is free of any organic solvents, such as alcohols, that might be present in the DNA preparation.
Substitution of Mg ²⁺ with other divalent cations (Mn ²⁺ , Cu ²⁺ , Co ²⁺ , Zn ²⁺)	Use Mg ²⁺ as the divalent cation. Other divalent cations may not fit correctly into the active site of the restriction enzyme, possibly interfering with proper recognition.

Note: The relative significance of each of these altered conditions will vary from enzyme to enzyme.

New England Biolabs recommends setting up restriction enzyme digests in a 50 µl reaction volume. However, different methods may require smaller reaction volumes. When performing restriction enzyme digests in smaller reaction volumes, extra care must be taken to follow the steps listed above to avoid star activity. Alternatively, using our line of **High Fidelity (HF) restriction enzymes** will allow greater flexibility in reaction setup. Please visit www.neb.com/HF frequently to learn about new additions to the HF restriction enzyme product line.

References:

- Nasri, M. and Thomas, D. (1986) *Nucleic Acids Res.* 14, 811.
- Barany, F. (1988) *Gene*, 68, 149.
- Bitinaite, J. and Schildkraut, I. (2002) *Proc. Natl. Acad. Sci. USA*, 99, 1164–1169.
- Nasri, M. and Thomas, D. (1987) *Nucleic Acids Res.* 15, 7677.
- Tikhonchenko, T.I., *et al.* (1978) *Gene*, 4, 195–212.

TOOLS & RESOURCES

Visit NEBRestrictionEnzymes.com to find:

- Video tutorials on how to avoid star activity, and for setting up restriction enzyme digests
- The full list of HF enzymes available
- Troubleshooting guides



RESTRICTION ENZYME TYPES

- Type I enzymes are multisubunit proteins that cut DNA randomly at a distance from their recognition sequence.
- Type II enzymes cut DNA at defined positions close to or within their recognition sequence and are commonly used in the laboratory. There are over ten subtypes with different types of recognition sites, cleavage sites and cofactor requirements.
- The most common Type II enzymes cleave within their recognition site (e.g., BamHI, EcoRI); sites can be symmetric or asymmetric.
- Type IIS enzymes cleave outside their recognition sequence (e.g., FokI, AlwI) and are invaluable for emerging technologies in the biotechnology industry.
- Type IIM enzymes recognize methylated targets (e.g., DpnI).
- Type III enzymes are large, combination restriction-and-modification enzymes that cleave outside their recognition sequences and require two sequences in opposite orientations to cleave one DNA molecule.
- Type IV enzymes recognize modified DNA (methylated, hydroxymethylated, etc.). They require two sites and cleave non-specifically.
- Isoschizomers are restriction enzymes that recognize the same sequence as the prototype.
- Neoschizomers are isoschizomers with different cleavage sites.



Learn more about restriction enzyme types in our online tutorials.



Time-Saver Qualified Restriction Enzymes

Whether you are quickly screening large numbers of clones or setting up overnight digests, you will benefit from the high quality of our enzymes. Typically, a restriction digest involves the incubation of 1 µl of enzyme with 1 µg of purified DNA in a final volume of 50 µl for 1 hour. However, to speed up the screening process, choose one of NEB's enzymes that are Time-Saver qualified. **190 of our enzymes will digest 1 µg of substrate DNA in 5-15 minutes using 1 µl of enzyme under recommended reaction conditions, and can also be used safely in overnight digestions.** Unlike other suppliers, there is no special formulation, change in concentration or need to buy more expensive, new lines of enzymes to achieve digestion in 5-15 minutes. Nor do you have to worry if you incubate too long.

In an effort to provide you with as much information as possible, NEB has tested all of its enzymes on unit assay substrate, as well as plasmid substrate and PCR fragments. We recommend that this data be used as a guide, as it is not definitive for all plasmids. Restriction enzymes can often show site preference, presumably determined by the sequence flanking the recognition site. In addition, supercoiled DNA may have varying rates of cleavage. For more information, visit www.neb.com/TimeSaver. Note that there are some enzymes indicated below that can cut in 5-15 minutes, but cannot be incubated overnight. These are not Time-Saver qualified.

Since all of our enzymes are rigorously tested for nuclease contamination, you can also safely set up digests for long periods of time without sample degradation. Only NEB Time-Saver qualified enzymes offer power and flexibility – the power to digest in 5-15 minutes and the flexibility to withstand overnight digestions with no loss of substrate.

Chart Legend

- digests in 5 minutes
- digests in 15 minutes
- ▲ not completely digested in 15 minutes
- NT not tested

ENZYME	SUBSTRATE		
	UNIT ASSAY	PLASMID	PCR
AatII	■	▲	●
AccI	■	▲	▲
Acc65I	●	▲	●
AcII	●	●	●
AcII	●	■	▲
AcuI	■	▲	▲
AflII	●	●	●
AgeI-HF	●	●	●
AgeI-HF RE-Mix	●	●	■
AhdI	●	●	■
AluI	●	▲	●
AlwNI	●	●	▲
ApaI	●	●	●
ApaLI	●	●	▲
ApeKI	●	■	▲
ApoI	●	●	●
Ascl	●	●	NT
Ascl RE-Mix	●	●	NT
Asel	●	●	NT
AvaI	●	▲	▲
Avall	●	●	●
AvrII	●	▲	NT
BaeI	■	●	▲

ENZYME	SUBSTRATE		
	UNIT ASSAY	PLASMID	PCR
BaeGI	●	▲	▲
BamHI	●	●	▲
BamHI-HF	●	●	●
BbsI	■	▲	▲
BbvI	●	▲	▲
BccI	■	▲	▲
BceAI	■	■	▲
BciVI	●	■	▲
BclI	●	▲	▲
BcoDI	●	●	▲
BfuAI	●	●	▲
BfuCI	■	▲	●
BglII	●	●	▲
BgIII	●	■	▲
Bipl	●	●	●
BmgBI	●	●	▲
Bmrl	■	▲	■
BmtI-HF	●	●	▲
BpuEI	●	●	▲
BsaI	●	●	▲
BsaI-HF	●	●	▲
BsaAI	●	●	■
BsaHI	■	■	●

ENZYME	SUBSTRATE		
	UNIT ASSAY	PLASMID	PCR
BsaWI	■	▲	▲
BsaXI	●	▲	▲
BseRI	●	●	■
BsgI	●	●	▲
BsiEI	●	▲	▲
BsiWI	●	●	▲
BsII	●	■	■
BsmI	●	●	▲
BsmAI	●	▲	●
BsmBI	■	▲	▲
BsmFI	●	●	▲
BsoBI	●	■	●
Bsp1286I	●	●	▲
BspCNI	■	▲	▲
BspEI	●	▲	▲
BspHI	■	●	●
BspQI	●	●	▲
BsrI	●	■	▲
BsrBI	●	■	▲
BsrDI	●	■	▲
BsrGI	■	▲	▲
BsrGI-HF	●	●	▲
BssHII	●	▲	▲
BssKI	■	▲	●
BssS ⁺ I	●	▲	▲
BstBI	●	●	▲
BstEII	●	●	▲
BstEII-HF	●	●	●
BstEII-HF RE-Mix	●	●	■
BstNI	●	●	▲
BstUI	●	●	▲
BstXI	●	●	▲
BstYI	■	●	▲
BstZ17I	●	▲	▲
Bsu36I	■	▲	■
Bts ⁺ I	●	●	■
BtsCI	●	■	▲
Cac8I	■	▲	▲
Clal	●	●	▲
CspCI	●	●	▲
CviAII	■	●	●
CviQI	●	●	●
DdeI	●	■	■



ENZYME	SUBSTRATE		
	UNIT ASSAY	PLASMID	PCR
DpnI	●	●	▲
DpnII	■	▲	●
DraI	●	●	■
DraIII-HF	●	●	▲
DrdI	■	●	●
EagI	●	▲	▲
EagI-HF	●	●	▲
EarI	■	■	▲
Eco53KI	●	●	■
EcoNI	●	■	●
EcoO109I	●	▲	▲
EcoP15I	■	▲	▲
EcoRI	●	●	▲
EcoRI-HF	●	●	●
EcoRI-HF RE-Mix	●	●	▲
EcoRV	●	●	▲
EcoRV-HF	●	●	▲
EcoRV-HF RE-Mix	●	●	▲
Fnu4HI	●	■	■
FokI	●	●	●
FseI	●	●	▲
FspI	■	▲	■
HaeII	■	▲	▲
HaeIII	●	●	●
HgaI	■	▲	▲
HhaI	●	■	▲
HincII	■	▲	●
HindIII-HF	●	●	●
HinfI	●	●	●
HinP1I	●	▲	●
HpaII	●	●	▲
HphI	●	▲	▲
Hpy166II	●	●	●
HpyAV	●	●	NT
HpyCH4IV	●	●	●
HpyCH4V	●	●	●
KpnI-HF	●	●	●
KpnI-HF RE-Mix	●	●	▲
MboI	●	▲	●
MboII	●	●	●
MfeI	●	●	●
MfeI-HF	●	●	●
MfeI-HF RE-Mix	●	●	●
MluI	●	●	●
MluI-HF	●	●	▲
MluCI	●	●	▲
MlyI	●	▲	●
MmeI	●	●	▲
MnII	●	●	■
MseI	■	■	●

ENZYME	SUBSTRATE		
	UNIT ASSAY	PLASMID	PCR
MsiI	●	●	●
MspI	●	●	●
MspA1I	●	●	●
MwoI	■	▲	▲
NciI	●	●	●
NcoI	●	■	▲
NcoI-HF	●	●	●
NcoI-HF RE-Mix	●	●	●
NdeI	●	●	▲
NgoMIV	■	●	▲
NheI	●	■	▲
NheI-HF	●	●	■
NheI-HF RE-Mix	●	●	▲
NlaIII	■	▲	■
NmeAIII	●	▲	▲
NotI	●	●	▲
NotI-HF	●	●	●
NotI-HF RE-Mix	●	●	▲
NruI	●	■	▲
NruI-HF	●	■	▲
NsiI	●	●	●
NsiI-HF	●	●	■
NspI	●	■	▲
PacI	●	●	●
PacI-RE-Mix	●	●	NT
PaeR7I	●	▲	▲
PfiI	●	■	▲
PfiMI	●	▲	▲
PmeI	●	■	NT
PmlI	●	▲	▲
PpuMI	●	▲	▲
PshAI	■	■	■
PstI	●	●	●
PstI-HF	●	●	●
PvuI	●	▲	●
PvuI-HF	●	●	●
PvuII	●	●	▲
PvuII-HF	●	●	▲
RsaI	●	●	●
SacI	●	●	▲
SacI-HF	●	●	●
SacII	●	▲	▲
SalI	●	■	▲
SalI-HF	●	●	▲
SalI-HF RE-Mix	●	●	▲
SapI	■	▲	▲
SbfI	●	●	▲
SbfI-HF	●	●	▲
Scal-HF	●	●	▲
Scal-HF RE-Mix	●	●	▲

