Maxime PCR PreMix Kit (i-Taq)

for $20\mu\ell$ rxn / $50\mu\ell$ rxn

Cat. No. 25025(for $20\mu\ell$ rxn, 96 tubes) **Cat. No. 25026**(for $20\mu\ell$ rxn, 480 tubes)

DESCRIPTION

INtRON's *Maxime* PCR PreMix Kit has not only various kinds of PreMix Kit according to experience purpose, but also a 2X Master mix solution. *Maxime* PCR PreMix Kit (*i*-Taq) is the product what is mixed every component: *i*-Taq[™] DNA Polymerase, dNTP mixture, reaction buffer, and so on- in one tube for 1 rxn PCR. This is the product that can get the best result with the most convenience system. The first reason is that it has every components for PCR, so we can do PCR just add a template DNA, primer set, and D.W.. The second reason is that it has Gel loading buffer to do electrophoresis, so we can do gel loading without any treatment. In addition, each batches are checked by a thorough Q.C., so its reappearance is high. It is suitable for various sample's experience by fast and simple using method.

STORAGE

Store at -20°C; under this condition, it is stable for at least a year.

CHARACTERISTICS

- · High efficiency of the amplification
- · Ready to use: only template and primers are needed
- Stable for over 1 year at -20 °C
- · Time-saving and cost-effective

CONTENTS

• Maxime PCR PreMix (i-Taq, for 20 ₩ rxn)

96 (480) tubes

Maxime PCR PreMix (i-Taq, for 50^{µℓ} rxn)

96 tubes

Component in	20 μ reaction	50 μ reaction
i-Taq™ DNA Polymerase(5U/μℓ)) 2.5U	5U
dNTPs	2.5mM each	2.5mM each
Reaction Buffer(10x)	1x	1x
Gel Loading buffer	1x	1x

Note: The PCR process is covered by patents issued and applicable in certain countries. iNtRON Biotechnology does not encourage or support the unauthorized or Unlicensed use of the PCR process. Use of this product is recommended for persons That either have a license to perform PCR or are not required to obtain a license.

PROTOCOL

1. Add template DNA and primers into *Maxime* PCR PreMix tubes (*i*-Taq).

Note 1 : Recommended volume of template and primer : $3\mu\ell \sim 9\mu\ell$ Appropriate amounts of DNA template samples

- · cDNA: 0.5-10% of first RT reaction volume
- · Plasmid DNA: 10pg-100ng
- Genomic DNA: 0.1-1ug for single copy

Note 2: Appropriate amounts of primers

- Primer : 5-20pmol/ $\mu\ell$ each (sense and anti-sense)
- 2. Add distilled water into the tubes to a total volume of $20\,\mu\!\ell$ or $50\,\mu\!\ell$. Do not calculate the dried components

Total 20 μ or 50 μ reaction volume

PCR reaction mixture	Add	Add	
Template DNA	1 ~ 2 <i>µ</i> ℓ	2 ~ 4 <i>μ</i> ℓ	
Primer (F : 10pmol/ $\mu\ell$)	$1\mu\ell$	$2 \sim 2.5 \mu \ell$	
Primer (R : 10pmol/ $\mu\ell$)	$1\mu\ell$	2 ~ 2.5 µl	
Distilled Water	16 ~ 17 <i>μ</i> ℓ	44 ~ 41 $\mu\ell$	
Total reaction volume	20 με	50 μ l	

Note: This example serves as a guideline for PCR amplification. Optimal reaction conditions such as amount of template DNA and amount of primer, may vary and must be individually determined.

3. Dissolve the blue pellet by pipetting.

Note: If the mixture lets stand at RT for 1-2min after adding water, the pellet is easily dissolved.

4. (Option) Add mineral oil.

Note: This step is unnecessary when using a thermal cycler that employs a top heating method(general methods).

- 5. Perform PCR of samples.
- Load samples on agarose gel without adding a loading-dye buffer and perform electrophoresis.

SUGGESTED CYCLING PARAMETERS

PCR cycle		Temp.	PCR product size		
			100-500bp	500-1000bp	1Kb-5Kb
Initial	Initial denaturation		2min	2min	2min
30-40 Cycles	Denaturation	94℃	20sec	20sec	20sec
	Annealing	50-65℃	10sec	10sec	20sec
	Extension	65-72℃	20-30sec	40-50sec	1min/Kb
Fina	Final extension		Optional. Normally, 2-5min		

EXPERIMENTAL INFORMATION

· Comparison with different company kit

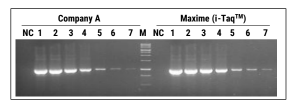


Fig.1. Comparison of *Maxime* PCR PreMix (*i*-Taq) and Company A's PreMix system by amplifying 1 Kb DNA fragment.

After diluting the λDNA as indicates, the PCR reaction was performed with Maxime PCR PreMix (i-Taq) and company's A product. Lane M, SiZer-1000 DNA Marker; lane 1, undiluted λDNA ; lane 2, 200 ng λDNA ; lane 3, 40 ng λDNA ; lane 4, 8 ng λDNA ; lane 5, 1.6 ng λDNA ; lane 6, 320 pg λDNA ; lane 7, 64 pg λDNA ; lane NC, Negative control

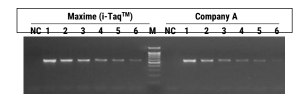


Fig.2. Comparison of *Maxime* PCR PreMix (i-Taq) and Company A's PreMix system by amplifying 570 bp DNA fragment (GAPDH).

Total RNA was purified from SNU-1 using easy-BLUETM Total RNA Extraction Kit (Cat. No. 17061). And then, the first strand of cDNA was synthesized using Power cDNA Synthesis Kit (Cat. No. 25011). After diluting the cDNA mixture as indicates, the RT-PCR reaction was performed.

lane M, SiZer-100 DNA Marker; lane 1, undiluted cDNA; lane 2, 1/2 diluted cDNA; lane 3, 1/4 diluted cDNA; lane 4, 1/8 diluted cDNA; lane 5, 1/16 diluted cDNA; lane 6, 1/32 diluted cDNA; lane NC, Negative control

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