EXact-Cut[™] Nrul Restriction Endonuclease

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

Catalog Number: EXNA028 Size: 500 Units



More information: info@exreprotein.com

Product Details Description EXact-Cut™ Nrul 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 enzvmes, our entire range of EXact-Cut™ restriction endonucleases are 100% active in our EXact-Cut™ buffer (included) and are optimized for single-tube reactions along with digestion and ligation protocols. **Restriction Enzyme Site** 5'...TCG ↓ CGA...3' 3'...AGC ↑ GCT...5' Isoschizomers: Bsp68I, BtuMI, Rrul (Isoschizomers may have different methylation sensitivities) Unit Definition One unit is defined as the amount of enzyme required to digest 1 μ g of λ DNA in 1 hour at 37°C in a total reaction volume of 50 µL. Recommended 1X EXact-Cut[™] Buffer **Reaction Conditions** Incubate at 37°C Refer to Protocol for reaction setup Heat Inactivation This enzyme cannot be heat inactivated. Please purify the digested product by phenol/chloroform treatment or a column-based purification kit. EXact-Cut[™] Nrul (10 Units/µL) Components 500 Units EXact-Cut™ 10X Buffer 1 mL 6X Gel Loading Dye, Purple 1 mL Storage and Preparation Shipped on blue ice. Shipping Stability and Storage Store at -20°C for up to 24 months.

Protocol

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_2O , make up to final volume indicated:	20 µL	30 µL	50 µL
Exact-Cut™ Nrul	10 Units	10 Units	30-50 Units

Note: DNA should be free of phenol, chloroform, ethanol, EDTA, detergents or high concentrations of salts. For compatibility with other common buffers, see the chart on page 2.

- 2. Gently mix or flick the tube to mix (do not vortex), then immediately follow with a quick spin-down in a microcentrifuge.
- 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: this enzyme cannot be heat inactivated. 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.



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Number of Rec	ognition	Sites in DNA							
λDNA	ФХ174	pBR322	pUC57	pUC18/19	SV40	M13mp18/19	Adeno2		
5	2	1	1	0	0	0	5		
Methylation Eff	ects on I	Digestion							
Dam		Dcm	С	CpG		I	EcoBI		
Blocked		No effect	Blo	Blocked		N	No effect		
Activity in Com	mon Bu	ffers							
		EXact-Cut™ Buffer		Takara QuickCut™ Buffer			NEB CutSmart® Buffer		
Activity		100%	10	100%			100%		
Application Not	tes								
Functional Test	i	A 20 μ L reaction in EXact-Cut Buffer containing 1 μ g of λ DNA (Dam) and 10 Units of EXact-Cut Nrul incubated for 15 minutes at 37°C results in complete digestion as determined by agarose gel electrophoresis.							
Digestion-Ligation	c T	At the optimal reaction temperature, the DNA was digested using 10 Units of EXact-Cut Nrul 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 Nrul.							
Non-Specific Endonuclease Acti Test	i vity v	At the optimal reaction temperature,10 Units of EXact-Cut NruI was incubated in 20 μ L reaction volume in EXact-Cut Buffer with 1 μ g of supercoiled plasmid DNA for 4 hours. Undigested, supercoiled plasmid DNA was detected using agarose gel electrophoresis.							



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