i-MAX II DNA Polymerase [for Enhancing PCR]

Cat. No. 25261 250 Units

DESCRIPTION

Taq DNA Polymerase is the most common PCR enzyme for amplifying up to 10kb IDNA templates and up to 3kb genomic templates. However, it is not able to amplify even longer DNA fragments. To overcome this constraint, variable PCR systems have been developed by various companies, which are containing Taq DNA Polymerase and thermostable DNA Polymerase with proofreading activity. Both i-MAX™ and i-MAX™ II DNA Polymerase are the PCR System which are developed for amplifying long and complex fragments. The first is designed for amplification of 5 10kb fragments from genomic DNA. The second (i-MAX™ II DNA Polymerase) can amplify even longer fragments up to 20kb from human genomic DNA, and up to 30kb from a IDNA template). Moreover, the second have improved amplification efficiency compared to i-MAX™ DNA Polymerase by improving enzyme activity. Therefore i-MAX™ II DNA Polymerase is a more versatile enzyme blend than i-MAX™ DNA Polymerase in amplifying various templates including short and long DNA fragment or simple and complex DNA, either.

STORAGE

Store at -20 $^{\circ}$ C, and then stable for at least one year.

CHARACTERISTICS

- Increased fidelity of PCR amplification, because the *i*-MAXTM II DNA Polymerase enzyme blend combines the proofreading activity of *Pfu* DNA Polymerase with the high processivity of *Taq* DNA Polymerase.
- Increased yield of PCR amplification, because the accuracy of the enzyme blend reduces the number of truncated amplification products formed.
- **Improved performance of long PCR**, because the reaction buffer and the enzyme blend are optimized for generation of certain length products.

APPLICATIONS

- Standard and long PCR
- PCR with difficult templates
- Cloning with TA and blunt ends

KIT CONTENTS

Label	25261 (250 Units)
$i ext{-MAX II DNA Polymerase} \ (5U/\mu\ell)$	250 Units
10X PCR Buffer* (w/20mM Mg ²⁺)	1.5 ml
10X Mg ²⁺ free PCR Buffer	1.5 ml
10mM dNTPs (2.5mM/each)	800 μℓ
25mM Mg ²⁺	1.5 ml

* 10× PCR BUFFER, 300 mM Tris-HCl(pH 9.0); 300 mM salts containing of K+ and NH₄+; 20 mM Mg²⁺; Enhancer solution

GENERAL REACTION MIXTURE for PCR (total $20\,\mu$ l reaction)

Template	1pg-1 <i>µ</i> g
Primer 1	5-10 pmoles
Primer 2	5-10 pmoles
<i>i</i> -MAX TM II DNA Polymerase (5U/ $\mu\ell$)	0.25 - $0.5\mu\ell$
10x PCR buffer	$2\mu\ell$
dNTP Mixture (2.5mM each)	2 <i>µ</i> ℓ
Sterilized distilled water	up to $20\mu\ell$

CYCLING STEPS for SHORT and LONG FRAGMENTS

Cycle program for fragments < 10kb			
	Temp.	Time	Cycle No.
Initial Denaturation	92-94℃	2-4min	1
Denaturation Annealing Extension*	94℃ 45-65℃ 72℃	15s-1min 15s-1min 1min/1-1.5kb	25-30
Final extension	72℃ 4℃	5-10min hold	1

*, Extension time for 30s-1min is sufficient for fragments < 1kb.

Cycle program for fragments > 10kb

	Temp.	Time	Cycle No.
Initial Denaturation	92-94℃	2-4min	1
Denaturation Annealing Extension	94℃ 45-65℃ 72℃	15s-1min 15s-1min 1min/1-1.5kb	10
Denaturation Annealing Extension	94℃ 45-65℃ 72℃	15s-1min 15s-1min 1min/1-1.5kb + 20s/cycle	15-20
Final extension	72℃ 4℃	5-10min hold	1

Note: This CYCLING PARAMETERS serves as a guideline for PCR amplification. optimal reaction conditions such as PCR cycles, annealing temperature, extension temperature and incubation times, may vary and must be individually determined.

PCR OPTIMIZATION

To produce high yields of specific DNA target sequences, individual reaction component concentrations (and time and temperature parameters) must be adjusted within suggested ranges for efficient amplification of specific targets.

Template DNA

For amplifying long target fragment, it is important that you use high quality, intact, and high pure template DNA.

Primers

PCR primers are oligonucleotide, typically 15-30 bases long, hybridizing to opposite strands and flanking the region of interest in the target DNA. A 40%-60% G+C content is recommended for both primers.



TECHNICAL INFORMATION

EXPERIMENTAL INFORMATION

Amplification

1) Amplification of 570bp Fragment

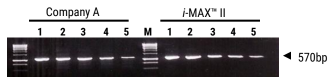


Figure 1. Comparison of *i*-MAX[™] II DNA Polymerase and Company A's PCR System by amplifying 570bp DNA fragment (GAPDH).

