

*n*Tag-Tenuto

 Cat.#
 Size
 Size

 P225A
 250 units
 5 units/µl

 P250A
 500 units
 5 units/µl

Store at -20℃

Supplied with: 10X nTaq-Tenuto Buffer (Mg²⁺ plus)

dNTP Mixture (2 mM each)

GC Melt I GC Melt II Sterile water

India Contact:

Life Technologies (India) Pvt. Ltd.

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Product description

nTaq-Tenuto is nTaq (Cat.# P025, P050) supplemented with 3' \rightarrow 5' proofreading activity and a PCR enhancing factor for improved efficiency and fidelity. nTaq-Tenuto DNA Polymerase can be used to amplify DNA longer than 10 kb, which is difficult with common Taq polymerases alone. Thus, this product is improved in both fidelity (> 2 fold) of PCR products and amplification efficiency of longer PCR products.

Characteristics

- Molecular weight: 94 kDa
- Error rate: 3.0 X 10⁻⁶
- Thermal stability: Half life of 40 min at 95℃
- A-tail formation at 3' ends of amplified DNA products.

Applications

- Amplification of long DNA fragments (>5~15 kb)
- Amplification of high-complexity template DNA such as cDNA and genomic DNA
- Primer extension
- Colony PCR
- Multiplex PCR
- Labeling of DNA fragments with radioactive-isotopes
- Nucleotide sequencing

For Research Use Only, Not for use in diagnostic procedures,

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with common Tag polymerases alone. Thus, this product is

improved in both fidelity (> 2 fold) of PCR products and

ISO9001 [ISO14001] ISO13485

Product description

Quality control

- Purity: >99% on SDS-PAGE
- Endonuclease-free
- Exonuclease-free
- RNase free
- Inhibitor-free

Unit definition

One unit is defined as the amount of enzyme required to incorporate 10 nmol of dNTP into acid insoluble materials in 30 min at 74°C in a 50-µl reaction mixture (20 mM Tris-HCl/pH 8.8, 50 mM KCl, 2.5 mM MgCl2, 10 mM 8-mercaptoethanol, 12.5 µg of calf thymus DNA),

Storage buffer

20 mM Tris-HCl (pH 7.9), 100 mM KCl, 0.1 mM EDTA, 1 mM DTT, 0.5% NP-40, 0.5% Tween-20, 50% glycerol,

10X nTaq-Tenuto Buffer (Mg²⁺ plus)

Mg²⁺ plus buffer: Containing 20 mM Mg²⁺

Life Technologies™

Your Molecular & Cell Technology Partner

Standard PCR conditions

- PCR mixture^a

10X nlag-lenuto Buffer (Mg ²⁺ plus)	2 µl
nTaq-Tenuto DNA Polymeraseb (5 units/µl)	0.2 µl
dNTP mixture (2 mM each, final conc., 200 μM each)	2 µl
Template DNA ^c (0.1~500 ng/µl)	1 µl
Primer 1 (5 pmole/µl)	1 µl
Primer 2 (5 pmole/µl)	1 µl
Sterile water	up to 20 μl
^a Assemble the reaction mixture on ice	
bAdd the PCR polymerase at the final step	
cPlasmid DNA, 0.1 ng~30 ng; genomic DNA,	50 ng~500 ng

PCR cycle

2 min			
Z 111111			
30 sec			
30~60 sec			
1 min/kb			
5 min			
When cycles are over, keep the reaction mixture at 4°C; may add 10 mM EDTA until use to prevent DNA degradation.			
^a Recommended annealing temperatures is 5 to 10℃ below the lower Tm of the two primers used.			



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°Plasmid DNA, 0.1 ng~30 ng; genomic DNA, !	50 ng~500 ng			

- PCR cycle

Initial denaturation	95℃	2 min		
Denaturation	95℃	30 sec		
Annealing ^a	55℃~65℃	30~60 sec		
Elongation	72℃	1 min/kb		
Number of cycles	25~35 times			
Final elongation	72℃	5 min		
When cycles are over, keep the reaction mixture at 4°C; may add 10 mM EDTA until use to prevent DNA degradation.				
^a Recommended annealing temperatures is 5 to 10℃ below				

the lower Tm of the two primers used.