

PRODUCT INFORMATION

BamHI

 #ER0051
 4000 U

 Lot: ____
 Expiry Date: ___

5'....**G**↓**G A T C C**....3'

3'...**C C T A G↑G**...5'

Concentration: Source: Supplied with: 10 U/µL Bacillus amyloliquefaciens H 2x1 mL of 10X Buffer BamHI 1 mL of 10X Buffer Tango







In total 4 vials.

BSA included

www.thermoscientific.com/onebio

RECOMMENDATIONS

1X Buffer BamHI (for 100% BamHI digestion) 10 mM Tris-HCI (pH 8.0), 5 mM MgCl₂, 100 mM KCI, 0.02% Triton X-100, 0.1 mg/mL BSA.

Incubation temperature

37°C.

Unit Definition

One unit is defined as the amount of BamHI required to digest 1 μ g of lambda DNA-Bsp120I fragments in 1 hour at 37°C in 50 μ L of recommended reaction buffer.

Dilution

Dilute with Dilution Buffer (#B19): 10 mM Tris-HCl (pH 7.4 at 25°C) 100 mM KCl, 1 mM EDTA, 1 mM DTT, 0.2 mg/mL BSA and 50% glycerol.

Double Digests

Thermo Scientific Tango Buffer is provided to simplify buffer selection for double digests. 98% of Thermo Scientific restriction enzymes are active in a 1X or 2X concentration of Tango[™] Buffer. Please refer to <u>www.thermoscientific.com/doubledigest</u> to choose the best buffer for your experiments. 1X Tango Buffer: 33 mM Tris-acetate (pH 7.9 at 37°C), 10 mM magnesium acetate, 66 mM potassium acetate, 0.1 mg/mL BSA.

Storage Buffer

BamHI is supplied in: 10 mM Tris-HCI (pH 7.4 at 25°C), 200 mM NaCl, 1 mM DTT, 0.1 mM EDTA, 0.15% Triton X-100, 0.2 mg/mL BSA and 50% glycerol.

Recommended Protocol for Digestion

• Add:

nuclease-free water16 μ L10X Buffer BamHI2 μ LDNA (0.5-1 μ g/ μ L)1 μ LBamHI0.5-2 μ L*,**

- Mix gently and spin down for a few seconds.
- Incubate at 37°C for 1-16 hours**.

The digestion reaction may be scaled either up or down.

Recommended Protocol for Digestion of PCR Products Directly after Amplification

• Add:

- Mix gently and spin down for a few seconds.
- Incubate at 37°C for 1-16 hours**.
- * This volume of the enzyme is recommended for preparations of standard concentrations (10 U/ μ L), whereas HC enzymes (50 U/ μ L) should be diluted with Dilution Buffer to obtain 10 U/ μ L concentration.
- ** See Overdigestion Assay.

Thermal Inactivation

Only small amounts of BamHI (up to 10 units) can be inactivated at 80°C in 20 min.

Inactivation Procedure

- To prepare the digested DNA for electrophoresis:
 - stop the digestion reaction by adding 0.5 M EDTA, pH 8.0 (#R1021), to achieve a 20 mM final concentration. Mix thoroughly, add an electrophoresis loading dye and load onto gel.
- To prepare DNA suitable for further enzymatic reactions:
 - extract with phenol/chloroform, precipitate with ethanol or isopropanol, wash the pellet with 75% cold ethanol and air-dry;
 - dissolve DNA in either nuclease-free water, TE buffer, or a buffer suitable for further applications;
 - check the DNA concentration in the solution.

For **ENZYME PROPERTIES** and **CERTIFICATE OF ANALYSIS**

see back page

ENZYME PROPERTIES

Enzyme Activity in Thermo Scientific REase Buffers, %

BamHI	В	G	0	R	Tango	2X Tango		
100	20-50**	100	20-50	50-100**	100**	50-100		
**Star activity appears at a greater than 5-fold overdigestion (5 U \times 1h).								

Methylation Effects on Digestion

Dam: completely overlaps - no effect.

Dcm: may overlap - no effect.

CpG: may overlap - no effect.

EcoKI: never overlaps - no effect.

EcoBI: never overlaps - no effect.

Stability during Prolonged Incubation

A minimum of 0.5 units of the enzyme is required for complete digestion of 1 μ g of lambda DNA in 16 hours at 37°C.

Digestion of Agarose-embedded DNA

A minimum of 5 units of the enzyme is required for complete digestion of 1 μg of agarose-embedded lambda DNA in 16 hours.

Compatible Ends

BcII, BgIII, Bsp143I, MboI, Psul

Number of Recognition Sites in DNA

λ	ФХ174	pBR322	pUC57	pUC18/19	pTZ19R/U	M13mp18/19
5	0	1	1	1	1	1

CERTIFICATE OF ANALYSIS

Overdigestion Assay

No detectable change in the specific fragmentation pattern is observed after an 80-fold overdigestion with BamHI (5 U/ μ g lambda DNA x 16 hours).

Ligation and Recleavage (L/R) Assay

The ligation and recleavage assay was replaced with L0 test after validating experiments showed L0 test ability to trace nuclease and phosphatase activities with sensitivity that is higher than L/R by a factor of 100.

Labeled Oligonucleotide (LO) Assay

No detectable degradation of single-stranded or doublestranded labeled oligonucleotides occurred during incubation with 10 units of BamHI for 4 hours.

Blue/White (B/W) Cloning Assay

The B/W assay was replaced with LO test after validating experiments showed LO test ability to detect nuclease and phosphatase activities with sensitivity that equals to that of B/W test.

Quality authorized by:



PRODUCT USE LIMITATION

This product is developed, designed and sold exclusively *for research purposes and in vitro use only.* The product was not tested for use in diagnostics or for drug development, nor is it suitable for administration to humans or animals.

Please refer to <u>www.thermoscientific.com/onebio</u> for Material Safety Data Sheet of the product.

© 2012 Thermo Fisher Scientific Inc. All rights reserved. All trademarks are the property of Thermo Fisher Scientific Inc. and its subsidiaries.