[Cat. No.] K-2036, K-2037

Introduction

AccuPower® PCR PreMix (Negative Dye) is the powerful technology for convenient and easy performance of DNA amplification. This product contains vacuum-dried components including *Top* DNA Polymerase, dNTPs, reaction buffer, and stabilizer. It simplifies preparation of reaction mixture by adding template DNA and primers without any extra process. After the reaction, loading-dye mixture must be added to samples, and then loaded on agarose gel for analysis.

Applications

- Conventional PCR
- Primer extension
- TA cloning
- · Gene sequencing

Features & Benefits

- Stability: Included stabilizer enables delivery at room temperature and provides increased stability compared to solution-type products.
- User-friendly: Reactants are individually packaged in each of the PCR tubes, it allows any user simply perform PCR by adding template DNA and primers.
- Sensitivity: Excellent sensitivity and amplification efficiency even with small amounts of DNA.
- Reproducibility: Mass production under ISO 9001 quality system allows minimized deviation between lots and reproducible results in replicated tests performed under same conditions and variation.

Composition

Composition	Concentration
Top DNA Polymerase	1 U
dNTPs (dATP, dCTP, dGTP, dTTP)	Each 250 μM
Reaction buffer with 1.5 mM MgCl ₂	1X
Stabilizer	0

Specifications

Top DNA Polymerase					
5' to 3' exonuclease activity	No				
3' to 5' exonuclease activity	No				
3'-A overhang	Yes				
Fragment size	Up to 10 kb				

Storage

Store at -20°C. If stored in the recommended temperature, this product will be stable until the expiration date printed out on the label.

Precautions

 This enzyme is specifically optimized for increasing base incorporation rate by inactivating 5' to 3' exonuclease activity.
 Therefore, this product is not recommended to use for real-time PCR using hydrolysis probe method.

Online Resources





Korean

Enalish

Visit our product page for additional information and protocols

Ordering Information

Descr	Cat. No.		
0.2 ml thin-wall 8-tube strips	96 tubes	20 µl/rxn (-dye)	K-2036
with attached cap	480 tubes	20 µl/rxn (-dye)	K-2037

Notice

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Explanation of Symbols





Biological Risks





















Experimental Procedures

Steps		Procedure Details					
		1. Add template DNA, primers, and nuclease-free water into <i>AccuPower</i> ® PCR PreMix (Negative Dye) tubes to make a total volume of 20 μl. Do not include the dried pellet.					
		·	Amount of template		A		
		Template DNA			Amount of template		
		Bacteriophage λ, Plasmid D	JNA	100 fg-200 ng			
4		Total genomic DNA 1-500 ng • Preparation of reaction mixture					
1			Components		reaction		
	Preparation of	Template DNA	·		e (1-10 µI)		
	reaction mixture	Forward primer (10 pmol/µl))	0.5	5-2 µl		
		Reverse primer (10 pmol/µl)	0.5	5-2 µl		
		Nuclease-free water		Va	riable		
		Total volume		2	0 μΙ		
		or vortexing, and br	iefly spin down.				
			3. Perform the reaction under the following conditions.				
	Incubate reactions in a thermal cycler	Step	Temperature	Time	Cycles		
2		Pre-denaturation	95°C	5 min	1 cycle		
		Denaturation	95°C	20 sec			
		Annealing	45-65°C	20 sec	25-35 cycles		
		Extension	72°C	0.5-1 min/kb			
	anorman cycle.	Final extension	72°C	3-5 min	1 cycle		
3	Analyze with gel electrophoresis	 4. After the reaction, maintain the reaction mixture at 4-8°C. The samples can be stored at -20°C until use. 5. Load samples on agarose gel with adding a loading-dye mixture (Cat. No. C-9029, not provided), and perform gel electrophoresis for analysis. 					
		If primer's Tm value is more than 65°C or PCR product size is more than the conditions as below.					
		Step	Temperature	Time	Cycles		
	([)	Pre-denaturation	95°C	5 min	1 cycle		
	\ • /	Denaturation	95°C	20 sec			
					20 25 0/0100		
	Option	Annealing/Extension	68°C	1 min/kb	30-35 cycles		