### **Product Datasheet**



5'...**CCTAGG**...3' 3'...**GGATCC**...5' Product No : RE1128 : 200u Quantity



Lot **Expiry Date** Concentration

Supplied with 1ml of 10X Buffer V4 1ml of 10X Buffer UB

0.5ml Diluent Viva Buffer A

(BSA included in all Reaction Buffer)

15u/μl

Store at -20°C



info@vivantechnologies.com

λDNA

(Hind III Digest) 0.7% Agarose

# **Reaction Conditions:**

Buffer V4,

10mM Tris-HCl (pH 8.5 at 30°C), 10mM MgCl<sub>2</sub>, 100mM KCl, and 100μg/ml BSA.

Incubate at 37°C.

Dilution: Viva Buffer A

10mM Tris-HCI (pH 7.4 at 25°C), 50mM KCI, 0.1mM EDTA,

1mM DTT, 200µg/ml BSA and 50% glycerol.

Thermal Inactivation: 80°C for 20 minutes

## Storage Buffer:

10mM Tris-HCl (pH 7.5), 100mM NaCl, 0.1mM EDTA, 100μg/ml BSA, 7mM 2-mercaptoethanol and 50% glycerol.

### Unit Definition:

1u is defined as the amount of enzyme that is required to digest 1μg of DNA in 1 hour at 37°C in 50µl of assay buffer.

# Quality Control Assays:

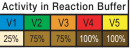
# Ligation/ Recutting Assay:

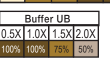
After 15-fold overdigestion with AspA2 I, more than 90% of the DNA fragments can be ligated and recut.

## Overdigestion assay:

An unaltered banding pattern was observed after 1µg of DNA was digested with 30u of AspA2 I for 16 hours at 37°C.







\* Buffer UB is provided for double digestion purpose.

#### NOTE:

- \* Total reaction volume dependent on experiment.
- $^{\star}\,$  The amount of enzyme to be used is very much dependent on the DNA template.
- For plasmid DNA, 5-10X more enzyme is required.

## **Example of Digestion Reaction**

1 unit Enzyme

Lambda (Hind III Digest) 0.3μg/μl : 3.33µl (1µg DNA)

10X Reaction Buffer : 5µl

Sterile Distilled Water : Up to 50µl

> Product Use Limitation This product is for research purposes and in vitro use only.