# Kpn I

### GGTAC^C AccuCut™ Restriction Endonuclease

· Cat. No.

E-1761

3000 Units

E-1762

15000 Units

Lot No.: 02I131491H8A3

Supplied with Enzyme

10X AccuCut™areeN Buffer : 1 mL

100 mM

pH 7.6 Tris-HCI

100 mM

MaCl<sub>2</sub> DTT

10 mM 1X Dilution Buffer

10 mM

: 1 mL

50 mM

pH 7.6 Tris-HCI

 $0.1 \, \text{mM}$ 

KCI

1 mM

**EDTA** DTT

 $200 \mu \text{ g/mL}$ 

Acetylated BSA

50%

Glycerol

Store at -20 ℃.

• Unit definition: One unit of restriction endonuclease activity is defined as the amount of enzyme required to completely digest 1µg of substrate DNA in a total reaction volume of 50 µL in one hour using the AccuCut™ buffer provided. Incubations are performed in 1.5 mL tubes at the appropriate incubation temperature as indicated in the Product Profile.

•Isoschizomer · None

•Neoschizomer: Acc65 I, Asp718, Asp718 I (G^GTACC)

- •Reactivity on methylated substrate DNA:Blocked by GGT m6ACC, GGTA m4CC, GGTA m5C m5C, GGTAC m4C. Not blocked by GGTA m5CC, GGTAC m5C
- •Ref)1.Kiss, A., Finta, C., Venetianer, P., (1991) Nucleic Acids Res., vol. 19, pp. 3460.
  - 2.Smith, D.I., Blattner, F.R., Davies, J., (1976) Nucleic Acids Res., vol. 3, pp. 343-353.
  - 3. Tomassini, J., Roychoudhury, R., Wu, R., Roberts, R.J., (1978) Nucleic Acids Res., vol. 5. pp. 4055-406

· Source : Klebsiella pneumonia.

• Concentration : 20 Units/uL

Reaction Condition

- 10X AccuCut™ greeN Buffer

Incubate at 37 ℃

Storage Buffer

20 mM 50 mM

pH 7.5, Tris-HCI KCI

1 mM FDTA 10 mM

• Heat inactivation: No.

2-mercaptoethanol

50% Glycerol

## **Quality Control**

#### · Overdigestion Assay:

No nonspecific activity was detected after incubation of 1  $\mu q$  of  $\lambda$  DNA with 50 units of Kpn I for 15 hours.

\* Conditions of low ionic strength, high enzyme concentration, glycerol concentration >5%, or pH >8.0 may result in star activity.

#### Nuclease Contamination Assay :

No altered pattern was detected after incubation of 1 μg of substrate DNA with Kpn I in 50 μL reaction volume with the supplied AccuCut™ buffer overnight.

#### Ligation and Recutting Assay :

This assay is used to test for exonuclease activity that would degrade the termini of restriction fragments. resulting in inhibition of ligation and of subsequent digestion of ligated fragments. After 40-fold overdigestion with Kpn I, 95% of the DNA fragments can be ligated and recut with Kpn I.