ENZYME	SUBSTRATE		
	UNIT ASSAY	PLASMID	PCR
SfiI	●	▲	▲
SfoI	●	●	●
SmaI	●	■	■
SpeI	●	●	●
SpeI RE-Mix	●	●	●
SphI	●	●	▲
SspI	●	●	▲
SspI-HF	●	●	▲
StuI	■	▲	▲
StyI	■	▲	▲
StyI-HF	●	●	▲
StyD4I	■	▲	▲
Swal	■	▲	▲
TaqI	●	●	▲
TfiI	■	●	▲
TseI	■	▲	▲
TspMI	●	■	▲
TspRI	●	■	▲
Tth111I	■	■	▲
XbaI	●	●	▲
XbaI-RE-Mix	●	●	■
XhoI	●	●	▲
XhoI RE-Mix	●	●	NT
XmaI	■	▲	■
XmnI	●	●	▲

TOOLS & RESOURCES

Visit www.neb.com/TimeSaver to find:

- The full list of Time-Saver qualified restriction enzymes available
- Video tutorials on how Time-Saver qualified enzymes speed up restriction enzyme digests



Optimizing Restriction Enzyme Reactions

There are several key factors to consider when setting up a restriction enzyme digest. Using the proper amounts of DNA, enzyme and buffer components in the correct reaction volume will allow you to achieve optimal digestion. By definition, 1 unit of restriction enzyme will completely digest 1 μg of substrate DNA in a 50 μl reaction in 60 minutes. This enzyme:DNA:reaction volume ratio can be used as a guide when designing reactions. However, most researchers follow the “typical” reaction conditions listed, where a 5–10 fold overdigestion is recommended to overcome variability in DNA source, quantity and purity. The Time-Saver protocol can be used for enzymes that are Time-Saver qualified and will digest DNA in 5–15 minutes (see page 6–7 for the full list). For additional convenience, a RE-Mix Restriction Enzyme Master Mix can also be used. NEB offers the following tips to help you to achieve maximal success in your restriction enzyme reactions.

Standard Protocol

Restriction Enzyme	1 μl (or 10 units)*
DNA	1 μg
10X NEBuffer	5 μl (1X)
Total Reaction Volume	50 μl
Incubation Temperature	Enzyme Dependent
Incubation Time	60 minutes

*Sufficient to digest all types of DNAs.

Time-Saver Protocol:

Restriction Enzyme	1 μl
DNA	1 μg
10X NEBuffer	5 μl (1X)
Total Reaction Volume	50 μl
Incubation Temperature	Enzyme Dependent
Incubation Time	5–15 minutes*

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

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 for genomic DNA
- NEB has introduced a line of High-Fidelity (HF) enzymes that provide added flexibility to reaction setup.
- If using a RE-Mix restriction enzyme master mix, see page 9 for protocol.

DNA

- Should be free of contaminants such as phenol, chloroform, alcohol, EDTA, detergents, nucleases or excessive salts
- Methylation of DNA can inhibit digestion with certain enzymes

Buffer

- Use at a 1X concentration
- BSA is included in NEBuffer 1.1, 2.1, 3.1 and CutSmart Buffer. No additional BSA is needed.

Reaction Volume

- A 50 μl reaction volume is recommended for digestion of 1 μg of substrate
- 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. The following guidelines can be used for techniques that require smaller reaction volumes.

Alternative Volumes for Restriction Digests

	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 Standard Protocol is 1 hour. Incubation for Time-Saver Protocol is 5–15 minutes.
- With many enzymes, it is possible to use fewer units and digest for up to 16 hours. For more information, visit www.neb.com.

Stopping a Reaction

If no further manipulation of DNA is required:

- Terminate with a stop solution (10 μ l per 50 μ l rxn) [50% glycerol, 50 mM EDTA (pH 8.0), and 0.05% bromophenol blue] (e.g., NEB #B7021) or Gel Loading Dye, Purple (6X) (NEB #B7024).

When further manipulation of DNA is required:

- Heat inactivation can be used (buffer chart indicates if the enzyme can be heat inactivated)
- If enzyme cannot be heat inactivated, remove by using a spin column or phenol/chloroform extraction

Storage

- Storage at -20°C is recommended for most restriction enzymes. For a few enzymes, storage at -70°C is recommended for periods longer than 30 days. Visit www.neb.com for storage information.
- 10X NEBuffers should also be stored at -20°C

Stability

- All enzymes are assayed for activity every 3–6 months. The expiration date is found on the label.
- Exposure to temperatures above -20°C should be minimized whenever possible

Control Reactions

For difficulty cleaving DNA substrate, we recommend the following controls:

- Control DNA (DNA with multiple known sites for the enzyme) with restriction enzyme to test enzyme viability
- If the control DNA is cleaved and the experimental DNA resists cleavage, the two DNAs can be mixed to determine if an inhibitor is present in the experimental sample. If an inhibitor (often salt, EDTA or phenol) is present, the control DNA will not cut after mixing.

TOOLS & RESOURCES

Visit NEBCutSmart.com to find:

- Video tutorials on setting up restriction enzyme reactions from NEB scientists



Star Activity

- Can occur when enzyme is used under sub-optimal conditions
- Star activity can be reduced by using a High-Fidelity (HF) enzyme, by reducing incubation time, by using a Time-Saver enzyme or by increasing reaction volume

Optimizing Restriction Enzyme Reactions Using RE-Mix Master Mixes

- RE-Mix master mixes include enzyme, buffer, BSA and loading dye. All that is required is the addition of DNA and water. For the full list of RE-Mix master mixes, visit NEBREmix.com.

Many of the optimization tips for restriction enzymes apply to RE-Mix. Additional tips include:

- RE-Mix Master Mixes should be used at 1X concentration
- A 20 μ l reaction volume is recommended for digestion with a RE-Mix Master Mix

- The recommended incubation time with a RE-Mix Master Mix is 15 minutes
- The RE-Master Mix includes a density agent and dye, and does not require addition of stop solution
- RE-Mix Master Mixes should be stored at -20°C

RE-Mix Protocol

DNA	X μ l (up to 1 μ g)
dH ₂ O	18 μ l–X
10X RE-Mix	2 μ l
Reaction Volume	20 μ l
Incubation Temperature	37 $^{\circ}\text{C}$
Incubation Time	15 minutes



Troubleshooting Guide

PROBLEM	CAUSE	SOLUTION
Few or no transformants	Restriction enzyme(s) didn't cleave completely	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
		When digesting a PCR fragment, make sure to have at least 6 nucleotides between the recognition site and the end of the DNA molecule
The digested DNA ran as a smear on an agarose gel	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
	Nuclease contamination	Use fresh, clean running buffer and a fresh agarose gel
		Clean up the DNA
Incomplete restriction enzyme digestion	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 3.1) may be salt sensitive, so clean up the DNA prior to digestion
		DNA purification procedures that use spin columns can result in high salt levels, which inhibit enzyme activity. 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
	Using the wrong buffer	Use the recommended buffer supplied with the restriction enzyme
	Too few units of enzyme used	Use at least 3–5 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.	
Incomplete restriction enzyme digestion	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
	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. Mini prep DNA is particularly susceptible to contaminants. Clean DNA with a spin column, resin or drop dialysis, or increase volume to dilute contaminant
Extra bands in the gel	If larger bands than expected are seen in the gel, this may indicate binding of the enzyme(s) to the substrate	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
	Star activity	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.
	Partial restriction enzyme digest	Try using a High-Fidelity (HF) restriction enzyme. HF enzymes have been engineered for reduced star activity.
		Enzymes that have low activity in salt-containing buffers (e.g., NEBuffer 3.1) may be salt sensitive. Make sure to clean up the DNA prior to digestion.
		DNA purification procedures that use spin columns can result in high salt levels, which inhibit enzyme activity. To prevent this, DNA solution should be no more than 25% of total reaction volume
		Clean-up the PCR fragment prior to restriction digest
	Use the recommended buffer supplied with the restriction enzyme	
	Use at least 3–5 units of enzyme per µg of DNA and digest the DNA for 1–2 hours	

FAQS

Q. Do restriction enzymes cleave single-stranded DNA?

A. Although some restriction enzymes have been reported to cleave ssDNA, it is unclear whether cleavage occurs on a ssDNA molecule or on two ssDNA molecules which transiently anneal at a region of partial homology (1–3). For this reason, we hesitate to make unreserved claims about a restriction enzyme's ability to cut ssDNA.

Q. How stable are restriction enzymes?

A. All restriction enzymes from NEB are assayed for activity every 3–6 months. Most are very stable when stored at -20°C in the recommended storage buffer. Exposure to temperatures above -20°C should be minimized whenever possible.

Q. Is extended digestion (incubation times > 1 hour) recommended?

A. The unit definition of our restriction enzymes is based on a 1 hour incubation. Incubation time may be shortened if additional units of restriction enzyme are added to the reaction or if a Time-Saver qualified restriction enzyme is used (5–15 minutes). Conversely, longer incubation times are often used to allow a reaction to proceed to completion with fewer units of enzyme. This is contingent on how long a particular enzyme can survive (maintain activity) in a reaction. Additional information on extended digestion can be found at www.neb.com.

References

- Blakesley, R.W., Wells, R.D. (1975) *Nature* 257, 421–422.
- Blakesley, R.W., et al. (1977) *J. Biol. Chem.* 252, 7300–7306.
- Yoo, O.J., Agarwal, K.L. (1980) *J. Biol. Chem.* 255, 10559–10562.



Double Digestion

Digesting a DNA substrate with two restriction enzymes simultaneously (double digestion) is a common timesaving procedure. Over 205 restriction enzymes are 100% active in CutSmart Buffer, making double digestion simple. If you are using an enzyme that is not supplied with CutSmart Buffer, the Performance Chart for Restriction Enzymes rates the percentage activity of each restriction endonuclease in the four standard NEBuffers.

