

PRODUCT INFORMATION

BseNI (BsrI)

#ER0882 5000 U

Lot: ___ Expiry Date: _

5'...**A** C T G G N↓...3'

3'... **T G A C**↑**C N** ...5'

Concentration: 10 U/µL

Source: Bacillus species N

Supplied with: 2 x 1 mL of 10X Buffer B

1 mL of 10X Buffer Tango

Store at -20°C









In total 4 vials.

BSA included

RECOMMENDATIONS

1X Buffer B (for 100% BseNI digestion)

10 mM Tris-HCl (pH 7.5), 10 mM MgCl₂, 0.1 mg/mL BSA.

Incubation temperature

65°C*.

Unit Definition

One unit is defined as the amount of BseNI required to digest 1 μ g of lambda DNA in 1 hour at 65°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 www.thermoscientific.com/doubledigest 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.

^{*} Incubation at 37°C results in less than 10% activity.

Storage Buffer

BseNI is supplied in: 10 mM Tris-HCI (pH 7.4 at 25°C), 100 mM KCI, 1 mM EDTA, 1 mM DTT, 0.2 mg/mL BSA and 50% glycerol.

Recommended Protocol for Digestion

Add:

nuclease-free water $16 \mu L$ 10X Buffer B $2 \mu L$ BseNI $0.5-2 \mu L$

- Mix gently and spin down for a few seconds.
- Incubate at 65°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:

PCR reaction mixture 10 μ L (~0.1-0.5 μ g of DNA) nuclease-free water 18 μ L 10X Buffer B 2 μ L BseNI 1-2 μ L

- Mix gently and spin down for a few seconds.
- Incubate at 65°C for 1-16 hours.

Thermal Inactivation

BseNI is inactivated by incubation at 80°C for 20 min.

ENZYME PROPERTIES

Enzyme Activity in Thermo Scientific REase Buffers, %

В	G	0	R	Tango	2X Tango
100	20-50	0-20	0-20	50-100	20-50

Methylation Effects on Digestion

Dam: never overlaps — no effect. Dcm: never overlaps — no effect. CpG: never overlaps — no effect.

EcoKI: may overlap — effect not determined. EcoBI: may overlap — effect not determined.

Stability during Prolonged Incubation

A minimum of 0.1 units of the enzyme is required for complete digestion of 1 µg of DNA in 16 hours at 65°C.

Number of Recognition Sites in DNA

λ	ФХ174	pBR322	pUC57	pUC18/19	pTZ19R/U	M13mp18/19
110	9	19	11	11	12	18

For **CERTIFICATE OF ANALYSIS** see back page

CERTIFICATE OF ANALYSIS

Overdigestion Assay

No detectable change in the specific fragmentation pattern is observed after a 160-fold overdigestion with BseNI (10 U/µg lambda DNA x 16 hours).

Ligation and Recleavage (L/R) Assay

The ligation and recleavage assay was replaced with LO test after validating experiments showed LO 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 BseNI for 4 hours.

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 www.thermoscientific.com/onebio 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.