EXact-Cut™ Pmel Restriction Endonuclease

Catalog Number: EXNA053

Size: 500 Units

For Research Use Only. Not Intended for Diagnostic or Therapeutic Use.

EXact-Cut™ PmeI (10 Units/µL)

EXact-Cut™ 10X Buffer

6X Gel Loading Dye, Purple



Product Details				
Description	EXact-Cut™ Pmel Restriction Endonuclease is engineered for high specificity, reduced star activity, and time saving DNA digestion in 5-15 minutes. To simplify experimental design using multiple restriction enzymes, our entire range of EXact-Cut™ restriction endonucleases are 100% active in or EXact-Cut™ buffer (included) and are optimized for single-tube reactions along with digestion and ligation protocols.			
Restriction Enzyme Site	5'GTTT↓AAAC3' 3'CAAA↑TTTG5' Isoschizomers: MssI (Isoschizomers may have different methylation sensitivities)			
Unit Definition	One unit is defined as the amount of enzyme required to digest 1 μg of λ DNA in 1 hour at 37°C in a total reaction volume of 50 μL .			
Recommended Reaction Conditions	1X EXact-Cut™ Buffer Incubate at 37°C Refer to Protocol for reaction setup			
Heat Inactivation	 Incubate at 80°C for 20 minutes Add appropriate volume of 6X Gel Loading Dye, according to the reaction system 			

Storage and Preparation			
Shipping	Shipped on blue ice.		
Stability and Storage	Store at -20°C for up to 24 months.		

500 Units

1 mL

1 mL

Protocol

Components

Protocol for Rapid DNA Digestion

1. Add the following components on ice in the indicated order:

	Plasmid DNA	PCR Product	Genomic DNA
DNA	≤ 1 µg	≤ 0.2 µg	≤ 5 µg
EXact-Cut™ 10X Buffer	2 μL	3 µL*	5 μL
ddH ₂ O, make up to final volume indicated:	20 μL	30 μL	50 μL
Exact-Cut™ Pmel	10 Units	10 Units	50 Units

^{*}This system is suitable for enzymatic digestion of purified PCR products. Unpurified PCR products have a certain ion strength. The amount of 10 × Buffer added can be appropriately reduced to 2 µL. However, due to the fact that DNA polymerase also has exonuclease activity, it can affect the cleavage products. Therefore, the following steps require cloning and other operations. It is recommended to purify the PCR products before cleavage.

Protocol is continued on page 2.



Protocol (continued)

Protocol for Rapid DNA Digestion (continued)

- 3. Incubate at 37°C for the indicated sample type: plasmid DNA (15 minutes), PCR product (15-30 minutes), or genomic DNA (30-60 minutes)
- 4. Optional: inactivate the enzyme at 80°C for 20 minutes and add an appropriate amount of 6X Gel Loading Dye, according to the reaction system.

Protocol for Multiple Digestion of DNA

- 1. Use 10 Units of each enzyme and scale up to the reaction conditions accordingly.
- 2. The combined volume of the enzymes in the reaction mixture **should not** exceed **1/10** of the total reaction volume.
- 3. If the enzymes require different reaction temperatures, start with the enzyme requiring the lowest temperature, followed by the next enzyme(s) and incubate at the higher temperature.

Note: For total reaction volumes > 20 µL, the incubation time should be increased accordingly in a water bath.

Number of Recognition Sites in DNA												
λDNA	ФХ174	pBR322	pUC57	pUC18/19	SV40	M13mp18/19	Adeno2					
2	0	0	0	0	0	0	1					
Methylation Effects on Digestion												
Dam		Dcm	CpG		EcoKI	EcoBI						
No effect	o effect No effect Blocked		cked	No effect	No effect							
Activity in Common Buffers												
		EXact-Cut™ Buffer		Takara QuickCut™ Buffer			NEB CutSmart® Buffer					
Activity		100%	% 100%		50%	100%						
Application N	otes											
Functional Test												
Digestion-Ligation	on Test	At the optimal reaction temperature, the DNA was digested using 10 Units of EXact-Cut Pmel and the digestion product was recovered. The DNA fragments were ligated using an appropriate amount of T4 DNA Ligase at 22°C. After the ligation product was recovered, it was able to be recut with EXact-Cut Pmel.										
Blue/White Scree Assay	ening	An appropriate vector containing the <i>lacZ</i> gene was digested using 10 Units of EXact-Cut Pmel. The digested product was ligated and transformed into <i>E.coli</i> cells plated on plates with X-Gal, IPTG and appropriate antibiotic. The successfully ligated <i>lacZ</i> gene expresses beta-galactosidase and gives rise to a blue colony, while an interrupted gene (due to degraded DNA end) gives rise to a white colony. Less than 1% white colonies were observed.										