Setting up a Double Digest

- Double digests with CutSmart restriction enzymes can be set up in CutSmart Buffer. Otherwise, choose an NEBuffer that results in the most activity for both enzymes. If star activity is a concern, consider using one of our High-Fidelity (HF) enzymes.
- Set up reaction according to recommended protocol. The final concentration of glycerol in any reaction should be less than 5% to minimize the possibility of star activity. For example, in a 50 μ l reaction, the total amount of enzyme added should not exceed 5 μ l.
- If two different incubation temperatures are necessary, choose the optimal reaction buffer and set up reaction accordingly. Add the first enzyme and incubate at the desired temperature. Then, heat inactivate the first enzyme, add the second enzyme and incubate at the recommended temperature.
- Depending on an enzyme's activity rating in a non-optimal NEBuffer, the number of units or incubation time may be adjusted to compensate for the slower rate of cleavage.

Setting up a Double Digest with a unique buffer

- NEB currently supplies three enzymes with unique buffers: EcoRI, SspI and DpnII. In most cases, DpnII requires a sequential digest. Note that EcoRI has an HF version which is supplied with CutSmart Buffer.

Setting up a Sequential Digest

- If there is no buffer in which the two enzymes both exhibit > 50% activity, a sequential digest can be performed.

- Set up a reaction using the restriction endonuclease that has the lowest salt concentration in its recommended buffer and incubate to completion.
- Adjust the salt concentration of the reaction (using a small volume of a concentrated salt solution) to approximate the reaction conditions of the second restriction endonuclease.
- Add the second enzyme and incubate to complete the second reaction.
- Alternatively, a spin column can be used to isolate the DNA prior to the second reaction.

Setting up a Double Digestion with RE-Mix Master Mixes

- RE-Mix master mixes can also be used in double digest reactions.

Double Digest Protocol using two RE-Mix Enzymes:

DNA	X μ l (up to 1 μ g)
dH ₂ O	36 μ l-X
RE-Mix 1	2 μ l
RE-Mix 2	2 μ l
Total Volume	40 μ l
Incubation Temperature	37°C
Incubation Time	15 minutes

Double Digest Protocol using One RE-Mix and One Standard Restriction Enzyme.*

DNA	X μ l (up to 1 μ g)
dH ₂ O	17 μ l-X
RE-Mix	2 μ l
Standard Enzyme	1 μ l
Total Volume	20 μ l
Incubation Temperature	37°C
Incubation Time	15 minutes (Time-Saver Enzymes)
	1 Hour (Standard Enzymes)

* Use only with standard restriction enzymes with 37°C incubation temperature.

TOOLS & RESOURCES

Visit www.neb.com/nebtools for:

- Help choosing double digest conditions using NEB's **Double Digest Finder** or **NEBCloner**®



TIPS FOR SETTING UP DOUBLE DIGESTS



DNA Methylation & Restriction Digests

DNA methyltransferases (MTases) that transfer a methyl group from S-adenosylmethionine to either adenine or cytosine residues are found in a wide variety of prokaryotes and eukaryotes. Methylation should be considered when digesting DNA with restriction endonucleases because cleavage can be blocked or impaired when a particular base in the recognition site is methylated.

Prokaryotic Methylation

In prokaryotes, MTases have most often been identified as elements of restriction/modification systems that act to protect host DNA from cleavage by the corresponding restriction endonuclease. Most laboratory strains of *E. coli* contain three site-specific DNA methyltransferases.

- Dam methyltransferases—methylation at the N⁶ position of the adenine in the sequence GATC (1,2).
- Dcm methyltransferases—methylation at the C⁵ position of cytosine in the sequences CCAGG and CCTGG (1,3).
- EcoKI methylase—methylation of adenine in the sequences AAC(N⁶A)GTGC and GCAC(N⁶A)GTT.

Some or all of the sites for a restriction endonuclease may be resistant to cleavage when isolated from strains expressing the Dam or Dcm MTase if the methylase recognition site overlaps the endonuclease recognition site. For example, plasmid DNA isolated from *dam*⁺ *E. coli* is completely resistant to cleavage by MboI, which cleaves at GATC sites.

Not all DNA isolated from *E. coli* is methylated to the same extent. While pBR322 DNA is fully modified (and is therefore completely resistant to MboI digestion), only about 50% of λ DNA Dam sites are methylated, presumably because the methylase does not have the opportunity to methylate the DNA fully before it is packaged into the phage head. As a result, enzymes blocked by Dam or Dcm modification will yield partial digestion patterns with λ DNA.

Restriction sites that are blocked by Dam or Dcm methylation can be un-methylated by cloning your DNA into a *dam*⁻, *dcm*⁻ strain of *E. coli*, such as *dam*⁻/*dcm*⁻ Competent *E. coli* (NEB #C2925).

Restriction sites can also be blocked if an overlapping site is present. In this case, part of the Dam or Dcm sequence is generated by the restriction enzyme sequence, followed by the flanking sequence. This situation should also be considered when designing restriction enzyme digests.

Eukaryotic Methylation

CpG MTases, found in higher eukaryotes (e.g., Dnmt1), transfer a methyl group to the C⁵ position of cytosine residues. Patterns of CpG methylation are heritable, tissue specific and correlate with gene expression. Consequently, CpG methylation has been postulated to play a role in differentiation and gene expression (4).

Note: The effects of CpG methylation are mainly a concern when digesting eukaryotic genomic DNA. CpG methylation patterns are not retained once the DNA is cloned into a bacterial host.

Methylation Sensitivity

The table below summarizes methylation sensitivity for NEB restriction enzymes, indicating whether or not cleavage is blocked or impaired by Dam, Dcm or CpG methylation if or when it overlaps each recognition site. This table should be viewed as a guide to the behavior of the enzymes listed rather than an absolute indicator. Consult REBASE (<http://rebase.neb.com/rebase/>), the restriction enzyme database, for more detailed information and specific examples upon which these guidelines are based.

KEY POINTS TO CONSIDER

- Genomic DNA directly isolated from a mammalian source is not Dcm or Dam methylated, and is therefore not an issue when digesting mammalian DNA.
- Mammalian and plant DNA that has been cloned into a methylating *E. coli* strain will be Dam/Dcm methylated. Most commonly used laboratory *E. coli* strains methylate DNA.
- Directly isolated mammalian and plant genomic DNA are CpG methylated. Some enzymes are inhibited by CpG methylation. (See www.neb.com for more information).
- Most bacterial DNA (including *E. coli* DNA) is not CpG methylated. Inhibition of enzyme activity by CpG methylation is not an issue for DNA prepared from *E. coli* strains.
- DNA amplified by PCR does not contain any methylated bases.
- To avoid Dam/Dcm methylation when subcloning in bacteria, NEB offers the methyltransferase deficient cloning strain *dam*⁻/*dcm*⁻ Competent *E. coli* (NEB #C2925) for propagation.

References

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2. Geier, G.E. and Modrich, P. (1979) *J. Biol. Chem.*, 254, 1408–1413.
3. May, M.S. and Hattman, S. (1975) *J. Bacteriol.*, 123, 768–770.
4. Siegfried, Z. and Cedar, H. (1997) *Curr. Biol.*, 7, r305–307.



Methylation Sensitivity – Dam, Dcm and CpG Methylation

Legend:

●	not sensitive
■	blocked
□ ol	blocked by overlapping
□ scol	blocked by some combinations of overlapping
◆	impaired
◇ ol	impaired by overlapping
◇ scol	impaired by some combinations of overlapping

Single Letter Code:

R = A or G	Y = C or T	M = A or C
K = G or T	S = C or G	W = A or T
H = A or C or T	B = C or G or T	V = A or C or G
D = A or G or T	N = A or C or G or T	

ENZYME	SEQUENCE	Dam	Dcm	CpG
AatII	GACGT/C	●	●	■
AccI	GT/MKAC	●	●	□ ol
Acc65I	G/GTACC	●	□ scol	□ scol
Acil	CCGC(-3/-1)	●	●	■
AclI	AA/CGTT	●	●	■
AcuI	CTGAAG(16/14)	●	●	●
AfeI	AGC/GCT	●	●	■
AfilI	C/TTAAG	●	●	●
AfilII	A/CRYGT	●	●	●
AgeI	A/CCGGT	●	●	■
AgeI-HF	A/CCGGT	●	●	■
AgeI-HF RE-Mix	A/CCGGT	●	●	■
AhdI	GACNNN/NGTCC	●	●	◇ scol
AleI	CACNN/NGGTG	●	●	◇ scol
AluI	AG/CT	●	●	●
AlwI	GGATC(4/5)	■	●	●
AlwNI	CAGNNN/CTG	●	□ ol	●
ApaI	GGGCC/C	●	□ ol	□ ol
ApalI	G/TGCAC	●	●	□ ol
ApeKI	G/CWGC	●	●	□ ol
ApoI	R/AATY	●	●	●
AscI	GG/CGCGCC	●	●	■
AscI RE-Mix	GG/CGCGCC	●	●	■
Asel	AT/TAAT	●	●	●
AsiSI	GCGAT/CGC	●	●	■
AvaI	C/YCGRG	●	●	■
AvaII	G/GWCC	●	□ ol	□ ol
AvrII	C/CTAGG	●	●	●
BaeI	(10/15)ACNNNNGTAYC(12/7)	●	●	□ scol
BaeGI	GKGCM/C	●	●	●
BamHI	G/GATCC	●	●	●
BamHI-HF	G/GATCC	●	●	●
BanI	G/GYRCC	●	□ scol	□ scol
BanII	GRGCY/C	●	●	●
BbsI	GAAGAC(2/6)	●	●	●
BbvI	GCAGC(8/12)	●	●	●
BbvCI	CCTCAGC(-2/-5)	●	●	◇ ol
BccI	CCATC(4/5)	●	●	●
BceAI	ACGGC(12/14)	●	●	■
BcgI	(10/12)CGANNNNNNTGC(12/10)	◇ ol	●	□ scol
BcoDI	GTCTC(1/5)	●	●	□ scol

