

# **BsiHKCI**

5'-C Py C G Pu G-3' 3'-G Pu G C Py C-5'

Cat. No.	size
E2109-01	10000 units
E2109-02	50000 units

Reaction Temperature: 65°C

Inactivation Temperature (20 min): -

Prototype: Aval

Source: Bacillus stearothermophilus HKC

**Note 1:** Purified from E.coli strain that carries the cloned bsiHKCRI gene from Bacillus stearothermophilus HKC.

### **Package Contents:**

- BsiHKCI
- 10 x ONE Buffer
- BSA [100x]

Added as separate component to prevent reaction buffer precipitation.

#### Dilution Buffer # 1

Added for enzymes exceeding 10  $U/\mu I$  in concentration. High protein concentration warrants optimal stability during prolonged storage. Use dilution buffer to prepare short term working stocks (5-10  $U/\mu I$ , non-freezing at -20°C).

Storage Conditions: Store at -20°C.

# Double Digestion - Buffer Compatibility:

ONE Buffer is compatible with most EURx restriction enzymes.

\*The enzyme exhibits star activity under certain conditions.

# **Restriction Enzyme Buffer Compatibility:**

Both, enzyme and buffers are fully compatible to restrictases and buffer systems from other manufacturers and can be used along in double digestions. To obtain best results, consult the corresponding manuals of all involved products.

### **DNA Methylation:**

No Inhibition: dam, dcm, EcoKI Inhibition (Blocked): CpG

# Standard Reaction Protocol (for 50 µl volume):

Mix the following reaction components:

1-2 μg pure DNA or 10 μl PCR product (=~0.1-2 μg DNA)

5 μl 10 x ONE Buffer

0.5 μl BSA [100x]

1-2 U BsiHKCI (use 1 U per µg DNA, < 10% React. Volume!)

*Tips*: Add enzyme as last component. Mix components well before adding enzyme. After enzyme addition, mix gently by pipetting. Do not vortex. High (excess) amounts of enzyme can greatly speed up the reaction.

# add sterile $H_2O$ to $50 \mu I$ final volume

#### Incubate for 1 h at 65°C

To obtain complete digestion of high molecular weight DNA, (e.g. plant genomic DNA), add excess amounts of enzyme and prolong the incubation time

#### **Stop** reaction by alternatively

- (a) Addition of 2.1  $\mu$ l EDTA pH 8.0 [0.5 M], final 20 mM or
- (b) Heat Inactivation

(not applicable for this enzyme) or

(c) Spin Column DNA Purification

(e.g. EURx PCR/DNA Clean-Up Kit, Cat.No. E3520) or

(d) Gel Electrophoresis and Single Band Excision

(e.g. EURx Agarose-Out DNA Kit, Cat.No. E3540) or

(e) Phenol-Chloroform Extraction or Ethanol Precipitation.

# Non-optimal buffer conditions:

To compensate for the lack of enzyme activity, increase the amount of enzyme and/ or reaction time accordingly. The following values may serve as orientation:

- 1. Enzyme amount: Instead of 1 U enzyme, use ~4 U of enzyme in buffers providing 25% rel. activity, ~2 U in 50%, ~1.5 U in 75% or ~1 U in 100%, respectively.
- 2. Reaction time: Increase by ~1.3-fold (75% rel. activity), ~2-fold (50%) or ~4-fold (25%).

## **Unit Definition:**

One unit is the amount of enzyme required to completely digest 1  $\mu g$  of Lambda DNA in 1 hr. Total reaction volume is 50  $\mu$ l. Enzyme activity was determined in the recommended reaction buffer.

## **Reaction Buffer:**

1 x ONE Buffer

To be supplemented with 100  $\mu$ g/ml bovine serum albumin.

## **Storage Buffer:**

10 mM Tris-HCl (pH 7.5 at 22°C), 50 mM KCl, 1 mM dithiothreitol, 0.1 mM EDTA, 500  $\mu$ g/ml bovine serum albumin and 50% (v/v)glycerol.

## **Quality Control:**

All preparations are assayed for contaminating endonuclease, 3'-exonuclease, 5'-exonuclease/5'-phosphatase, as well as for nonspecific single- and double-stranded DNase activities.