



DONGSHENG BIOTECH

Optimus™ Hotstart Taq

DNA Polymerase

#P1041, 250U

Concentration: 5U/μl

Contents:

Optimus™ Hotstart Taq DNA Polymerase (5U/ul)	50 μl
10× Hotstart Buffer (Mg ²⁺ Plus)	1.25 ml
6× Loading Buffer	1 ml

Store at -20°C

For research use only.

In total 3 vials.

Description

Optimus™ Hotstart Taq DNA Polymerase is a hot-start polymerase with chemical modification, which brings higher specificity by reducing non-specific products as the enzyme activity is temperature-dependent and is inhibited at room temperature. The amplification length and speed can reach to 5 kb (simple template) and 2min/kb (simple template up to 20s/kb) separately. Hotstart Taq has 5'-3' polymerase activity, but no 3'-5' exonuclease activity. The product of Hotstart Taq has overhanged dA at 3'-end. Optimus™ Hotstart Taq DNA Polymerase has zero animal source pollution by being produced with advanced chemical modification. And it is much more stable than antibody-modified hot-start polymerase. Its efficiency is higher than most chemical-modified polymerase and the initial-denaturation time can be reduced to 3 minutes.

Optimus™ Hotstart Taq DNA polymerase is an innovative and useful product.

Unit Definition

One unit is defined as the amount of the enzyme required to catalyze the incorporation of 10 nmole of dNTPs into an acid-insoluble form in 30 minutes at 70°C using hering sperm DNA as substrate.

PRODUCT USE LIMITATION.

This product is developed, designed and sold exclusively *for research purposes and in vitro use only*. The product was not tested for use in diagnostics or for drug development, nor is it suitable for administration to humans or animals.

Storage Buffer

20mM Tris-HCl, 1mM DTT, 0.1mM EDTA, 100mM KCl, 0.5% Tween20, 50% (V/V) glycerol, 0.5% Triton X -100

10× Hotstart Buffer with Mg²⁺

50mM KCl, 100mM Tris-HCl, 200mM NH₄Cl, 20mM MgCl₂.

Applications

- High-specificity amplification & multiplex PCR: hot-start polymerase with chemical modification
- Thermostable: half-life over 40 min at 95°C incubation
- Generates 3'-dA overhangs PCR products

Basic PCR Protocol

The following basic protocol serves as a general guideline and a starting point for any PCR amplification. Optimal reaction conditions (incubation time and temperature, concentration of *Optimus*TM Taq DNA Polymerase, primers, Mg²⁺, and template DNA) vary and need to be optimized.

1. Add the following components to a sterile microcentrifuge tube sitting on ice:

1.1 Recommended PCR assay with Hotstart Buffer (Mg²⁺ plus)

Reagent	Quantity	Final concentration
Sterile deionized water	variable	-
10× Hotstart Buffer (Mg ²⁺ plus)	5 µl	1×

dNTPs (10mM each)	1 µl	0.2mM each
Primer I	variable	0.4-1µM
Primer II	variable	0.4-1µM
<i>Optimus</i> TM Taq DNA Polymerase (5U/µl)	0.25-0.5 µl	1.25-2.5U/50 µl
Template DNA	variable	10pg-1µg
Total		50 µl

1.2 Recommended PCR assay with Hotstart Buffer (Mg²⁺ free)

Reagent	Quantity	Final concentration
Sterile deionized water	variable	-
10× Hotstart Buffer (Mg ²⁺ free)	5 µl	1×
dNTPs (10mM each)	1 µl	0.2mM each
Primer I	variable	0.4-1µM
Primer II	variable	0.4-1µM
25mM Mg ²⁺	variable	1.0-4.0mM
<i>Optimus</i> TM Taq DNA Polymerase (5U/µl)	0.25-0.5 µl	1.25-2.5U/50 µl
Template DNA	variable	10pg-1µg
Total		50 µl

Table for selection volume of 25 mM MgCl₂ solution in a 50 µl reaction mix:

Final Mg ²⁺ Conc. (mM)	1.0	1.5	2.0	2.5	3.0	4.0
Mg ²⁺ (25mM)	2 µl	3 µl	4 µl	5 µl	6 µl	8 µl



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10× Hotstart Buffer with Mg²⁺

50mM KCl, 100mM Tris-HCl, 200mM NH₄Cl, 20mM MgCl₂.