ENZYME	SEQUENCE	Dam	Dcm	CpG
BciVI	GTATCC(6/5)	●	●	●
BclI	T/GATCA	■	●	●
Bfal	C/TAG	●	●	●
BfuAI	ACCTGC(4/8)	●	●	◇ ol
BfuCI	/GATC	●	●	□ ol
BglI	GCCNNNN/NGGC	●	●	□ scol
BglII	A/GATCT	●	●	●
BlpI	GC/TNAGC	●	●	●
BmgBI	CACGTC(-3/-3)	●	●	■
BmrI	ACTGGG(5/4)	●	●	●
BmtI	GCTAG/C	●	●	●
BmtI-HF	GCTAG/C	●	●	●
BpmI	CTGGAG(16/14)	●	●	●
Bpu10I	CCTNAGC(-5/-2)	●	●	●
BpuEI	CTTGAG(16/14)	●	●	●
BsaI	GGTCTC(1/5)	●	◇ scol	□ scol
BsaI-HF	GGTCTC(1/5)	●	□ ol	□ scol
BsaAI	YAC/GTR	●	●	■
BsaBI	GATNN/NNATC	□ ol	●	□ scol
BsaHI	GR/CGYC	●	□ scol	■
BsaJI	C/CNNGG	●	●	●
BsaWI	W/CCGGW	●	●	●
BsaXI	(9/12)ACNNNNNCTCC(10/7)	●	●	●
BseRI	GAGGAG(10/8)	●	●	●
BseYI	CCCAGC(-5/-1)	●	●	□ ol
BsgI	GTGCAG(16/14)	●	●	●
BsiEI	CGRY/CG	●	●	■
BsiHKA1	GWGCW/C	●	●	●
BsiWI	C/GTACG	●	●	■
BsII	CCNNNNN/NGGG	●	□ scol	□ scol
BsmI	GAATGC(1/-1)	●	●	●
BsmAI	GTCTC(1/5)	●	●	□ scol
BsmBI	CGTCTC(1/5)	●	●	■
BsmFI	GGGAC(10/14)	●	□ ol	□ ol
BsoBI	C/YCGRG	●	●	●
Bsp1286I	GDGCH/C	●	●	●
BspCNI	CTCAG(9/7)	●	●	●
BspDI	AT/CGAT	□ ol	●	■
BspEI	T/CCGGA	□ ol	●	◆
BspHI	T/CATGA	◇ ol	●	●
BspMI	ACCTGC(4/8)	●	●	●



METHYLATION SENSITIVITY

ENZYME	SEQUENCE	Dam	Dcm	CpG
BspQI	GCTCTTC(1/4)	●	●	●
BsrI	ACTGG(1/-1)	●	●	●
BsrBI	CCGCTC(-3/-3)	●	●	□ scol
BsrDI	GCAATG(2/0)	●	●	●
BsrFI	R/CCGGY	●	●	■
BsrGI	T/GTACA	●	●	●
BsrGI-HF	T/GTACA	●	●	●
BssHII	G/CGCGC	●	●	■
BssKI	/CCNGG	●	□ ol	□ ol
BssS ^{nl}	CACGAG(-5/-1)	●	●	●
BstAPI	GCANNNN/NTGC	●	●	□ scol
BstBI	TT/CGAA	●	●	■
BstEII	G/GTNACC	●	●	●
BstEII-HF	G/GTNACC	●	●	●
BstEII-HF RE-Mix	G/GTNACC	●	●	●
BstNI	CC/WGG	●	●	●
BstUI	CG/CG	●	●	■
BstXI	CCANNNN/NTGG	●	□ scol	●
BstYI	R/GATCY	●	●	●
BstZ17I	GTA/TAC	●	●	□ scol
Bsu36I	CC/TNAGG	●	●	●
BtgI	C/CRYGG	●	●	●
BtgZI	GCGATG(10/14)	●	●	◆
Bts ^{nl}	GCAGTG(2/0)	●	●	●
BtsIMutI	CAGTG(2/0)	●	●	●
BtsCI	GGATG(2/0)	●	●	●
CacBI	GCN/NGC	●	●	□ scol
Clal	AT/CGAT	□ ol	●	■
CspCI	(11/13)CAANNNNNGTGG(12/10)	●	●	●
CviAII	C/ATG	●	●	●
CviKI-1	RG/CY	●	●	●
CviQI	G/TAC	●	●	●
Ddel	C/TNAG	●	●	●
Dpnl	GA/TC	●	●	□ ol
DpnII	/GATC	■	●	●
Dral	TTT/AAA	●	●	●
DrallI-HF	CACNNN/GTG	●	●	◇ scol
DrdI	GACNNNN/NGTGC	●	●	□ scol
EaeI	Y/GGCCR	●	□ ol	□ ol
EagI	C/GGCCG	●	●	■
EagI-HF	C/GGCCG	●	●	■
EarI	CTCTTC(1/4)	●	●	◇ ol
Ecil	GGCGGA(11/9)	●	●	□ scol
Eco53kl	GAG/CTC	●	●	□ scol
EcoNI	CCTNN/NNNAGG	●	●	●
EcoO109I	RG/GNCCY	●	□ ol	●
EcoP15I	CAGCAG(25/27)	●	●	●
EcoRI	G/AATTC	●	●	□ scol
EcoRI-HF	G/AATTC	●	●	□ scol
EcoRI-HF RE-Mix	G/AATTC	●	●	□ scol
EcoRV	GAT/ATC	●	●	◇ scol
EcoRV-HF	GAT/ATC	●	●	◇ scol
EcoRV-HF RE-Mix	GAT/ATC	●	●	◇ scol

ENZYME	SEQUENCE	Dam	Dcm	CpG
FatI	/CATG	●	●	●
FauI	CCC GC(4/6)	●	●	■
Fnu4HI	GC/NGC	●	●	□ ol
FokI	GGATG(9/13)	●	◇ ol	◇ ol
FseI	GGCCGG/CC	●	◇ scol	■
FspI	TGC/GCA	●	●	■
FspEI	C5mC>NNNNNNNNNN	●	●	●
Haell	RGCGC/Y	●	●	■
HaellI	GG/CC	●	●	●
Hgal	GACGC(5/10)	●	●	■
HhaI	GCG/C	●	●	■
HincII	GTY/RAC	●	●	□ scol
HindIII	A/AGCTT	●	●	●
HindIII-HF	A/AGCTT	●	●	●
Hinfl	G/ANTC	●	●	□ scol
HinP1I	G/CGC	●	●	■
HpaI	GTT/AAC	●	●	□ scol
HpaII	C/CGG	●	●	■
HphI	GGTGA(8/7)	■	■	●
Hpy99I	CGWCG/	●	●	■
Hpy166II	GTN/NAC	●	●	□ ol
Hpy188I	TCN/GA	□ ol	●	●
Hpy188III	TC/NNGA	□ ol	●	□ ol
HpyAV	CCTTC(6/5)	●	●	◇ ol
HpyCH4III	ACN/GT	●	●	●
HpyCH4IV	A/CGT	●	●	■
HpyCH4V	TG/CA	●	●	●
KasI	G/GCGCC	●	●	■
KpnI	GGTAC/C	●	●	●
KpnI-HF	GGTAC/C	●	●	●
KpnI-HF RE-Mix	GGTAC/C	●	●	●
LpnPI	C5mCDGNNNNNNNN	●	●	●
Mbol	/GATC	■	●	◇ ol
MbolI	GAAGA(8/7)	□ ol	●	●
MfeI	C/AATTG	●	●	●
MfeI-HF	C/AATTG	●	●	●
MfeI-HF RE-Mix	C/AATTG	●	●	●
MluI	A/CGCGT	●	●	■
MluI-HF	A/CGCGT	●	●	■
MluCI	/AATT	●	●	●
MlyI	GAGTC(5/5)	●	●	●
MmeI	TCCRAC(20/18)	●	●	□ ol
MnlI	CCTC(7/6)	●	●	●
MscI	TGG/CCA	●	□ ol	●
MseI	T/TAA	●	●	●
MslI	CAYNN/NNRTG	●	●	●
MspI	C/CGG	●	●	●
MspA1I	CMG/CKG	●	●	□ ol
MspJ	5mC>NNNNNNNNNN	●	●	●
MwoI	GCNNNN/NGC	●	●	□ scol
NaeI	GCC/GGC	●	●	■
NarI	GG/CGCC	●	●	■
Nb.BbvCI	CCTCAGC (none/-2)	●	●	●



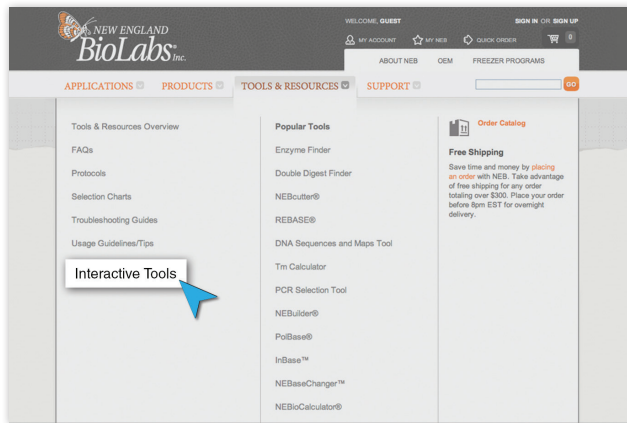
ENZYME	SEQUENCE	Dam	Dcm	CpG
Nb.BsmI	GAATGC (none/-2)	●	●	●
Nb.BsrDI	GCAATG (none/0)	●	●	●
Nb.BtsI	GCAAGT	●	●	●
NciI	CC/SGG	●	●	◇ ol
NcoI	C/CATGG	●	●	●
NcoI-HF	C/CATGG	●	●	●
NcoI-HF RE-Mix	C/CATGG	●	●	●
NdeI	CA/TATG	●	●	●
NgoMIV	G/CCGGC	●	●	■
NheI	G/CTAGC	●	●	□ scol
NheI-HF	G/CTAGC	●	●	□ scol
NheI-HF RE-Mix	G/CTAGC	●	●	□ scol
NlaIII	CATG/	●	●	●
NlaIV	GGN/NCC	●	□ ol	□ ol
NmeAIII	GCCGAG(21/19)	●	●	●
NotI	GC/GGCCCG	●	●	■
NotI-HF	GC/GGCCCG	●	●	■
NotI-HF RE-Mix	GC/GGCCCG	●	●	■
NruI	TCG/CGA	□ ol	●	■
NruI-HF	TCG/CGA	□ ol	●	■
Nsil	ATGCA/T	●	●	●
Nsil-HF	ATGCA/T	●	●	●
NspI	RCATG/Y	●	●	●
Nt.AlwI	GGATC(4/-5)	■	●	●
Nt.BbvCI	CCTCAGC(-5/none)	●	●	□ scol
Nt.BsmAI	GTCTC(1/none)	●	●	■
Nt.BspQI	GCTCTTC(1/none)	●	●	●
Nt.BstNBI	GAGTC(4/none)	●	●	●
Nt.CviPII	(0/-1)CCD	●	●	■
PacI	TTAAT/TAA	●	●	●
PacI RE-Mix	TTAAT/TAA	●	●	●
PaeR7I	C/TCGAG	●	●	■
PciI	A/CATGT	●	●	●
PfiFI	GACN/NNGTC	●	●	●
PfiMI	CCANNNN/NTGG	●	□ ol	●
PhoI	GG/CC	●	◇ scol	◇ scol
PleI	GAGTC(4/5)	●	●	□ scol
PluTI	GGCGC/C	●	●	■
PmeI	GTTT/AAAC	●	●	□ scol
PmlI	CAC/GTG	●	●	■
PpuMI	RG/GWCCY	●	□ ol	●
PshAI	GACNN/NNGTC	●	●	□ scol
PsiI	TTA/TAA	●	●	●
PspGI	/CCWGG	●	■	●
PspOMI	G/GGCC	●	◇ scol	□ ol
PspXI	VC/TCGAGB	●	●	◆
PstI	CTGCA/G	●	●	●
PstI-HF	CTGCA/G	●	●	●
PvuI	CGAT/CG	●	●	■
PvuI-HF	CGAT/CG	●	●	■
PvuII	CAG/CTG	●	●	●
PvuII-HF	CAG/CTG	●	●	●
RsaI	GT/AC	●	●	□ scol

