



Maxime™ PCR PreMix Kit (i-MAXII)

QUICK GUIDE

96 tube : 25265

1. KIT DESCRIPTION

iNtRON's Maxime™ PCR PreMix Kit is an optimized, ready-to-use PCR master mix system containing i-MAX™ II DNA Polymerase, dNTP mixture, reaction buffer, and a gel loading buffer in a single PCR tube for a 1-reaction scale. This system ensures maximum convenience and reproducibility by minimizing pipetting steps. Users simply need to add template DNA, primer sets, and sterile distilled water to execute amplification. Furthermore, the inclusion of a tracking dye/gel loading buffer allows direct gel loading onto agarose electrophoresis gels without additional sample preparation. Every manufacturing lot undergoes rigorous Quality Control (QC) to guarantee exceptional batch-to-batch consistency and high amplification performance.

2. CHARACTERISTICS

- **All-in-One System** : Contains i-StarMAX™ II DNA Polymerase, dNTP mixture, reaction buffer, and a tracking dye in a single pre-aliquoted tube for maximum convenience and reproducibility.
- **High-Fidelity Amplification** : The enzyme blend combines proofreading activity with efficient DNA synthesis to support accurate PCR amplification.
- **Long PCR capability** : Suitable for amplification of both short and long DNA fragments.
- **Direct Gel Loading** : PCR products can be loaded directly onto an agarose gel without additional loading dye.
- **Versatile Applications** : Suitable for plasmid DNA, phage DNA, genomic DNA, and other DNA templates.

3. KIT CONTENTS AND STORAGE

Contents	25265
Maxime PCR PreMix (i-MAXII)	12 Strips/pk (96 Tube)

- **Storage Condition** : Store strictly at -20°C immediately upon receipt.
- **Shelf Life & Stability** : Guaranteed to maintain stable performance through the indicated Expiration Date when stored and handled under recommended conditions.

4. CONSIDERATIONS BEFORE USE

- **Research Use Only (RUO)** : For research use only. Not for diagnostic or clinical procedures.
- **Storage & Handling** : Store strictly at -20°C. Avoid repeated freeze-thaw cycles.
- **Contamination Control** : Use clean gloves and filter pipette tips to prevent cross-contamination.
- **PreMix Dissolution** : Ensure complete pellet dissolution; briefly spin down if necessary prior to use.
- **Template Integrity** : Use high-purity DNA free of inhibitors (phenol, ethanol, EDTA).

5. PROTOCOLS

- 1) Prepare the reaction mixture by adding components into the pre-aliquoted PCR tube as follows :

- ⚠ **Template DNA** : 1 ~ 2 µl (Recommended : 1 pg-1 µg)
- ⚠ **Forward Primer (10 pmol/µl)** : 1 µl
- ⚠ **Reverse Primer (10 pmol/µl)** : 1 µl
- ⚠ **Sterile Distilled Water (D.W.)** : 16-17 µl (final vol : 20 µl)

⚠ **NOTE** : Since the PreMix already contains enzyme and buffer components, adjust the volume of distilled water according to the template and primer volumes.

- 2) 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.

3) Perform PCR of samples.

[SUGGESTED CYCLING PARAMETERS]

Tube	Temp.	Time	
		≤ 10 kb	> 10kb
Initial denaturation	94°C	2-4min	
Denaturation	94°C	15sec-1min	
25-30 Cycles	Annealing	Tm-dependent*	15sec-1min
	Extension	72°C	1min/ 1-1.5kb 11-30 cycles: +20 sec/cycle*
Final extension	72°C	5-10min	

*Annealing temperature: 3-5°C below the lower primer Tm.

*For targets >10 kb, use the standard extension time for the first 10 cycles, then increase the extension time by 20 sec at each successive cycle for the remaining 15-20 cycles.

- 4) Load the PCR products directly onto an agarose gel without adding loading dye, and perform electrophoresis.

6. TROUBLESHOOTING GUIDE

Refer to the following troubleshooting guide for possible causes and recommended actions when unexpected PCR results are observed.

Observation	Possible Cause	Recommendation
Little or no PCR product	Insufficient template or poor template quality	Check template concentration and quality
	Primer concentration not optimal	Optimize primer concentration
	Incorrect cycling conditions	Optimize annealing temperature, cycle number, and extension time
	Insufficient extension time	Increase extension time according to amplicon length
Non-specific bands	Annealing temperature too low	Increase annealing temperature
	Excessive template	Reduce template amount
	Excessive primer concentration	Reduce primer concentration
Primer-dimer formation	Primer concentration/design not optimal	Optimize primer concentration or primer design
Smeared bands	Poor template quality or excessive DNA	Use purified DNA and reduce template amount
	Excessive cycle number	Reduce the number of PCR cycles

7. PCR OPTIMIZATION NOTES

- Annealing temperature should be optimized according to primer T_m.
- Extension time should be adjusted according to amplicon length.
- For targets >10 kb, increase the extension time by 20 sec at each successive cycle after the first 10 cycles.
- Template and primer concentrations may require optimization depending on the target.

8. CAUTIONS

- Wear appropriate personal protective equipment when handling samples and reagents.
- Use aerosol-resistant filter tips to minimize cross-contamination.
- Use separate areas for PCR setup and post-PCR analysis whenever possible.
- Avoid contamination of reagents with amplified DNA.
- Dispose of biological samples and PCR products according to applicable laboratory procedures.

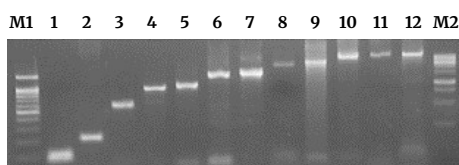
9. ADDITIONAL REQUIRED EQUIPMENT

- Thermal cycler
- Micropipettes and aerosol-resistant tips
- PCR-grade water
- Forward and reverse primers
- Agarose gel electrophoresis equipment

10. EXPERIMENTAL INFORMATION

1) Broad-range amplification performance

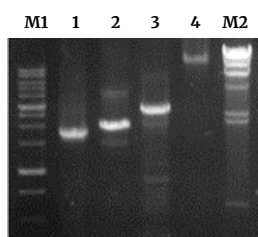
Maxime™ PCR PreMix Kit (i-MAX™ II) was evaluated using various DNA templates and target sizes ranging from short fragments to long PCR products. Efficient amplification was observed across targets ranging from 87 bp to 20 kb.



Lane M1, 100bp Ladder DNA Marker; lane 1, 87bp; lane 2, 200bp; lane 3, 570bp; lane 4, 1Kb; lane 5, 1.3Kb; lane 6, 1.8kb; lane 7, 2Kb; lane 8, 2.7Kb; lane 9, 4.5Kb; lane 10, 9Kb; lane 11, 17.5Kb; lane 12, 20Kb; lane M2, 1Kb Ladder DNA Marker

2) Long-range amplification from genomic DNA

Maxime™ PCR PreMix Kit (i-MAX™ II) successfully amplified DNA fragments up to 17.5 kb from human genomic DNA.



Lane M1, 1Kb Ladder DNA Marker; lane M2, λ/Hind III Marker; lane 1, 1.8Kb; lane 2, 2Kb; lane 3, 2.7Kb; lane 4, 17.5Kb

10. EXPLANATION OF SYMBOLS

LOT	Batch code	REF	Catalogue number
	Contains sufficient for <n> tests		Temperature limit
	Date of manufacture		Manufacturer
	Use-by date	RUO	Research Use Only
	Consult instructions for use		Caution

Visual Protocol

