V *i* V a n t *i* S RESTRICTION ENDONUCLEASE

Product Datasheet



Reaction Conditions:

Buffer V5

30mM Tris-acetate (pH 7.9 at 30°C), 10mM Mg-acetate, 60mM K-acetate, and 100µg/ml BSA. Incubate at 50°C.

Dilution: Viva Buffer A

10mM Tris-HCl (pH 7.4 at 25°C), 50mM KCl, 0.1mM EDTA, 1mM DTT, 200 $\mu g/ml$ BSA and 50% glycerol.

Thermal Inactivation: None

Storage Buffer:

10mM Tris-HCI (pH 7.5), 100mM NaCl, 0.1mM EDTA, 7mM 2-mercaptoethanol,100µg/ml BSA 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 50°C in 50µl of assay buffer.

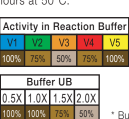
Quality Control Assays:

Ligation/ Recutting Assay:

After 50-fold overdigestion with **BstHH 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 100u of **BstHH I** for 16 hours at 50°C.



* Buffer UB is provided for double digestion purpose.

NOTE:

- * Blocked by CpG-methylation.
- * Total reaction volume dependent on experiment.
- * 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		
Enzyme	:	1 unit
Lambda 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. V \vec{l} V a n t \vec{l} S | www.vivantechnologies.com

λDNA

1.0% Agarose

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