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1.1 Recommended PCR assay with Hotstart Buffer (Mg²⁺ plus)

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Sterile deionized water	variable	-
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dNTPs (10mM each)	1 µl	0.2mM each
Primer I	variable	0.4-1µM
Primer II	variable	0.4-1µM
<i>Optimus</i> TM Taq DNA Polymerase (5U/µl)	0.25-0.5 µl	1.25-2.5U/50 µl
Template DNA	variable	10pg-1µg
Total		50 µl

1.2 Recommended PCR assay with Hotstart Buffer (Mg²⁺ free)

Reagent	Quantity	Final concentration
Sterile deionized water	variable	-
10× Hotstart Buffer (Mg ²⁺ free)	5 µl	1×
dNTPs (10mM each)	1 µl	0.2mM each
Primer I	variable	0.4-1µM
Primer II	variable	0.4-1µM
25mM Mg ²⁺	variable	1.0-4.0mM
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Table for selection volume of 25 mM MgCl₂ solution in a 50 µl reaction mix:

Final Mg ²⁺ Conc. (mM)	1.0	1.5	2.0	2.5	3.0	4.0
Mg ²⁺ (25mM)	2 µl	3 µl	4 µl	5 µl	6 µl	8 µl

Recommendation amounts of template DNA in a 50 µl reaction mix:

Human genomic DNA	0.1µg-1µg
Plasmid DNA	0.5ng-5ng
Phage DNA	0.1ng-10ng
E.coli genomic DNA	10ng-100ng

2. Mix contents of tube. Cap tubes and centrifuge briefly to collect the contents to the bottom.

When using a thermal cycler that does not contain a heated lid, overlay the reaction mixture with 25 µl mineral oil.

3. Perform 25-35 cycles of PCR amplification as follows:

Initial Denaturation	94°C	3 minutes
25-35 Cycles	94°C	30 seconds
	55-68°C	30 seconds
	72°C	1-10 minutes
Final Extension	72°C	10 minutes

4. Incubate for an additional 10 min at 72°C and maintain the reaction at 4°C. The samples can be stored at -20°C until use.

5. Analyze the amplification products by agarose gel electrophoresis and visualize by nucleic acid dye staining. Use appropriate molecular weight standards.

Notes on cycling conditions

- Optimus™ Hotstart Taq DNA Polymerase adopts improved chemical modification technology, so it relies on temperature to activate the polymerase activity, which can effectively inhibit non-specific bindings, and the reaction system can be configured at room temperature.
- The half-life of enzyme is >40 minutes at 95°C.
- Optimus™ Hotstart Taq DNA Polymerase accepts modified nucleotides (e.g. biotin-, digoxigenin-, fluorescent-labeled nucleotides) as substrates for the DNA synthesis.
- The number of PCR cycles depends on the amount of template DNA in the reaction mix and on the expected yield of the PCR product. 25-35 cycles are usually sufficient for the majority PCR reaction. Low amounts of starting template may require 40 cycles.

Guidelines for preventing contamination of PCR reaction

During PCR more than 10 million copies of template DNA are generated. Therefore, care must be taken to avoid contamination with other templates and amplicons that may be present in the laboratory environment. General recommendations to lower the risk of contamination are as follows:

- Prepare your DNA sample, set up the PCR mixture, perform thermal cycling and analyze PCR products in separate areas.
- Set up PCR mixtures in a laminar flow cabinet equipped with an UV lamp.

- Wear fresh gloves for DNA purification and reaction set up.
- Use reagent containers dedicated for PCR. Use positive displacement pipettes, or use pipette tips with aerosol filters to prepare DNA samples and perform PCR set up.
- Always perform “no template control” (NTC) reactions to check for contamination.

[3H]-RNA (40000cpm/μg) for 4 hours at 37°C and 70°C.

Quality Control

The absence of endodeoxyribonucleases, exodeoxyribonucleases and ribonucleases is confirmed by appropriate quality tests. Functionally tested in amplification of a single-copy gene from human genomic DNA.

Endodeoxyribonuclease Assay

No detectable conversion of covalently closed circular DNA to a nicked DNA was observed after incubation of 10U *Optimus*TM Hotstart Taq DNA Polymerase with 1μg pBR322 DNA for 4 hours at 37°C and 70°C.

Exodeoxyribonuclease Assay

No detectable degradation of lambda DNA-HindIII fragments was observed after incubation of 10U *Optimus*TM Hotstart Taq DNA Polymerase with 1μg digested DNA for 4 hours at 37°C and 70°C.

Ribonuclease Assay

0% of the total radioactivity was released into trichloroacetic acid-soluble fraction after incubation of 10U *Optimus*TM Hotstart Taq DNA Polymerase with 1μg E.coli

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