



## **MBT060C**

# Tag Polymerase (Recombinant) (3 units/µl)

### Components

Reagents provided	МВТО6ОС			
	500 Units	1000 Units	5000 Units	20 x 500 Units
Taq Polymerase (3 U/μl)	170 μΙ	335 μl	1.7 ml	3.4ml (20 x 170 μl)
10X HiBuffer A (With 25mM MgCl <sub>2</sub> )	2 ml	4 ml	20 ml	40 ml (20 x 2 ml)
10X HiBuffer S (With 17.5 mM MgCl <sub>2</sub> )	2 ml	4 ml	20 ml	40 ml (20 x 2 ml)
50mM MgCl <sub>2</sub>	1 ml	2 ml	10 ml	20 ml (20 x 1 ml)

## **Description:**

Taq DNA Polymerase is a thermostable DNA polymerase of thermophillic bacterium *Thermus aquaticus*. The enzyme catalyzes 5'-3' synthesis of DNA, has no detectable 3'-5' exonuclease (proof-reading) activity, but maintains the 5'-3' exonuclease activity. In addition, Taq DNA Polymerase exhibits deoxynucleotidyl transferase activity, resulting in addition of extra adenines at 3'-end of PCR products.

#### Features:

- Thermostable enzyme of approximately 94 kDa from Thermus aquaticus
- Ultra pure recombinant protein
- Replicates DNA at 74°C and exhibits a half-life of 40 minutes at 95°C
- Generates mostly 3' dA overhang PCR products which are suitable for TA cloning
- DNA sequencing and labeling
- Incorporates modified nucleotides (EX: biotin, deoxygenin, fluroscently-labelled nucleotides)
- Supplied with two buffers- 10X Hibuffer A with KCl and 10X Hibuffer S with (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> [(NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> allows for PCR at wide range of Mg<sup>+2</sup> concentrations and decrease unspecific priming]

Concentration: 3 U/µl

**Source:** *E.coli* cells with a pol gene from *Thermus aquaticus* 

Molecular weight: 94 kDa monomer

### **Unit Definition:**

1u is defined as amount of enzyme that is required to catalyze the incorporation of 10 nmoles of dNTP into acid-insoluble material in 30 minutes at 74°C.

### **Reaction Buffer:**

## 10X HiBuffer A (With 25mM MgCl<sub>2</sub>):

**500mM KCI**, 100mM Tris-HCl (pH 9.1 at  $20^{\circ}$ C) and 0.1% Triton X -100. The buffer is optimized for use with 0.1-0.2mM of each dNTP.









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#### 10X HiBuffer S:

**160mM (NH<sub>4</sub>)<sub>2</sub>SO**<sub>4</sub>, 500mM Tris-HCl (pH 9.2 at 22°C), **17.5mM MgCl<sub>2</sub>** and 0.1% Triton X-100. The buffer is optimized for use with 0.35mM of each dNTP.

## **Storage Buffer:**

200 mM Tris-HCl (pH 8.0 at 22°C), 100 mM KCl, 0.5% Tween 20, 0.5% Nonidet-P40, 0.1 mM EDTA, 1mM DTT and 50% Glycerol. Store at  $-20^{\circ}$ C

### Guidelines for PCR optimization using HiMedia's Taq Polymerase:

### DNA Template

- 1. Use high quality, purified DNA templates
- 2. Approximately 10<sup>4</sup> copies are required to detect the amplification in 25-30 PCR cycles
- 3. Use higher DNA concentration when few PCR cycles are desired

#### Primers

- 1.Generally 20-30 bp in size
- 2.GC content between 40-60% ideally
- 3.Melting temperatures should be between 42-65°C
- 4. Final concentration to be used 0.1-0.5 

  µM of each primer

## • Magnesium Concentration

- 1. Ideal for Taq Polymerase is 1.5-2.0mM
- 2. Optimum concentration depends on template, buffer and dNTPs
- 3. Higher than optimal concentration yields undesired products and if concentration is too low the concentration, no amplification products are detected

#### dNTPs

- 1. Typical concentration to be used is 200μM
- 2. Higher than optimal concentration of dNTPs yields higher yield but low fidelity

### Taq Polymerase

1. Typical concentration to be used is 0.5 to 2 units per 50µl of reaction

#### PCR reaction

- 1. Thaw all reaction components on ice
- 2. To PCR reaction, add Taq Polymerase at the end
- 3. Once the reaction is set, immediately transfer the tubes to pre-heated thermal cycler.
- 4. Start the reaction with desired cycling conditions with annealing temperature set to 5°C difference of melting temperature between forward and reverse primers.

### **Quality Control:**

All preparations are assayed for contaminating endonuclease, exonuclease, and non-specific DNase activities. Functionally tested in DNA amplification.

### **Buffers recommended for different sizes of template DNA**

Buffers	Size of template DNA to be amplified				
	100bp-5kb	5kb-8kb	8kb-20kb		
HiBuffer S (1X)	-	+	+		
HiBuffer A (1X)	+	-	-		
MgCl <sub>2</sub>	+	+	+		

Key: + Indicates recommended buffer

## **Inhibition and Inactivation:**

- Inhibitors: ionic detergents (deoxycholate, sarkosyl and SDS) at concentrations higher than 0.06, 0.02, and 0.01% respectively.
- Inactivated by phenol/chloroform extraction.

#### NOTE:

The error rate for Taq Polymerase, which lacks proof-reading activity is approximately 1 to  $2 \times 10^{-5}$  errors (or mutation frequency) per nucleotide per duplication. Accordingly, the accuracy of PCR is 4.5  $\times 10^{4}$ . Accuracy is an inverse of the error rate and shows an average number of correct nucleotides incorporated before an error occurs.

**Storage conditions:** The Taq Polymerase should be stored at -20°C. When stored under the recommended conditions, the product is stable for 2 years.

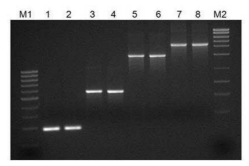


Figure representing amplification of different amplicon sizes using Taq Polymerase with HiBuffer A and HiBuffer S.

### **Technical Assistance**

At HiMedia, we pride ourselves on the quality and availability of our technical support. For any kind of technical assistance, mail at mb@himedialabs.com.

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