ENZYME	SEQUENCE	Dam	Dcm	CpG
RsrII	CG/GWCCG	●	●	■
SacI	GAGCT/C	●	●	●
SacI-HF	GAGCT/C	●	●	●
SacII	CCGC/GG	●	●	■
Sall	G/TCGAC	●	●	■
Sall-HF	G/TCGAC	●	●	■
Sall-HF RE-Mix	G/TCGAC	●	●	■
SapI	GCTCTTC(1/4)	●	●	●
Sau3AI	/GATC	●	●	□ ol
Sau96I	G/GNCC	●	□ ol	□ ol
SbfI	CCTGCA/GG	●	●	●
SbfI-HF	CCTGCA/GG	●	●	●
Scal-HF	AGT/ACT	●	●	●
Scal-HF RE-Mix	AGT/ACT	●	●	●
ScrFI	CC/NGG	●	□ ol	□ ol
SexAI	A/CCWGGT	●	■	●
SfaNI	GCATC(5/9)	●	●	◇ scol
Sfcl	C/TRYAG	●	●	●
SfiI	GGCCNNNN/NGGCC	●	◇ ol	□ scol
SfoI	GGC/GCC	●	□ scol	■
SgrAI	CR/CCGGYG	●	●	■
SmaI	CCC/GGG	●	●	■
SmlI	C/TYRAG	●	●	●
SnaBI	TAC/GTA	●	●	■
SpeI	A/CTAGT	●	●	●
SpeI RE-Mix	A/CTAGT	●	●	●
SpeI-HF	A/CTAGT	●	●	●
SphI	GCATG/C	●	●	●
SphI-HF	GCATG/C	●	●	●
Sspl	AAT/ATT	●	●	●
Sspl-HF	AAT/ATT	●	●	●
StuI	AGG/CCT	●	□ ol	●
StyI	C/CWWGG	●	●	●
StyI-HF	C/CWWGG	●	●	●
StyD4I	/CCNGG	●	□ ol	◇ ol
Swal	ATTT/AAAT	●	●	●
TaqI	T/CGA	□ ol	●	●
TfiI	G/AWTC	●	●	□ scol
TseI	G/CWGC	●	●	□ scol
Tsp45I	/GTSAC	●	●	●
TspMI	C/CCGGG	●	●	■
TspRI	NNCASTGNN/	●	●	●
Tth111I	GACN/NNGTC	●	●	●
XbaI	T/CTAGA	□ ol	●	●
XbaI RE-Mix	T/CTAGA	□ ol	●	●
XcmI	CCANNNNN/NNNTGG	●	●	●
XhoI	C/TCGAG	●	●	◆
XhoI RE-Mix	C/TCGAG	●	●	◆
XmaI	C/CCGGG	●	●	◆
XmnI	GAANN/NNTTC	●	●	●
ZraI	GAC/GTC	●	●	■



Online Tools

The Tools & Resources tab, accessible on our homepage, contains a selection of interactive technical tools for use with restriction enzymes. These tools can also be accessed directly in the footer of every web page.



NEB Tools for Restriction Enzymes

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.

NEBioCalculator®



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

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. It is also very helpful with double digests! While you are there, you can also, find other relevant tools and resources to enable protocol optimization.

NEBcutter® V2.0



Identify restriction sites within your DNA sequence using NEBcutter. Choose between Type II and commercially available Type III enzymes to digest your DNA. NEBcutter V2.0 indicates cut frequency and methylation sensitivity.

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.

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 CutSmart, 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.



Cleavage Close to the Ends of DNA Fragments

To simulate cloning reactions, a selection of NEB restriction enzymes have been tested for their ability to cleave close to the end of a DNA fragment. Reaction conditions are described below. Note that the data reported represents the minimum number of bases that will work, and will not necessarily result in maximum cleavage. As a general rule, 6 base pairs should be added on either side of a restriction enzyme recognition site to cleave efficiently.

Experimental: Linearized vectors were incubated with the indicated enzymes (10 units/μg) for 60 minutes at the recommended reaction conditions for each enzyme. Following ligation and transformation, cleavage efficiencies were determined by dividing the number of transformants from the digestion reaction by the number obtained from religation of the linearized DNA (typically 100–500 colonies) and subtracting from 100%. “Base Pairs from End” refers to the number of double-stranded base pairs between the recognition site and the terminus of the fragment; this number does not include the single-stranded overhang from the initial cut.

TOOLS & RESOURCES

Visit www.neb.com for:

- Technical information including additional charts, protocols and technical tips related to restriction enzymes

ENZYME	BASE PAIRS FROM END	% CLEAVAGE EFFICIENCY	VECTOR	INITIAL CUT
AatII	3	88	LITMUS 29	NcoI
	2	100	LITMUS 28	NcoI
	1	95	LITMUS 29	PinAI
Acc65I	2	99	LITMUS 29	SpeI
	1	75	pNEB193	SacI
AfIII	1	13	LITMUS 29	StuI
AgeI	1	100	LITMUS 29	XbaI
	1	100	LITMUS 29	AatII
Apal	2	100	LITMUS 38	SpeI
AscI	1	97	pNEB193	BamHI
AvrII	1	100	LITMUS 29	SacI
BamHI	1	97	LITMUS 29	HindIII
BglIII	3	100	LITMUS 29	NsiI
BsiWI	2	100	LITMUS 29	BssHII
BspEI	2	100	LITMUS 39	BsrGI
	1	8	LITMUS 38	BsrGI
BsrGI	2	99	LITMUS 39	SphI
	1	88	LITMUS 38	BspEI
BssHII	2	100	LITMUS 29	BsiWI
EagI	2	100	LITMUS 39	NheI
EcoRI	1	100	LITMUS 29	XhoI
	1	88	LITMUS 29	PstI
	1	100	LITMUS 39	NheI
EcoRV	1	100	LITMUS 29	PstI
HindIII	3	90	LITMUS 29	NcoI
	2	91	LITMUS 28	NcoI
	1	0	LITMUS 29	BamHI
KasI	2	97	LITMUS 38	NgoMIV
	1	93	LITMUS 38	HindIII
KpnI	2	100	LITMUS 29	SpeI
	2	100	LITMUS 29	SacI
	1	99	pNEB193	SacI
MluI	2	99	LITMUS 39	EagI

ENZYME	BASE PAIRS FROM END	% CLEAVAGE EFFICIENCY	VECTOR	INITIAL CUT
MunI	2	100	LITMUS 39	NgoMIV
NcoI	2	100	LITMUS 28	HindIII
NgoMIV	2	100	LITMUS 39	MunI
NheI	1	100	LITMUS 39	EcoRI
	2	82	LITMUS 39	EagI
NotI	7	100	Bluescript SK-	SpeI
	4	100	Bluescript SK-	KspI
	1	98	Bluescript SK-	XbaI
NsiI	3	100	LITMUS 29	BssHII
	3	77	LITMUS 29	BglIII
	2	95	LITMUS 28	BssHII
PacI	1	76	pNEB193	BamHI
	1	94	pNEB193	PstI
PstI	3	98	LITMUS 29	EcoR V
	2	50	LITMUS 39	HindIII
	1	37	LITMUS 29	EcoRI
SacI	1	99	LITMUS 29	AvrII
Sall	3	89	LITMUS 39	SpeI
	2	23	LITMUS 39	SphI
SfiI*	1	61	LITMUS 38	SphI
	9	81	LITMUS 38	BamHI
	4	97	LITMUS 38	MluI
SpeI	1	93	LITMUS 38	EcoRI
	2	100	LITMUS 29	Acc65I
	2	100	LITMUS 29	KpnI
SphI	2	99	LITMUS 39	Sall
	2	97	LITMUS 39	BsrGI
	1	92	LITMUS 38	Sall
XbaI	1	99	LITMUS 29	AgeI
	1	94	LITMUS 29	PinAI
XhoI	1	97	LITMUS 29	EcoRI
XmaI	2	98	pNEB193	AscI
	2	92	pNEB193	BssHII

* A modified version of LITMUS 38 with an introduced SfiI site was used for this test.



Performance Chart for Restriction Enzymes

New England Biolabs supplies > 200 restriction enzymes that are 100% active in a single buffer, CutSmart. 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 BSA is now included in all NEBuffers, and is no longer provided as a separate tube. In addition, this performance chart shows recommended reaction temperature, heat-inactivation temperature, recommended diluent buffer, methylation sensitivity and whether the enzyme is Time-Saver™ qualified (i.e., cleaves substrate in 5–15 minutes under recommended conditions, and can be used overnight without degradation of DNA).

