

Max Taq DNA Polymerase (recombinant)

Product No : PL2201
Quantity : 200u



Lot :
Expiry Date :
Concentration : 5u/μl
Supplied with : 2ml of 10X ViBuffer A
1ml of 10X ViBuffer S
1ml of 50mM MgCl₂

Store at - 20°C

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Description:

Max Taq DNA Polymerase is a modified and optimized thermostable enzyme blend containing Taq DNA Polymerase, Pfu DNA Polymerase and enhancing factors. It exhibits the 3' to 5' proofreading activity, resulting in considerably higher amplification fidelity than possible with unmodified Taq DNA Polymerase. **Recommended for use in amplification to obtain DNA products up to 20kb.**

Features:

- Ultra pure recombinant protein.
- Excellent for multiplex amplification as it exhibits wider tolerance for Mg²⁺ and salt concentrations, pH, template contaminations and has increased half-life in comparison to unmodified Taq DNA polymerase.
- Improves amplification results with critical templates, such as those containing GC-rich regions, palindromes or multiple repeats.
- Increased amplification product yields and purity.

Unit Definition :

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

Reaction Buffer:

10X ViBuffer A (without MgCl₂):

500mM KCl, 100mM Tris-HCl (pH 9.1 at 20°C) and 0.1% TritonTMX-100. The buffer is optimized for use with 0.1-0.2mM of each dNTP.

10X ViBuffer S:

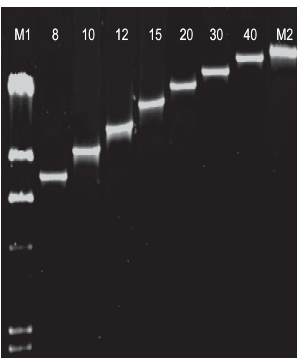
160mM (NH₄)₂SO₄, 500mM Tris-HCl (pH 9.2 at 22°C), 17.5mM MgCl₂ and 0.1% TritonTMX-100. The buffer is optimized for use with 0.35mM of each dNTP.

Storage Buffer:

20mM Tris-HCl (pH 8.0 at 22°C), 100mM KCl, 0.5% TweenTM 20, 0.5% Nonidet P-40, 0.1mM EDTA, 1mM DTT and 50% glycerol.

Quality Control:

All preparation are assayed for contaminating endonuclease, 3'-exonuclease, and non-specific DNase activities. Functionally tested in DNA amplification.



Amplification Using Vivantis Max Taq DNA Polymerase

Lane M1 : VC Lambda / Hind III Marker
Lane 8kb : 8kb amplification products generated using 0.25mM dNTPs, 2u Vivantis Max Taq DNA Polymerase and 3% formamide.
Lane 10kb-20kb : 10kb, 12kb, 15kb, and 20kb amplification products generated using 0.36mM dNTPs, 2u Vivantis Max Taq DNA Polymerase and 3% formamide.
Lane 30kb and 40kb : 30kb and 40kb amplification products generated using 0.36mM dNTPs, 2u Vivantis Max Taq DNA Polymerase and 3% formamide.
Lane M2 : Lambda DNA (indicates 48kb)

SUGGESTED INITIAL PCR CONDITIONS FOR VARIOUS PCR PRODUCT SIZES WITH VIVANTIS DNA POLYMERASE (#PL1201 - 06 / #PL2201 - 06 / #PL3201 - 06 / #PL4201 - 06) REACTION MIX (FINAL CONCENTRATION) :

Primers : 0.2 - 1μM Template: Plasmid (0.02 - 2ng) Lambda (0.1 - 150ng) Genomic (0.05 - 5μg)	Product Size	100bp - 5kb	5kb - 8kb	8kb - 20kb
	dNTP Mix	100μM	200μM	360μM
	ViBuffer (1X)	A	A	S
	Ultrapure DMSO or formamide	-	3%	3%
DNA Polymerase				
Refer to below Table (A)				

Product Size	100bp - 5kb	5kb - 8kb	8kb - 20kb
Denaturation	94°C, 2 min	94°C, 2 min	94°C, 2 min
Denaturation	94°C, 30 s	94°C, 12 s	94°C, 12 s
Annealing*	50 - 68°C, 30 s	50 - 68°C, 30 s	50 - 68°C, 30 s
Extension / 1kb	72°C, 30 s	72°C, 45 s	68°C, 1 min
Cycles	25 - 35	25 - 35	25 - 35
Final Extension	72°C, 7 min	72°C, 7 min	68°C, 7 min

* Primer dependent

TABLE (A) : RECOMMENDED UNITS FOR SPECIFIC VIVANTIS DNA POLYMERASES PER 50μL REACTION VOLUME :

Product Size	Taq (#PL1201 - 06)	Max Taq (#PL2201 - 06)	At Taq (#PL3201 - 06)	AtMax Taq (#PL4201 - 06)
0.1 - 5.0kb	2.0	2.0	2.0	2.0
5.0 - 8.0kb	2.5	2.0	2.5	2.0
8.0 - 20.0kb	2.5	2.0	2.5	2.0
>20.0kb	-	2.0	-	2.0

* This protocol is subjected to changes depending on the template DNA

Product Use Limitation

This product is for research purpose an *in vitro* use only