# **AllBio Sal I-Fast Cut**



Cat. No. ABTGRES301

Storage: at -20°C for two years Concentration: 20,000 units/ml Recognition Site

5'...GTCGAC...3' 3'...CAGCTG...5'

### Description

AllBio Sal I-Fast Cut is expressed and purified from *E.coli* that carries the recombinant Sal I gene. The molecular weight is 36.3 kDa, with the recognition site at G^TCGAC The reaction is conducted for 5-15 minutes at 37°C, and heat-inactivated at 65°C for 20 minutes. This enzyme is not sensitive to dam, dcm, but sensitive to mammalian CpG methylation.

#### **Enzyme Properties**

Fast digestion in 5-15 minutes with high fidelity

#### Application

Genomic DNA, plasmid DNA, PCR product

#### Kit Contents

Component	ABTGRES301-01	ABTGRES301-02
AllBio Sal I-Fast Cut	1,000 units	2,000 units
10×FastCut Buffer	250 ul	500 ul
10×DNA Loading Buffer	1 ml	1 ml

#### Unit Definition

One unit is defined as the amount of enzyme required to digest 1 µg of  $\lambda$  DNA in 1 hour at 37°C in a total reaction volume of 50 µl.

#### Quality Control

**Ligation and re-cutting:** After 10-fold overdigestion with AllBio Sal I, more than 95% of the DNA fragments can be ligated with T4 DNA ligase at 25°C. Of these ligated fragments, more than 95% can be recut.

**16-Hour incubation:** A 50 μl reaction containing 1 μg of DNA and 10 units of enzyme incubated for 16 hours results in the same pattern of DNA bands as a reaction incubated for 1 hour with 1 unit of enzyme.

Blue/White screening (Terminal integrity): A DNA vector is digested at a unique site within  $lacZ\alpha$  gene with a 10-fold excess of enzyme, and then ligated, transformed and plated on X-gal/IPTG plate. Successful expression of the  $\beta$ -galactosidase indicates that  $lacZ\alpha$  gene remains integrity after cloning. A blue colony represents an intact gene, and a white colony represents an interrupted gene. To be Blue/White certified, enzymes must produce fewer than 3% white colonies.

**Exonuclease activity:** After incubation for 4 hours at 37°C, a 50  $\mu$ l reaction containing 100 units of enzyme and 1  $\mu$ g <sup>3</sup>H DNA releases less than 0.1% radioactive substance.

**Endonuclease activity:** After incubation for 4 hours at 37°C, a 50 μl reaction containing 15 units of enzyme with 1 μg pBR322 RFI DNA results in less than 10% conversion from RFI to RFII.

#### Storage Buffer

20 mM Tris-HCl pH7.4, 250 mM NaCl, 0.1 mM EDTA, 1.5 mM DTT, 400 μg/ml BSA, 50% Glycerol

#### 10×FastCut Buffer

500 mM Tris-Ac pH7.9, 1 M KAc, 120 mM MgAc,, 1 mg/ml BSA

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## **Reaction Components**

Component	20 μl Volume	50 μl Volume
DNA	≤1 μg	1-2 μg
10×FastCut Buffer	2 µl	5 μl
AllBio Sal I-Fast Cut	0.5 μl	1 μ1
ddH <sub>2</sub> O	to 20 μl	to 50 μl

Prior to use, please completely mix the FastCut Buffer. Increase the volume of enzyme, in case of digestion of  $>2 \mu g$  DNA or incomplete digestion, but the total volume of enzyme should be less than 1/10 of the reaction system.

Incubation for 5-15 minutes at 37°C, Enzyme is inactivated by adding 10×DNA Loading Buffer to a final concentration at 1×, or by heating at 65°C for 20 minutes.

#### Notes

- Thaw the 10×FastCut Buffer completely and mix well before use.
- · Low ionic strength, high enzyme concentration, glycerol concentration > 5%, or pH > 8.0 may result in star activity.

FOR RESEARCH USE ONLY

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