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.	SAM	Supplied with a separate vial of S-adenosylmethionine (SAM). To obtain 100% activity, SAM should be added to the 1X reaction mix as specified on the product data card.
Rri	Recombinant	dcm	dcm methylation sensitivity
⌚	Time-Saver qualified	CpG	CpG methylation sensitivity
e	Engineered enzyme for maximum performance	Mix	RE-Mix Master Mix version available
dam	dam 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 LIGATION AND RECUTTING	
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.

NEBuffer Compositions (1X)

NEBuffer 1.1	10 mM Bis Tris Propane-HCl, 10 mM MgCl ₂ , 100 µg/ml BSA (pH 7.0 @ 25°C).
NEBuffer 2.1	10 mM Tris-HCl, 10 mM MgCl ₂ , 50 mM NaCl, 100 µg/ml BSA (pH 7.9 @ 25°C).
NEBuffer 3.1	50 mM Tris-HCl, 10 mM MgCl ₂ , 100 mM NaCl, 100 µg/ml BSA (pH 7.9 @ 25°C).
CutSmart Buffer	20 mM Tris-acetate, 10 mM magnesium acetate, 50 mM potassium acetate, 100 µg/ml BSA (pH 7.9 @ 25°C).
Diluent A	50 mM KCl, 10 mM Tris-HCl, 0.1 mM EDTA, 1 mM dithiothreitol, 200 µg/ml BSA (pH 7.4 @ 25°C).
Diluent B	300 mM NaCl, 10 mM Tris-HCl, 0.1 mM EDTA, 1 mM dithiothreitol, 500 µg/ml BSA, 50% glycerol (pH 7.4 @ 25°C).
Diluent C	50 mM KCl, 10 mM Tris-HCl, 0.1 mM EDTA, 1 mM dithiothreitol, 0.15% Triton X-100, 200 µg/ml BSA 50% glycerol (pH 7.4 @ 25°C).

ENZYME	SUPPLIED NEBUFFER	% ACTIVITY IN NEBUFFERS				INCUB. TEMP. (°C)	INACTIV. TEMP. (°C)	DIL.	SUBSTRATE	METHYLATION SENSITIVITY	NOTE(S)	
		1.1	2.1	3.1	CUTSMART							
Rri ⌚	AatII	CutSmart	< 10	50*	50	100	37°	80°	B	Lambda	CpG	
Rri	AbaSI	4	25	50	50	100	25°	65°	C	T4 wt Phage		e
Rri ⌚	AccI	CutSmart	50	50	10	100	37°	80°	A	Lambda	CpG	
Rri ⌚	Acc65I	3.1	10	75*	100	25	37°	65°	A	pBC4	dcm CpG	
Rri ⌚	AcII	CutSmart	< 10	25	100	100	37°	65°	A	Lambda	CpG	
Rri ⌚	AcII	CutSmart	< 10	< 10	< 10	100	37°	No	B	Lambda	CpG	
Rri ⌚	AcuI	CutSmart + SAM	50	100	50	100	37°	65°	B	Lambda		3, b, d
Rri	AfeI	CutSmart	25	100	25	100	37°	65°	B	pXba	CpG	
Rri ⌚	AfiIII	CutSmart	50	100	10	100	37°	65°	A	phiX174		
Rri	AfiIII	3.1	10	50	100	50	37°	80°	B	Lambda		
Rri	AgeI	1.1	100	75	25	75	37°	65°	C	Lambda	CpG	2
Rri ⌚ e Mix	AgeI-HF	CutSmart	100	50	10	100	37°	65°	A	Lambda	CpG	
Rri ⌚	AhdI	CutSmart	25	25	10	100	37°	65°	A	Lambda	CpG	a
Rri	AleI	CutSmart	< 10	< 10	< 10	100	37°	80°	B	Lambda	CpG	
Rri ⌚	AluI	CutSmart	25	100	50	100	37°	80°	B	Lambda		b
Rri	AlwI	CutSmart	50	50	10	100	37°	No	A	Lambda dam-	dam	1, b, d
Rri ⌚	AlwNI	CutSmart	10	100	50	100	37°	80°	A	Lambda	dcm	
Rri ⌚	ApaI	CutSmart	25	25	< 10	100	25°	65°	A	pXba	dam CpG	
Rri ⌚	ApaLI	CutSmart	100	100	10	100	37°	No	A	Lambda HindIII	CpG	
Rri ⌚	ApeKI	3.1	25	50	100	10	75°	No	B	Lambda	CpG	



RR	ENZYME	SUPPLIED NEBUFFER	% ACTIVITY IN NEBUFFERS				CUTSMART	INCUB. TEMP. (°C)	INACTIV. TEMP. (°C)	DIL.	SUBSTRATE	METHYLATION SENSITIVITY	NOTE(S)
			1.1	2.1	3.1								
RR	ApoI	3.1	10	75	100	75	50°	80°	A	Lambda			
RR	Mix	Ascl	CutSmart	< 10	10	10	100	37°	80°	A	Lambda	CpG	
RR	Asel	3.1	< 10	50*	100	10	37°	65°	B	Lambda		3	
RR	AsiSI	CutSmart	50	100	100	100	37°	80°	B	pXba (Xho digested)	CpG	2, b	
RR	AvaI	CutSmart	< 10	100	25	100	37°	80°	A	Lambda	CpG		
RR	Avall	CutSmart	50	75	10	100	37°	80°	A	Lambda	dcm CpG		
RR	AvrII	CutSmart	100	50	50	100	37°	No	B	Lambda HindIII			
RR	BaeI	CutSmart + SAM	50	100	50	100	25°	65°	A	Lambda	CpG	e	
RR	BaeGI	3.1	75	75	100	25	37°	80°	A	Lambda			
RR	BamHI	3.1	75*	100*	100	100*	37°	No	A	Lambda		3	
RR	BamHI-HF	CutSmart	100	50	10	100	37°	No	A	Lambda			
RR	BanI	CutSmart	10	25	< 10	100	37°	65°	A	Lambda	dcm CpG	1	
RR	BanII	CutSmart	100	100	50	100	37°	80°	A	Lambda		2	
RR	BbsI	2.1	100	100	25	75	37°	65°	B	Lambda			
RR	BbvI	CutSmart	100	100	25	100	37°	65°	B	pBR322		3	
RR	BbvCI	CutSmart	10	100	50	100	37°	No	B	Lambda	CpG	1, a	
RR	BccI	CutSmart	100	50	10	100	37°	65°	A	pXba		3, b	
RR	BceAI	3.1	100*	100*	100	100*	37°	65°	A	pBR322	CpG	1	
RR	BcgI	3.1 + SAM	10	75*	100	50*	37°	65°	A	Lambda	dam CpG	e	
RR	BciVI	CutSmart	100	25	< 10	100	37°	80°	C	Lambda		b	
RR	BclI	3.1	50	100	100	75	50°	No	A	Lambda dam-	dam		
RR	BcoDI	CutSmart	50	75	75	100	37°	No	B	Lambda	CpG		
RR	Bfal	CutSmart	< 10	10	< 10	100	37°	80°	B	Lambda		2, b	
RR	BfuAI	3.1	< 10	25	100	10	50°	65°	B	Lambda	CpG	3	
RR	BfuCI	CutSmart	100	50	25	100	37°	80°	B	Lambda	CpG		
RR	BglI	3.1	10	25	100	10	37°	65°	B	Lambda	CpG		
RR	BglIII	3.1	10	10	100	< 10	37°	No	A	Lambda			
RR	BlnI	CutSmart	50	100	10	100	37°	No	A	Lambda		d	
RR	BmgBI	3.1	< 10	10	100	10	37°	65°	B	Lambda	CpG	3, b, d	
RR	Bmri	2.1	75	100	75	100*	37°	65°	B	Lambda HindIII		b	
RR	BmtI	3.1	100	100	100	100	37°	65°	B	pXba		2	
RR	BmtI-HF	CutSmart	50	100	10	100	37°	65°	B	pXba			
RR	Bpml	3.1	75	100	100	100	37°	65°	B	Lambda		2	
RR	Bpu10I	3.1	10	25	100	25	37°	80°	B	Lambda		3, b, d	
RR	BpuEI	CutSmart + SAM	50*	100	50*	100	37°	65°	B	Lambda		d	
RR	BsaI	CutSmart	75*	75	100	100	37°	65°	B	pXba	dcm CpG	3	
RR	BsaI-HF	CutSmart	50	100	25	100	37°	65°	B	pXba	dcm CpG		
RR	BsaAI	CutSmart	100	100	100	100	37°	No	C	Lambda	CpG		
RR	BsaBI	CutSmart	50	100	75	100	60°	80°	B	Lambda dam-	dam CpG	2	
RR	BsaHI	CutSmart	50	100	100	100	37°	80°	A	Lambda	dcm CpG		
RR	BsaJI	CutSmart	50	100	100	100	60°	80°	A	Lambda			
RR	BsaWI	CutSmart	10	100	50	100	60°	80°	A	Lambda			
RR	BsaXI	CutSmart	50*	100*	10	100	37°	No	B	Lambda		e	
RR	BseRI	CutSmart	100*	100	75	100	37°	80°	A	Lambda		d	
RR	BseYI	3.1	10	50	100	50	37°	80°	B	Lambda	CpG	d	
RR	BsgI	CutSmart + SAM	25	50	25	100	37°	65°	B	Lambda		d	
RR	BsiEI	CutSmart	25	50	< 10	100	60°	No	A	Lambda	CpG		
RR	BsiHKA1	CutSmart	25	100	100	100	65°	No	B	Lambda			
RR	BsiWI	3.1	25	50*	100	25	55°	65°	B	phiX174	CpG		
RR	BsII	CutSmart	50	75	100	100	55°	No	A	Lambda	dcm CpG	b	
RR	BsmI	CutSmart	25	100	< 10	100	65°	80°	A	pBR322			
RR	BsmAI	CutSmart	50	100	100	100	55°	No	B	Lambda	CpG		



