

Lambda DNA BshFI digest, 1.4 % agarose



5'...GG[▼]CC...3' 3'...CC_▲GG...5'

Content:

Ref No.	250110S	color
BshFl 10 U/μL	7000 units	blue
10x buffer A*	1x 1 mL	red
10x buffer K	1x 1 mL	yellow
Datasheet		

We recommend the use of buffer K as universal buffer (BSA included).

Storage: -20 °C

Concentration: 10 U/µL

Source: BshFl is a restriction enzyme purified from *Bacillus sphaericus*.

Enzyme Properties:

1x buffer A composition: 20 mM Tris-acetate (pH 7.9 at 25 °C), 50 mM potassium acetate,

10 mM magnesium acetate, 1 mM Dithiothreitol.

General reaction mixture: 10 U BshFl 1 μL

 $\begin{array}{lll} \text{10x buffer A* or K} & \text{2 } \mu\text{L} \\ \text{DNA substrate} & \text{<1 } \mu\text{g} \\ \text{Sterile ultrapure water} & \text{Up to 20 } \mu\text{L} \\ \end{array}$

Incubate for 15 min at 37 °C

Heat inactivation: 80 °C for 20 minutes.

Methylation Sensitivity: dam methylation: Not sensitive

dcm methylation: Not sensitive CpG methylation: Not sensitive

Storage buffer: 10 mM Tris-HCl (pH 7.4), 50 mM KCl, 0.1 mM EDTA, 1 mM dithiothreitol,

200 μg/ml BSA and 50 % glycerol. Store at -20 °C

Absence of contaminants: 200 units of BshFl do not produce any unspecific cleavage products after

16 hrs incubation with 1 μg of Lambda DNA at 37 °C. After 50-fold overdigestion with BshFI, greater than 95 % of the DNA fragments can be

ligated and recut with this enzyme.

Unit definition:One unit is defined as the amount of enzyme required to produce a complete

digest of 1 μg Lambda DNA (dam) in 60 minutes in a total reaction volume of

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15

22

0.05 mL under assay conditions.

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λ Ad-2 Φx174 pUC18 M13mp18 pBR322
Frequency of Cutting:

216

Percent Activity in BIORON Buffers:

L* M* H* SH* A* K

50-75 75-100 75 50-75 100 100

*we recommend the addition of BSA to a final concentration of 100 µg/mL.

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