ZINZYME

Tinzyme Co., Limited

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Bell

Product Number: BC01

Shipping and Storage

Stored at -20°C, valid for two years.

Components

Component	BC01
	125µl
BelI	125µl
10×Cut Buffer	1ml
Easy-Load 10×Cut Buffer	1ml

Description

BcII is a high-quality restriction endonuclease that has been genetically engineered and can quickly complete DNA cleavage using only one buffer within 5-15 minutes. Suitable for rapid enzymatic digestion of plasmid DNA, PCR products, or genomic DNA.

- 1. **Enzyme activity detection:** At the optimal reaction temperature, in a 20μl reaction system, 1μl of BcII can completely digest 1μg of λDNA(Dam-) within 15 minutes.
- Long term enzyme digestion detection: Incubate 1μl BclI with 1μg λDNA(Dam-) for 3 hours at the optimal reaction
 temperature, and no non-specific degradation of the substrate caused by other nucleases contamination or star activity was
 detected. Delayed enzyme digestion may result in star activity.
- 3. **Enzyme digestion ligation re digestion detection:** At the optimal reaction temperature, use 1µl BclI to digest the substrate, recover the enzyme digestion product, and use an appropriate amount of T4 DNA Ligase at 22 °C to reconnect the enzyme digestion product. After recovering the ligation product again, use the same endonuclease to cleave the ligation product again.

Basic information

Recognition	Isoschizomer	Enzyme digestion	Deactivation	Methylation
sequence		temperature	conditions	interference?
5'-T^GATCA-3'	BsiQI, FbaI,	37°C	80°C 20min	Sometimes
3'-ACTAG^T -5'	Ksp22I			

The activity (buffer compatibility) in different reaction buffers is as follows:

10×Cut	Easy-Load 10×Cut	Thermo FastDigest	NEB CutSmart®	Takara QuickCut™
Buffer	Buffer	Buffer	Buffer	Buffer
100%	100%	100%	100%	100%

$\label{lem:please refer to the table below for the methylation effects of BcII \ recognition \ sites:$

Dam	Dcm	CpG	EcoKI	EcoBI
completely overlaps – blocked.	No effect	No effect	No effect	may overlap – blocked.

Features

- 1. Enzymatic cleavage can be completed within 5-15 minutes;
- 2. All endonucleases share a single enzyme digestion buffer, Cut Buffer, greatly simplifying the enzyme digestion reaction system and facilitating double or multiple enzyme digestion;
- 3. In response to the issue of differences in activity of different enzymes in Cut Buffer, the concentrations of different enzymes were adjusted to uniformly add 1µl of enzyme per 20µl of system for enzyme digestion reaction;
- 4. Many modifying enzymes, such as Alkaline Phosphatase, Antarctic Phosphatase, T4 DNA Ligase, T4 Polynucleotide Kinase, T4 PNK (3 'phosphatase minus), etc., are 100% compatible with Cut Buffer, making reaction systems such as "enzyme cut connect" and "enzyme cut modify connect" compatible and supporting single tube reactions;

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5. Good enzyme activity redundancy makes it easy to cope with substrate excess or difficult template enzyme digestion.

Note

- 1. Endonucleases should be stored in an ice box or on an ice bath during use, and should be immediately stored at -20°C after use.
- 2. If it is found that the expected enzyme cleavage site cannot be cleaved, please confirm whether there is methylation interference.
- 3. Homolytic enzymes may have different sensitivities to different methylation modifications, and when encountering potential methylation interference issues, they can be attempted.
- 4. This product is only for scientific research by professionals and should not be used for clinical diagnosis or treatment, food or medicine, or stored in ordinary residential areas.
- 5. For your safety and health, please wear lab coats and disposable gloves when operating.

Protocol

1. When performing single enzyme digestion, the following reaction system can be referred to and operated on an ice bath.

Reagent	Plasmid DNA	PCR Product	Genomic DNA
Ultrapure Water	(17-x)µl	(26-x)µl	(40-x)μl
10×Cut Buffer or Easy-Load 10×Cut Buffer	2μ1	$3\mu l$	5μ1
Substrate DNA	$x\mu l(up to 1\mu g)$	$x\mu l(\sim 0.2\mu g)$	xµl(5µg)
BclI	1μl	$1\mu l$	5μ1
Total volume	20μ1	30μ1	50μ1
Incubate at 37°C	15min	15-30min	30-60min

Note: The above reaction system is applicable for enzyme digestion of purified PCR products. The unpurified PCR product has a certain ionic strength and pH, and the amount of $10\times\text{Cut}$ Buffer added can be appropriately reduced to $2\mu\text{L}$. However, due to the simultaneous exonuclease activity of many DNA polymerases, it can affect the cleavage products. Therefore, the following steps require ligation, cloning, and other operations. It is recommended to purify the PCR products before cleavage.

- 1.1. After adding various liquids in sequence according to the table above, use a pipette to gently suck or tap the tube wall to mix (do not vortex), and then centrifuge instantly to settle the liquid to the bottom of the tube.
- 1.2. Incubate at 37°C for 15 minutes (plasmid), 15-30 minutes (PCR product), or 30-60 minutes (genomic DNA). It is recommended to use a water bath as the preferred method for enzyme digestion reactions, as the reaction temperature is usually more constant.
- 1.3. Incubate at 80°C for 20 minutes to inactivate the enzyme and stop the reaction (optional).
- When performing double or multiple enzyme digestion, the reaction system can be set according to the following principles based on the enzyme digestion reaction system settings in the reference form.
 - 2.1. The dosage of each rapid endonuclease is 1µl, and the reaction system should be appropriately expanded as needed.
 - 2.2. The total volume of all rapid endonucleases must not exceed 1/10 of the total reaction system.
 - 2.3. If the optimal reaction temperatures for the several rapid endonucleases used are different, the enzyme with the lower optimal temperature should be used first for digestion, and then the enzyme with the higher optimal temperature should be added for digestion reaction at its optimal reaction temperature.

Related products

Product Number	Product Name
APA01	ApaLI
AS01	AscI
AV01	AvrII
BAM01	BamHI



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BC01	BelI	
BG01	BglII	
BSA01	BsaI	
BSN01	BspQI	
BSN02	BstBI	
BSN03	BstEII	
CF01	Cfr9I	
CL01	ClaI	
DPN01	DpnI	
DPN02	DpnII	
DRA01	DraI	
EA01	EagI	
EA02	EarI	
EC02	EcoRI	
EC01	EcoRV	
FSP01	FspI	
HID01	HindIII	
HIN01	HinfI	
HPA01	HpaI	
KAS01	KasI	
KPN01	KpnI	
ML01	MluI	
RS02	MnlI	
MSP01	MspI	
NC01	NcoI	
ND01	NdeI	
NH01	NheI	
NO01	NotI	
NR01	NruI	
NS01	NsiI	
PA01	PacI	
PM01	PmeI	
PS01	PstI	
PV01	PvuII	
SA02	SacI	
SA05	SacII	
SA03	SalI	
SA04	SapI	
SB02	SbfI	
SC02	ScaI	
SF01	SfiI	
SM01	SmaI	
SP01	SpeI	
SP02	SphI	



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SSP01	SspI
STE01	StuI
TAQ01	TaqI
XB01	XbaI
XC01	XcmI
XH01	XhoI
XM01	XmaI