PERFORMANCE CHART

ENZYME	SUPPLIED NEBUFFER	% ACTIVITY IN NEBUFFERS				CUTSMART	INCUB. TEMP. (°C)	INACTIV. TEMP. (°C)	DIL.	SUBSTRATE	METHYLATION SENSITIVITY	NOTE(S)
		1.1	2.1	3.1								
	BsmBI	3.1	10	50*	100	25	55°	80°	B	Lambda	CpG	
	BsmFI	CutSmart	25	50	50	100	65°	80°	A	pBR322	dam CpG	1
	BsoBI	CutSmart	25	100	100	100	37°	80°	A	Lambda		
	Bsp1286I	CutSmart	25	25	25	100	37°	65°	A	Lambda		3
	BspCNI	CutSmart + SAM	100	75	10	100	25°	80°	A	Lambda		b
	BspDI	CutSmart	25	75	50	100	37°	80°	A	Lambda	dam CpG	
	BspEI	3.1	< 10	10	100	< 10	37°	80°	B	Lambda dam-	dam CpG	
	BspHI	CutSmart	< 10	50	25	100	37°	80°	A	Lambda	dam	
	BspMI	3.1	10	50*	100	10	37°	65°	B	Lambda		
	BspQI	3.1	100	100	100	100	50°	80°	B	Lambda		3
	BsrI	3.1	< 10	50	100	10	65°	80°	B	phiX174		b
	BsrBI	CutSmart	50	100	100	100	37°	80°	A	Lambda	CpG	d
	BsrDI	2.1	10	100	75	25	65°	80°	A	Lambda		3, d
	BsrFI	CutSmart	10	100*	100*	100	37°	No	C	pBR322	CpG	1
	BsrGI	2.1	25	100	100	25	37°	80°	A	Lambda		
	BsrGI-HF	CutSmart	10	100	100	100	37°	80°	A	Lambda		
	BssHII	CutSmart	100	100	100	100	50°	65°	B	Lambda	CpG	
	BssKI	CutSmart	50	100	100	100	60°	80°	A	Lambda	dam CpG	b
	BssS ^{cl}	CutSmart	10	25	< 10	100	37°	No	B	Lambda		
	BstAPI	CutSmart	50	100	25	100	60°	80°	A	Lambda	CpG	b
	BstBI	CutSmart	75	100	10	100	65°	No	A	Lambda	CpG	
	BstEII	3.1	10	75*	100	75*	60°	No	A	Lambda		3
	BstEII-HF	CutSmart	< 10	10	< 10	100	37°	No	A	Lambda		
	BstNI	3.1	10	100	100	75	60°	No	A	Lambda		a
	BstUI	CutSmart	50	100	25	100	60°	No	A	Lambda	CpG	b
	BstXI	3.1	< 10	50	100	25	37°	80°	B	Lambda	dam	3
	BstYI	2.1	25	100	75	100	60°	No	A	Lambda		
	BstZ17I	CutSmart	75	100	100	100	37°	No	B	Lambda	CpG	3, b
	Bsu36I	CutSmart	25	100	100	100	37°	80°	A	Lambda HindIII		b
	BtgI	CutSmart	50	100	100	100	37°	80°	B	pBR322		
	BtgZI	CutSmart	10	25	< 10	100	60°	80°	A	Lambda	CpG	3, b, d
	Bts ^{cl}	CutSmart	100	100	25	100	55°	No	A	Lambda		
	BtsIMutI	CutSmart	100	50	10	100	55°	80°	A	pUC19		b
	BtsCI	CutSmart	10	100	25	100	50°	80°	B	Lambda		
	Cac8I	CutSmart	50	75	100	100	37°	65°	B	Lambda	CpG	b
	Clal	CutSmart	10	50	50	100	37°	65°	A	Lambda dam-	dam CpG	
	CspCI	CutSmart + SAM	10	100	10	100	37°	65°	A	Lambda		e
	CviAII	CutSmart	50	50	10	100	25°	65°	C	pUC19		
	CviKI-1	CutSmart	25	100	100	100	37°	No	A	pBR322		1, b
	CviQI	3.1	75	100*	100	75*	25°	No	C	Lambda		b
	Ddel	CutSmart	75	100	100	100	37°	65°	B	Lambda		
	Dpnl	CutSmart	100	100	75	100	37°	80°	B	pBR322	dam CpG	b
	DpnII	U	25	25	100*	25	37°	65°	B	Lambda dam-	dam	
	Dral	CutSmart	75	75	50	100	37°	65°	A	Lambda		
	DraIII-HF	CutSmart	< 10	50	10	100	37°	No	B	Lambda	CpG	b
	DrdI	CutSmart	25	50	< 10	100	37°	65°	A	pUC19	CpG	3, b
	EaeI	CutSmart	10	50	< 10	100	37°	65°	A	Lambda	dam CpG	b
	EagI	3.1	10	25	100	10	37°	65°	C	pXba	CpG	
	EagI-HF	CutSmart	25	100	100	100	37°	65°	B	pXba	CpG	
	EarI	CutSmart	50	10	< 10	100	37°	65°	B	Lambda	CpG	b, d
	EcoI	CutSmart	100	50	50	100	37°	65°	A	Lambda	CpG	2
	Eco53kI	CutSmart	100	100	< 10	100	37°	65°	A	pXba	CpG	3, b

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.



ENZYME	SUPPLIED NEBUFFER	% ACTIVITY IN NEBUFFERS				CUTSMART	INCUB. TEMP. (°C)	INACTIV. TEMP. (°C)	DIL.	SUBSTRATE	METHYLATION SENSITIVITY	NOTE(S)
		1.1	2.1	3.1								
EcoNI	CutSmart	50	100	75	100	37°	65°	A	Lambda		b	
EcoO109I	CutSmart	50	100	50	100	37°	65°	A	Lambda HindIII	dam	3	
EcoP15I	3.1 + ATP	75	100	100	100	37°	65°	A	pUC19		e	
EcoRI	U	25	100*	50	50*	37°	65°	C	Lambda	CpG		
EcoRI-HF	CutSmart	10	100	< 10	100	37°	65°	C	Lambda	CpG		
EcoRV	3.1	10	50	100	10	37°	80°	A	Lambda	CpG		
EcoRV-HF	CutSmart	25	100	100	100	37°	65°	B	Lambda	CpG		
FatI	2.1	10	100	50	50	55°	80°	A	pUC19			
Faul	CutSmart	100	50	10	100	55°	65°	A	Lambda	CpG	3, b, d	
Fnu4HI	CutSmart	< 10	< 10	< 10	100	37°	No	A	Lambda	CpG	a	
FokI	CutSmart	100	100	75	100	37°	65°	A	Lambda	dam CpG	3, b, d	
FseI	CutSmart	100	75	< 10	100	37°	65°	B	Adenovirus-2	dam CpG		
FspI	CutSmart	10	100	10	100	37°	No	C	Lambda	CpG	b	
FspEI	4 + BSA	< 10	< 10	< 10	100	37°	80°	B	pBC4		2, e	
HaeII	CutSmart	25	100	10	100	37°	80°	A	Lambda	CpG		
HaeIII	CutSmart	50	100	25	100	37°	80°	A	Lambda			
HgaI	1.1	100	100	25	100	37°	65°	A	phiX174	CpG	1	
HhaI	CutSmart	25	100	100	100	37°	65°	A	Lambda	CpG		
HincII	3.1	25	100	100	100	37°	65°	B	Lambda	CpG		
HindIII	2.1	25	100	50	50	37°	80°	B	Lambda		2	
HindIII-HF	CutSmart	10	100	10	100	37°	80°	B	Lambda			
HinfI	CutSmart	50	100	100	100	37°	80°	A	Lambda	CpG		
HinP1I	CutSmart	100	100	100	100	37°	65°	A	Lambda	CpG		
HpaI	CutSmart	< 10	75*	25	100	37°	No	A	Lambda	CpG	1	
HpaII	CutSmart	100	50	< 10	100	37°	80°	A	Lambda	CpG		
HphI	CutSmart	50	50	< 10	100	37°	65°	B	Lambda	dam dcm	b, d	
Hpy99I	CutSmart	50	10	< 10	100	37°	65°	A	Lambda	CpG		
Hpy166II	CutSmart	100	100	50	100	37°	65°	C	pBR322	CpG		
Hpy188I	CutSmart	25	100	50	100	37°	65°	A	pBR322	dam	1, b	
Hpy188III	CutSmart	100	100	10	100	37°	65°	B	pUC19	dam CpG	3, b	
HpyAV	CutSmart	100	100	25	100	37°	65°	B	Lambda	CpG	3, b, d	
HpyCH4III	CutSmart	100	25	< 10	100	37°	65°	A	Lambda		b	
HpyCH4IV	CutSmart	100	50	25	100	37°	65°	A	pUC19	CpG		
HpyCH4V	CutSmart	50	50	25	100	37°	65°	A	Lambda			
KasI	CutSmart	50	100	50	100	37°	65°	B	pBR322	CpG	3	
KpnI	1.1	100	75	< 10	50*	37°	No	A	pXba		1	
KpnI-HF	CutSmart	100	25	< 10	100	37°	No	A	pXba			
LpnPI	4 + BSA	< 10	< 10	< 10	50	37°	65°	B	pBR322		2, e	
MboI	CutSmart	75	100	100	100	37°	65°	A	Lambda dam-	dam CpG		
MboII	CutSmart	100*	100	50	100	37°	65°	C	Lambda dam-	dam	b	
MfeI	CutSmart	75	50	10	100	37°	No	A	Lambda		2	
MfeI-HF	CutSmart	75	25	< 10	100	37°	No	A	Lambda			
MluI	3.1	10	50	100	25	37°	80°	A	Lambda	CpG		
MluI-HF	CutSmart	25	100	100	100	37°	No	A	Lambda	CpG		
MluCI	CutSmart	100	10	10	100	37°	No	A	Lambda			
MlyI	CutSmart	50	50	10	100	37°	65°	A	Lambda		b, d	
MmeI	CutSmart + SAM	50	100	50	100	37°	65°	B	phiX174	CpG	b, c	
MnlI	CutSmart	75	100	50	100	37°	65°	B	Lambda		b	
MscI	CutSmart	25	100	100	100	37°	80°	B	Lambda	dam		
MseI	CutSmart	75	100	75	100	37°	65°	A	Lambda			
MspI	CutSmart	50	50	< 10	100	37°	80°	A	Lambda			
MspI	CutSmart	75	100	50	100	37°	No	A	Lambda			

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.



