

Tinzyme Co., Limited

Email: sales@tinzyme.com Website: www.tinzyme.com Tel: +86-755-86134126 WhatsApp/Facebook/Twitter: +86-189-22896756

FlashCut[™] MunI (MfeI)

Product Number: RE0547

Shipping and Storage

-20°C.

Components

Component	Specifications
FlashCut [™] MunI (MfeI)	25µL
10× CutOne™ Buffer	2×1mL
10× CutOne™ Color Buffer	2×1mL

Description

FlashCutTM Rapid endonucleases are a series of genetically engineered restriction endonucleases that are suitable for rapid enzymatic cleavage of plasmid DNA, PCR products, or genomic DNA. All FlashCutTM Rapid endonucleases have excellent activity in both CutOne TM and CutOne TM Color Buffer, and can complete enzyme cleavage within 5-15 minutes. In addition, Baishimei dephosphorylation and ligation reagents are available on CutOne TM Buffer has 100% activity and supports one tube reaction, enhancing the experience of "enzyme digestion modification connection".

CutOne[™] Color Buffer includes red and yellow tracer dyes, which can be directly used for gel electrophoresis. The migration rate of red dye of CutOne[™] Color Buffer and 2500 bp double stranded DNA fragment in 1% agarose gel is similar; The migration rate of yellow dye and 10 bp double stranded DNA fragment in 1% agarose gel is similar.

Suggested reaction conditions

- 1. $1 \times CutOne^{TM}$ Buffer solution.
- 2. Incubate at 37°C.
- 3. Prepare the reaction system according to the "DNA rapid enzyme digestion process".

Inactivation conditions

Cannot be thermally deactivated, please use phenol chloroform extraction or column purification.

Quality control

- 1. Functional activity detection: At 37°C, in a 20µL universal CutOne[™] reaction system, 1µL FlashCut[™] MunI (MfeI) can completely digest 1µg λ DNA within 15 minutes.
- In the 20µL universal CutOne[™] reaction system, 1µL FlashCut[™] MunI (MfeI) was incubated with 1µg λ DNA for 3 hours, and no non-specific degradation of the substrate caused by other nuclease contamination or star activity was detected. Longer enzyme digestion may result in star activity.
- 3. Enzyme digestion ligation re digestion detection:At 37°C, use FlashCut[™] MunI (MfeI) with 10 times the enzyme amount digests DNA substrates, recovers enzyme cleavage products, and at 22°C, T4 DNA Ligase (Fast) can reconnect over 95% of the enzyme cleavage products. After recycling the connecting product again, more than 95% of the connecting product can be re cut using the same endonuclease.

Icon annotation

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5'...CAATTG...3' 3'...GTTAAC...5'

🗲 37 🖉 🛨

- 1. 📝 Rapid endonuclease can complete the reaction within 5-15 minutes.
- 2. 37 The optimal reaction temperature is 37° C.
- 3. Cannot be thermally deactivated.
- 4. 😿 3 hours of incubation did not show star activity, and longer enzyme digestion may result in star activity.

Protocol

1. DNA rapid enzymatic digestion process

1.1. Prepare the reaction system on ice according to the recommended sample addition sequence as follows:

	plasmid DNA	PCR products	Genomic DNA
ddH ₂ O	15µL	16µL	30µL
10×CutOne [™] Buffer or 10×CutOne [™]	2µL	3µL	5µL
Color Buffer			
Substrate DNA	2µL(up to 1µg)	10µL(~0.2µg)	10µL(5µg)
FlashCut™ MunI (MfeI)	1µL	1µL	5µL
Total	20µL	30µL	50µL

This system is suitable for enzyme digestion of purified PCR products. The unpurified PCR product has a certain ionic strength, $10 \times \text{CutOne}^{TM}$ The amount of buffer added can be appropriately reduced to 2μ L. However, due to the simultaneous exonuclease activity of DNA polymerase, it can affect the cleavage products. Therefore, the following steps require cloning and other operations. It is recommended to purify the PCR products before enzyme cleavage.

- 1.2. Gently suck or tap the tube wall to mix well (without vortexing), then centrifuge instantly to collect hanging wall droplets.
- 1.3. Incubate at 37°C for 15 minutes (plasmid), or 15-30 minutes (PCR product), or 30-60 minutes (genomic DNA).
- 1.4. Phenol chloroform extraction or column purification (optional).
- 1.5. If using CutOne[™] Color Buffer is used for enzyme digestion reaction, and the resulting product can be directly subjected to sample electrophoresis.

2. Double enzyme digestion or multi enzyme digestion

- 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 enzyme digestion, and then the enzyme with the higher optimal temperature should be added for enzyme digestion reaction at its optimal reaction temperature.

3. Expansion reaction system suitable for plasmids

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DNA	1µg	2µg	3µg	4µg	5µg
FlashCut™ MunI (MfeI)	1µL	$2\mu L$	3µL	4µL	5µL
10×CutOne [™] Buffer or 10×CutOne [™]	1µL	2µL	3µL	4µL	5µL
Color Buffer					
Total	20µL	20µL	30µL	40µL	50µL

Note: If the total reaction system is greater than $20\mu L$, the incubation time should be appropriately increased, and water bath, metal bath or sand bath should be used as much as possible.

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The number of enzyme cleavage sites in different DNA

λDNA	ФХ174	pBR322	pUC57	pUC18/19	SV40	M13mp18/19	Adeno2
8	1	0	0	0	4	0	4

The influence of methylation modification

Dam	Dcm	CpG	EcoKI	EcoBI
no effect				

Activity in different reaction buffers

	CutOne™	Thermo Scientific	NEB	Takara
	Buffer	FastDigest Buffer	CutSmart® Buffer	QuickCut [™] Buffer
reactivity	100%	12.5%	100%	12.5%

Note: The activity data comes from the detection under the restriction enzyme standard reaction system.

The activity of DNA modifying enzymes in CutOne[™] Buffer and CutOne[™] Color Buffer

Alkaline Phosphatase (Fast)	100%
T4 DNA Ligase (Fast)	100%

Note: The activity data comes from the detection under standard reaction system, and T4 DNA Ligase (Fast) requires ATP as a cofactor.