

Product Information					
Taq-B DNA Polymerase					
Part Number	P7250L				
Concentration	5,000 U/mL				
Unit Size	10,000 U				
Storage Temperature	-25ºC to -15ºC				
Lot Number					
Reference Number					

Product Specifications P7250L Rev 03

Product Description: Taq-B DNA Polymerase is a thermally stable, processive, $5' \rightarrow 3'$ DNA polymerase. The 94 kDa protein possesses an inherent $5' \rightarrow 3'$ nick-translation moiety and lacks a $3' \rightarrow 5'$ proofreading function.

Product Specifications						
P7030						
Assay SDS Purity	Specific	SS	DS Exonuclease	DS	E. coli DNA	
	SDS Purity	Activity	Exonuclease	DS Exonuclease	Endonuclease	Contamination
Units Tested	n/a	n/a	50	50	50	50
Creation	>99%	74,625 U/mg	<5.0%	<1.0%	No Conversion	<10 copies
Specification			Released	Released		

<u>Source of Protein</u>: A recombinant E. coli strain carrying the Taq DNA polymerase gene from the thermophilic organism Thermus Aquaticus YT-1.

<u>Unit Definition</u>: 1 unit is defined as the amount of enzyme that will incorporate 10 nmol of dNTP into acid-insoluble material in 30 minutes at 75°C.

Molecular weight: 93,910 Daltons

Quality Control Analysis:

Unit Activity is measured using a 2-fold serial dilution method. Dilutions of enzyme were made in 1X reaction buffer and added to 50 µL reactions containing Calf Thymus DNA, 25 mM TAPS (pH 9.3), 50 mM KCl, 2.0mM MgCl2, 1 mM DTT, 3H-dTTP and 100 µM dNTPs. Reactions were incubated 10 minutes at 75°C, plunged on ice, and analyzed using the method of Sambrook and Russell (Molecular Cloning, v3, 2001, pp. A8.25-A8.26).

Protein Concentration (OD280) is determined by OD280 absorbance.

Physical Purity is evaluated by SDS-PAGE of concentrated and diluted enzyme solutions followed by silver stain detection. Purity is assessed by comparing the aggregate mass of contaminant bands in the concentrated sample to the mass of the protein of interest band in the diluted sample.

Single-Stranded Exonuclease is determined in a 50 μ L reaction containing a radiolabeled single-stranded DNA substrate and 10 μ L of enzyme solution incubated for 4 hours at 37°C.

Double-Stranded Exonuclease is determined in a 50 μ l reaction containing a radiolabeled double-stranded DNA substrate and 10 μ L of enzyme solution incubated for 4 hours at 37°C.

Double-Stranded Endonuclease is determined in a 50 μ L reaction containing 0.5 μ g of plasmid DNA and 10 μ L of enzyme solution incubated for 4 hours at 37°C.



E.coli 16S rDNA Contamination is evaluated using 5 µL replicate samples of enzyme solution denatured and screened in a TaqMan qPCR assay for the presence of contaminating *E.coli* genomic DNA using oligonucleotide primers corresponding to the 16S rRNA locus.

<u>Supplied in:</u> 20mM Tris-HCl, 100mM NaCl, 1mM DTT, 0.1mM EDTA, Stabilizer, 50% glycerol pH 7.5 @ 25⁰C. <u>Supplied with:</u>

10X PCR Buffer I (B7030): 100mM Tris-HCl, 500mM KCl, 15mM MgCl₂ pH 8.3 @ 25°C.

Notes:

Taq DNA Polymerase is the original and most commonly used PCR enzyme. Taq excels at amplifying shorter (<5 kb) sequences from lowcomplexity template sources and produces robust yields with little or no optimization of reaction conditions. Consider the following guidelines when designing PCR strategies using Taq DNA Polymerase.

1. **DNA Template:** Although extensive purification of PCR templates is typically not necessary, care should be taken with crude or partially purified DNA sources as handling and chemical agents can adversely affect the PCR process. Exposure to short-wave UV light or other DNA damaging agents should be avoided, as should high ionic strength, detergents such as SDS, loading dyes and phenol. In order to prevent contamination from previous PCR reactions, consider setting up reactions in a positive-pressure hood and with aerosol barrier pipet tips. In a typical 25 cycle PCR, 10⁴ copies of target sequence will yield reproducible amplification product. This corresponds to roughly 0.1-1 ng/ml (final concentration) of plasmid DNA, and 1-10 μg/ml of genomic DNA. The use of lower DNA concentrations typically produces less non-specific product, while higher concentrations can allow for fewer cycles and lower mutation rates.

2. Primer Design: Ideally, oligonucleotide primers are 15-30 bases in length, nearly 50% G+C, and have equal (+/- 3° C) annealing temperatures. The use of software to detect self-complementary or hairpin-prone regions is advised and is offered as a service by some synthesis providers. Note that although the 5'-terminus of the primer may contain untemplated sequence, the 3' end must match perfectly. Typical oligonucleotide concentration in the reaction is 0.1-0.5 μ M.

3. Magnesium: Magnesium is a critical component of the PCR reaction though its concentration can be modulated to promote various effects. Generally, 1.5-2.0 mM Mg2+ is targeted, but higher concentrations (up to 5 mM) may be used to stimulate the yield of reactions at the expense of fidelity. The converse is also true – lower magnesium concentrations will promote higher- fidelity products with a lower overall amplification yield. Note that certain reaction components, in particular template DNA and oligonucleotides, may contribute chelating agents to the reaction which could lower the effective magnesium concentration and starve the reaction.

4. dNTPs: Generally, a final concentration of 100-200 μM dNTPs is employed, though higher concentrations may stimulate yields (particularly with longer targets) and lower may offer increases in fidelity. Taq DNA Polymerase can also incorporate and read through deoxy Uridine and Inosine, two analogs used in certain applications.

5. Taq Polymerase: 1 unit/50 μ L reaction (20 U/mL) is typical, though additional enzyme may be added to stimulate yields. Taq DNA Polymerase extends a DNA template at approximately 2000 nucleotides/minute, so it is recommended that 45-60 seconds of extension time be provided per cycle. Appropriate extension temperatures range from 66-72°C

Usage Instructions:

Typical 50 µL Reaction

On ice, prepare each of following master mixes, combine, and place in heated (to 94°C) thermal cycler:

2X DNA/Oligonucleotide Master Mix:	General Cycling Conditions				
	94°C	3 minutes	Initial Denaturation		
1.0 μL 10 mM dNTPs	25 Cycl	25 Cycles:			
1.0 μL 10 μM Forward Primer	94°C	30 seconds	Denaturation		
1.0 μL 10 μM Reverse Primer	55°C	30 seconds	Annealing		
1.0 μL 500 ng/μL genomic DNA	68°C	30 seconds	500 bp extension		
21 μL Type I Water	68°C	5 minutes	Final Extension		
2X Enzyme/Buffer Master Mix:					

5.0 μL 10X eGrade Reaction Buffer 0.2 μL 5 U/μL Taq DNA Polymerase 19.8 μL Type I Water

Legal Disclaimers:

Patents

Certain applications in which this product can be used may be covered by patents issued and applicable in the United States and abroad. Purchase of this product does not include a license to perform any patented application, therefore it is the sole responsibility of users of this product to determine whether they may be required to engage a license agreement depending upon the particular application in which the product is used.

Limitations of Use

This product was developed, manufactured, and sold for *in vitro* use only. The product is not suitable for administration to humans or animals. SDS sheets relevant to this product are available upon request.

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