PERFORMANCE CHART

RR	ENZYME	SUPPLIED NEBUFFER	% ACTIVITY IN NEBUFFERS				INCUB. TEMP. (°C)	INACTIV. TEMP. (°C)	DIL.	SUBSTRATE	METHYLATION SENSITIVITY	NOTE(S)
			1.1	2.1	3.1	CUTSMART						
RR	MspA1I	CutSmart	10	50	10	100	37°	65°	B	Lambda	CpG	
RR	MspJI	4 + BSA	< 10	< 10	< 10	50	37°	65°	B	pBR322		2, e
RR	Mwol	CutSmart	< 10	100	100	100	60°	No	B	Lambda	CpG	
RR	Nael	CutSmart	25	25	< 10	100	37°	No	A	pXba	CpG	b
RR	NarI	CutSmart	100	100	10	100	37°	65°	A	pXba	CpG	
RR	Nb.BbvCI	CutSmart	25	100	100	100	37°	80°	A	pUB		e
RR	Nb.BsmI	3.1	< 10	50	100	10	65°	80°	A	pBR322		e
RR	Nb.BsrDI	CutSmart	25	100	100	100	65°	80°	A	pUC19		e
RR	Nb.BtsI	CutSmart	75	100	75	100	37°	80°	A	phiX174		e
RR	NciI	CutSmart	100	25	10	100	37°	No	A	Lambda	CpG	b
RR	NcoI	3.1	100	100	100	100	37°	80°	A	Lambda		
RR	NcoI-HF	CutSmart	50	100	10	100	37°	80°	B	Lambda		
RR	NdeI	CutSmart	75	100	100	100	37°	65°	A	Lambda		
RR	NgoMIV	CutSmart	100	50	10	100	37°	No	A	pXba	CpG	1
RR	NheI	2.1	100	100	10	100	37°	65°	C	Lambda HindIII	CpG	
RR	NheI-HF	CutSmart	100	25	< 10	100	37°	80°	C	Lambda HindIII	CpG	
RR	NlaIII	CutSmart	< 10	< 10	< 10	100	37°	65°	B	phiX174		
RR	NlaIV	CutSmart	10	10	10	100	37°	65°	B	pBR322	dcm CpG	
RR	NmeAIII	CutSmart + SAM	10	10	< 10	100	37°	65°	B	phiX174		c
RR	NotI	3.1	< 10	50	100	25	37°	65°	C	pBC4	CpG	
RR	NotI-HF	CutSmart	25	100	25	100	37°	65°	A	pBC4	CpG	
RR	NruI	3.1	< 10	10	100	10	37°	No	A	Lambda	dam CpG	b
RR	NruI-HF	CutSmart	0	25	50	100	37°	No	A	Lambda	dam CpG	
RR	NsiI	3.1	10	75	100	25	37°	65°	B	Lambda		
RR	NsiI-HF	CutSmart	< 10	20	< 10	100	37°	80°	B	Lambda		
RR	NspI	CutSmart	100	100	< 10	100	37°	65°	A	Lambda		
RR	Nt.AlwI	CutSmart	10	100	100	100	37°	80°	A	pUC101 dam-dcm-	dam	e
RR	Nt.BbvCI	CutSmart	50	100	10	100	37°	80°	A	pUB	CpG	e
RR	Nt.BsmAI	CutSmart	100	50	10	100	37	65°	A	pBR322	CpG	e
RR	Nt.BspQI	3.1	< 10	25	100	10	50°	80°	B	pUC19		e
RR	Nt.BstNBI	3.1	0	10	100	10	55°	80°	A	T7		
RR	Nt.CviPII	CutSmart	< 10	100	25	100	37°	65°	A	pUC19	CpG	e
RR	Pacl	CutSmart	100	75	10	100	37°	65°	A	pNEB193		
RR	PaeR7I	CutSmart	25	100	10	100	37°	No	A	Lambda HindIII	CpG	
RR	PciI	3.1	50	75	100	50*	37°	80°	B	pXba		
RR	PfiFI	CutSmart	25	100	25	100	37°	65°	A	pBC4		b
RR	PfiMI	3.1	0	100	100	50	37°	65°	A	Lambda	dcm	3, b, d
RR	PleI	CutSmart	25	50	25	100	37°	65°	A	Lambda	CpG	b
RR	PluTI	CutSmart	100	25	< 10	100	37°	65°	A	pXba	CpG	
RR	PmeI	CutSmart	< 10	50	10	100	37°	65°	A	Lambda	CpG	
RR	PmlI	CutSmart	100	50	< 10	100	37°	65°	A	Lambda HindIII	CpG	
RR	PpuMI	CutSmart	< 10	< 10	< 10	100	37°	No	B	Lambda HindIII	dcm	
RR	PshAI	CutSmart	25	50	10	100	37°	65°	A	Lambda	CpG	
RR	PsiI	CutSmart	10	100	10	100	37°	65°	B	Lambda		3
RR	PspGI	CutSmart	25	100	50	100	75°	No	A	T7	dcm	3
RR	PspOMI	CutSmart	10	10	< 10	100	37°	65°	B	pXba	dcm CpG	
RR	PspXI	CutSmart	< 10	100	25	100	37°	No	B	Lambda HindIII	CpG	
RR	PstI	3.1	75	75	100	50*	37°	80°	C	Lambda		
RR	PstI-HF	CutSmart	10	75	50	100	37°	No	C	Lambda		
RR	PvuI	3.1	< 10	25	100	< 10	37°	80°	B	pXba	CpG	
RR	PvuI-HF	CutSmart	25	100	100	100	37°	No	B	pXba	CpG	

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.



ENZYME	SUPPLIED NEBUFFER	% ACTIVITY IN NEBUFFERS				CUTSMART	INCUB. TEMP. (°C)	INACTIV. TEMP. (°C)	DIL.	SUBSTRATE	METHYLATION SENSITIVITY	NOTE(S)
		1.1	2.1	3.1								
	PvuII	3.1	50	100	100	100*	37°	No	B	Lambda		
e	PvuII-HF	CutSmart	< 10	< 10	< 10	100	37°	80°	B	Lambda		
	RsaI	CutSmart	25	50	< 10	100	37°	No	A	Lambda	CpG	
	RsrII	CutSmart	25	75	10	100	37°	65°	C	Lambda	CpG	
	SacI	1.1	100	50	10	100	37°	65°	A	Lambda HindIII		
e	SacI-HF	CutSmart	10	50	< 10	100	37°	65°	A	Lambda HindIII		
	SacII	CutSmart	10	100	10	100	37°	65°	A	pXba	CpG	
	Sall	3.1	< 10	< 10	100	< 10	37°	65°	A	Lambda HindIII	CpG	
e Mix	Sall-HF	CutSmart	10	100	100	100	37°	65°	A	Lambda HindIII	CpG	
	SapI	CutSmart	75	50	< 10	100	37°	65°	B	Lambda		
	Sau3AI	1.1	100	50	10	100	37°	65°	A	Lambda	CpG b	
	Sau96I	CutSmart	50	100	100	100	37°	65°	A	Lambda	dcm CpG	
	SbfI	CutSmart	50	25	< 10	100	37°	80°	A	Lambda	3	
e	SbfI-HF	CutSmart	50	25	< 10	100	37°	80°	B	Lambda		
e Mix	Scal-HF	CutSmart	100	100	10	100	37°	80°	B	Lambda		
	ScrFI	CutSmart	100	100	100	100	37°	65°	C	Lambda	dcm CpG 2, a	
	SexAI	CutSmart	100	75	50	100	37°	65°	A	pBC4 dcm-	dcm 3, b, d	
	SfiNI	3.1	< 10	75	100	25	37°	65°	B	phiX174	CpG 3, b	
	Sfci	CutSmart	75	50	25	100	37°	65°	B	Lambda	3	
	SfiI	CutSmart	25	100	50	100	50°	No	C	pXba	dcm CpG	
	SfoI	CutSmart	50	100	100	100	37°	No	B	Lambda HindIII	dcm CpG	
	SgrAI	CutSmart	100	100	10	100	37°	65°	A	Lambda	CpG 1	
	SmaI	CutSmart	< 10	< 10	< 10	100	25°	65°	B	Lambda HindIII	CpG b	
	SmlI	CutSmart	25	75	25	100	55°	No	A	Lambda	b	
	SnaBI	CutSmart	50	50	10	100	37°	80°	A	T7	CpG 1	
Mix	SpeI	CutSmart	75	100	25	100	37°	80°	C	pXba-XbaI digested		
e	SpeI-HF	CutSmart	25	50	10	100	37°	80°	C	pXba-XbaI digested		
	SphI	2.1	100	100	50	100	37°	65°	B	Lambda	2	
e	SphI-HF	CutSmart	50	25	10	100	37°	65°	B	Lambda		
	SspI	U	50	100	50	50	37°	65°	C	Lambda		
e	SspI-HF	CutSmart	25	100	< 10	100	37°	No	B	Lambda		
	StuI	CutSmart	50	100	50	100	37°	No	A	Lambda	dcm	
	StyD4I	CutSmart	10	100	100	100	37°	65°	B	Lambda	dcm CpG	
	StyI	3.1	10	25	100	10	37°	65°	A	Lambda	b	
e	StyI-HF	CutSmart	25	100	25	100	37°	65°	A	Lambda		
	Swal	3.1	10	10	100	10	25°	65°	B	pUPS	b, d	
	Taq ^{ql}	CutSmart	50	75	100	100	65°	80°	B	Lambda	dam	
	TfiI	CutSmart	50	100	100	100	65°	No	C	Lambda	CpG	
	TseI	CutSmart	75	100	100	100	65°	No	B	Lambda	CpG 3	
	Tsp45I	CutSmart	100	50	< 10	100	65°	No	A	Lambda		
	TspMI	CutSmart	50*	75*	50*	100	75°	No	B	pUCAdeno	CpG d	
	TspRI	CutSmart	25	50	25	100	65°	No	B	Lambda		
	Tth111I	CutSmart	25	100	25	100	65°	No	B	pBC4	b	
Mix	XbaI	CutSmart	< 10	100	75	100	37°	65°	A	Lambda HindIII dam-	dam	
	XcmI	2.1	10	100	25	100	37°	65°	C	Lambda	2	
Mix	XhoI	CutSmart	75	100	100	100	37°	65°	A	Lambda HindIII	CpG b	
	XmaI	CutSmart	25	50	< 10	100	37°	65°	A	pXba	CpG 3	
	XmnI	CutSmart	50	75	< 10	100	37°	65°	A	Lambda	b	
	ZraI	CutSmart	100	25	10	100	37°	80°	B	Lambda	CpG	

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