

AapCas12b Nuclease Instruction Manual

Product Number	AapCas12b Nuclease	Product Number	EDE0006
Molecular Weight 130.5 KDa		Form	Liquid

I. Product Description

AapCas12b nuclease (also known as C2c1) is derived from Alicyclobacillus acidophilus. It is a crRNA-mediated nuclease that recognizes and cleaves double-stranded DNA (dsDNA) in the presence of a PAM (Protospacer Adjacent Motif) sequence (TTN). For single-stranded DNA (ssDNA) cleavage, the PAM sequence is not required. Upon binding to complementary ssDNA or dsDNA, the trans-cleavage activity of AapCas12b on non-specific ssDNA is activated. AapCas12b has an optimal reaction temperature of 60°C, making it more heat-resistant compared to Cas13a and Cas12a. This property makes it highly suitable for field nucleic acid detection applications when combined with loop-mediated isothermal amplification (LAMP).

II. Product Information

·Product Components

Component	EDE0006-50	EDE0006-500	EDE0006-1000
AapCas12b Nuclease	5 μM*10 μL (50pmol)	5 μM*100 μL (500 pmol)	5 μM*200 μL (1000 pmol)
AapCas12b Cleavage Buffer (10×)	40 µL*1 tube	400 µL*1 tube	800 μL*1 tube

Storage Conditions and Shelf Life

The product is stable for 1 year when stored at -20°C. For long-term storage, it is recommended to store at -80°C. It is advised to aliquot the product based on the frequency of use to avoid repeated freeze-thaw cycles.

Product Features

The product is prepared using a one-step purification process, retaining maximum enzymatic activity. It has been tested to show significantly higher activity compared to similar products.

Activity Definition

In a 20 µL reaction system at 60°C, the amount of Cas12b enzyme required to cleave 1 pmol of ssDNA probe within 1 minute defines 1 transU. For example, if a batch of AapCas12b enzyme exhibits trans-cleavage activity of 30 transU/pmol, it indicates that 1 pmol of this batch of AapCas12b enzyme can cleave 30 pmol of ssDNA probe under the specified reaction conditions within 1 minute.

Quality Assurance

Sample Purity: ~95% (verified by SDS-PAGE).

III. Detection Steps

Required Reagents:

1. ssDNA reporter probe (labeled with FAM at the 5' end and BHQ1 at the 3' end). Note: You can use the ssDNA reporter from our company or design your own.

Product Name	Product Number	
ssDNA reporter (DNA probe)	EDD0001	

2. crRNA/gRNA: Forms a functional complex with Cas12b, which is specifically activated by the target sequence.

Note: Our company offers crRNA synthesis or you can custom-design crRNA.

Name	Specification	Product Number
crRNA Biosynthesis	20D	EDR0002
crRNA Chemical Synthesis	20D	EDR0003

For free design consultation, please contact: info@editxor.com

3. Isothermal Amplification Reaction Kit

Reaction System

Component	Final Concentration	Volume (µL)
10× Cleavage Buffer	1×	$2 \mu L$
5 μM AapCas12b Nuclease	150-250 nM	0.8 μL (200 nM)
500 nM crRNA	150-250 nM	8 μL (200 nM)
$4 \mu M$ ssDNA Reporter	150-250 nM	1 μL (200 nM)
$1 \mu M$ DNA target	150-250 nM	4 μL(200 nM)
DEPC H ₂ O		
Total	20 µL	20 µL

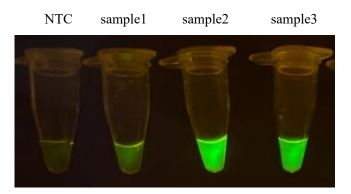
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Reaction Conditions:

Use a real-time fluorescence quantitative PCR machine or an isothermal amplification instrument to detect the fluorescence signal. The reaction should occur at 60°C, and the fluorescence signal should be collected every 30 seconds. Alternatively, the fluorescence signal can be directly observed under a UV light.

Photograph Example



Precautions

1. To prevent contamination by RNase, please keep the experimental area clean and tidy. Wear clean gloves and masks during operations. All consumables, such as pipette tips and centrifuge tubes, should be RNase-free.

2. Cas12 enzymes are prone to inactivation; store the enzyme at -20°C immediately after use.

Publishing Requirements

When using this product in publications, please acknowledge our company: Guangzhou Editgene Co. Ltd, China. Or EDITGENE CO.LTD if used within U.S. or Europe territory.