

ET190122

# **Taq Plus DNA Polymerase**

Cat. no. 4992769/4992770

Storage: -20°C.

Concentration: 2.5 U/µl

Product size

Product Components	4992769	4992770
Taq Plus DNA Polymerase	250 U	500 U
10x Tag Plus Buffer	1 8 ml	1.8 ml

#### Introduction

Taq Plus DNA Polymerase is a special blend of Taq polymerase and Pfu polymerase. It possesses both 5'-3' and 3'-5' exonuclease activity. The advantages of Taq Plus DNA Polymerase has high productivity and fidelity. Comparing to Taq polymerase, Taq Plus can efficiently amplify large DNA fragments (20 Kb for simple templates and 10 Kb for complex templates). Besides, it possesses higher extension rate and amplification efficiency than Pfu Polymerase.

*Taq* Plus DNA Polymerase generates PCR products with 3'-dA overhangs that can be directly used in TAcloning. To obtain higher cloning efficiency, however, PCR products could be purified and added 3'-dA overhangs before TA cloning procedures.

#### **Unit Definition**

One unit of *Taq* Plus DNA Polymerase is defined as the amount that incorporates 10 nmol of dNTPs into acid-insoluble substrates within 30 min at 74°C with activated salmon sperm DNA as the template/primer.

### **Quality Control**

The purity of SDS-PAGE is more than 99%; No activity of exogenous nuclease is detected; Single gene in human genome could be amplified effectively; No significant activity change when stored at room temperature for one week.

## Storage Buffer

20 mM Tris-HCl (pH 8.0), 0.1 mM EDTA, 1 mM DTT, 100 mM KCl, Stabilizers, 50% Glycerol

# 10× Taq Plus Buffer

200 mM Tris-HCl (pH9.0), 200 mM KCl, 100 mM (NH4) $_2$ SO $_4$ , 15 mM MgCl $_2$  and other components.

# Applications

Amplify DNA fragments from complex templates (e.g. Genome) with high fidelity, for applications such as gene cloning, Site-directed mutagenesis, SNP Analysis etc.

### Example

Note: The following example is only for reference, user must set up optimal reaction system according to different reaction conditions such as different templates or primers etc.

1. For 50 µl PCR reaction system: 1 kb fragment of human genomic DNA was amplified (If use different reaction system, please proportionally increase or decrease the amount of reaction components referring to this system).

Template	< 1 µg
Primer 1 (10 μM)	1 μl
Primer 2 (10 μM)	1 μl
10× Taq Plus Buffer	5 μl
dNTP Mixture (2.5 mM)	4 μΙ
Taq Plus (2.5 U/μl)	0.5-1 μΙ
ddH₂O	up to 50 μ

2. PCR cycle set-up:

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94°C 3 min
94°C 30 sec
55°C 30 sec
72°C 1 min
72°C 5 min
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3. Result detection: Load 5  $\mu l$  PCR products to agarose gel for PCR detecting.

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