Comparison of $i\text{-}\text{MAX}^{TM}$ II DNA Polymerase and Company A's PCR System by amplifying 570bp DNA fragment from variable amounts of cDNA Aliquots of $5\,\mu$ l in $20\,\mu$ l reaction are loaded on 8% agarose gel. The $i\text{-}\text{MAX}^{TM}$ II DNA Polymerase have comparable sensitivity and amplifying performance with company A's longrange PCR System in amplifying 570bp DNA fragment. Lanes M, 100bp Ladder DNA marker (iNtRON's); lanes 1, 2^{-3} diluted cDNA; lane 3, 2^{-5} diluted cDNA; lane 4, 2^{-6} diluted cDNA; lane 5, 2^{-7} diluted cDNA

2) Amplification of 1.3Kb Fragment

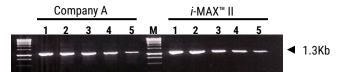
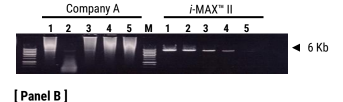


Figure 2. Comparison of *i*-MAX[™] II DNA Polymerase and Company A's PCR System by amplifying 1.3 kb DNA fragment.

Comparison of $i\text{-MAX}^{\text{TM}}$ II DNA Polymerase and Company A's PCR System by amplifying 1.3 kb DNA fragment from variable amounts of human genomic DNA Aliquots of $5\mu\ell$ in $20\mu\ell$ reaction are loaded on 8% agarose gel. The $i\text{-MAX}^{\text{TM}}$ II DNA Polymerase have comparable sensitivity and amplifying performance with company A's long-range PCR System in amplifying 1.3kb DNA fragment. Lanes M, 1kb Ladder DNA marker (iNtRON's); lanes 1, 300 fg; lane 2, 150 fg; lane 3, 75fg; lane 4, 37.5fg; lane 5, 18.75fg

3) Amplification of 6Kb / 20Kb Fragment

[Panel A]



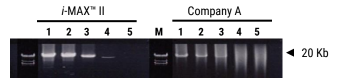


Figure. 3. Comparison of *i*-MAX[™] II DNA Polymerase and Competitor A's PCR System in amplifying 6kb and 20kb fragment

Comparison of $i\text{-MAX}^{\text{TM}}$ II DNA Polymerase and Company A's PCR System by amplifying 6Kb, 20Kb DNA fragment from variable amounts of λ DNA. Aliquots of $5\mu\ell$ in $20\mu\ell$ reaction are loaded on 8% agarose gel. The $i\text{-MAX}^{\text{TM}}$ II DNA Polymerase have comparable sensitivity and amplifying performance with company A's long-range PCR System in amplifying 6Kb, 20Kb DNA fragment.

[Pannel A] 6 Kb product; [Pannel B] 20kb product [Pannel A] Lanes M, λDNA EcoRl digest; lanes 1, 300 fg; lane 2, 150 fg; lane 3,

75 fg; lane 4, 37.5 fg; lane 5, 18.75 fg

[Pannel B] Lanes M, λDNA EcoRI digest; lanes 1, 1 ng; lane 2, 100 pg; lane 3, 10 pg; lane 4, 1 pg; lane 5, 100 fg

TROUBLESHOOTING GUIDE

Problem	Possible Cause	Recommendation
Little or no product	Primer	- Primer concentration not optimal or primers degraded or primers degraded
no product	Problems with starting template	- Check the concentrations, storage conditions, and quality of the starting template.
	Incorrect annealing	templater
	temperature or time	-Decrease annealing temperature by $2{}^{\circ}\!$
	Extension time too short	- Annealing time should be between 30 and 60s.
	Hot start may be necessay	-Increase the extension time by increments of 1 $\!$ min.
Product is multi-banded		-Perform manual hotstrat PCR or iNtRON's hotstart PCR enzyme (i -StarMAX $^{\text{TM}}$).
or smeared	Annealing temperature too low	-Including the points mentioned above, check the facts below.
	Hot start may be necessary	-Increase annealing temperature in $2^\circ\!\!\mathbb{C}$ increments.
	Primer design not optimal	
	Too much start template	

RELATED PRODUCTS

Product Name	Cat. No.
G-spin™ Total DNA Extraction Mini Kit	17045/17046
i-genomic Plant DNA Extraction Kit	17371
i-genomic BYF DNA Extraction Kit	17361
DNA-spin™ Plasmid DNA Purification Kit	17096 / 17097 / 17098
MEGAquick-spin™ Total Fragment DNA Purification Kit	17286 / 17287 / 17288
Maxime RT PreMix Kit (Oligo[dT] ₁₅ Primer)	25081
Maxime RT PreMix Kit (Random Primer)	25082
Maxime PCR PreMix Kit(i-StarMAX II)	25281
RevoScript™ RT PreMix Kit(Random Primer)	25085 / 25086
RevoScript™ RT PreMix Kit(Oligo dT ₁₅ Primer)	25083 / 25084
RealMOD™ Real-time PCR Master mix Kit(2X)	25341 / 25342
RealMOD™ Green Real-time PCR Master mix Kit(2X)	25343 / 25344