

# Prime Taq DNA Polymerase

Cat. No.	Size	Remark	
G-1000	250 Units	with dNTPs mix., +MgCl <sub>2</sub> buffer	

## Package Information

G-1000	- Prime Taq DNA Polymerase (5 Units/#l): 50#l - 10X Reaction buffer (with MgCl <sub>2</sub> ): 1.0 ml - 10 mM dNTP Mixture (2.5 mM of each dNTPs): 0.5 ml
--------	---

#### Description

Prime Taq DNA Polymerase is a high quality recombinant enzyme and catalyzes  $5' \rightarrow 3'$  synthesis of DNA. The enzyme has no detectable  $3' \rightarrow 5'$  proofreading exonuclease activity.

It is provided with 10X reaction buffer that contains PCR enhancers. This reaction buffer will enable or improve sub-optimal PCR caused by templates that have a high degree of secondary structure or that are GC-rich.

### **Buffer and Reagents**

Storage Buffer

20 mM Tris-HCl (pH 8.0), 100 mM KCl, 0.5 mM EDTA,

0.1 mM DTT, 0.5% Tween 20, 50% Glycerol

10X Reaction Buffer (with MgCl<sub>2</sub>)

Contains Tris-HCl (pH 9.0), 20 mM MgCl<sub>2</sub>, (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> and PCR enhancers. 10 mM dNTP mixture

2.5 mM of each dATP, dCTP, dGTP and dTTP

#### **Usage Information**

A DNA fragment which is amplified by Prime Taq DNA Polymerase has A-overhang, and it enables you to do cloning by using T-vectors.

This product is sold for research purpose only. This is not to be used for

human diagnostic or drug purposes.

All claims must be brought within expired date.

#### Protocol

Optimal reaction conditions, such as reaction time, temperature and amount of template DNA, may vary and must be individually determined.

1. Thaw 10X Reaction buffer.

2. Prepare a master mix.

Components	Volume	Final Conc.
10X Reaction buffer	5 <i>µ</i> l	1X
10 mM dNTP Mix. (2.5 mM of each)	1 ~ 5 <i>µl</i>	0.2 ~ 1.0 mM
Upstream Primer	Variable	0.1 ~ 1.0 μM
Downstream Primer	Variable	0.1 ~ 1.0 μM
Prime Taq DNA Pol. (5Unit/µl)	0.6 ~ 1.25 <i>µ</i> l	1.2 ~ 2.5 U
Template DNA	Variable	10 fg~1 <i>µ</i> g
Sterilized D.W	Variable	-
Total Volume	50 <i>µ</i> l	-

# Made by ISO 9001 & ISO 13485 system

- \* Amount of template:
- Bacteriophage lambda, cosmid, plasmid: 10 fg ~ 300 ng
- Total genomic DNA: 100 ng ~ 1 μg

3. Mix the master mix and dispense appropriate volumes into PCR tubes. Centrifuge the reactions in a micro-centrifuge for 10 seconds.

4. Perform PCR using your standard parameters (3-step cycling).

Star	Temp. & Time		Curles
Step	Temp.	Time	Cycles
Initial denaturation	95°C	5 min.	1
Amplification	95℃ 50~60℃ 72℃	30 sec. 30 sec. 30~60 sec.	25 ~ 45
Final extension	72℃	5 min.	1

 $\star$  For PCR products longer than 3~4kb, use an extension time of approximately 1 min per Kb DNA.

5. Separate the PCR products by agarose gel electrophoresis and visualize with EtBr or any other means.

Research Use Only

Store at -20℃

Avoid multiple freeze-thaw cycles and exposure to frequent temperature changes.