[Cat. No.] Please refer to the Ordering Information

Introduction

AccuPower® Taq PCR PreMix is the powerful technology for convenient and easy performance of DNA amplification. This product contains vacuum-dried components including *Taq* DNA Polymerase, dNTPs, reaction buffer, stabilizer, and tracking dye. It simplifies preparation of reaction mixture by adding template DNA and primers without any extra process. After the reaction, samples can be applied directly 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.
- Sensitivity: Excellent sensitivity and amplification efficiency even with small amounts of DNA.
- 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.
- 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
Taq DNA Polymerase	1 U
dNTPs (dATP, dCTP, dGTP, dTTP)	Each 250 µM
Reaction buffer with 1.5 mM $MgCl_2$	1X
Stabilizer and tracking dye	0

Specifications

Taq DNA Polymerase				
5' to 3' exonuclease activity	Yes			
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.

Online Resources





English

Visit our product page for additional information and protocols

Ordering Information

Description		Cat. No.	
	96 tubes	20 µl/rxn	K-2601
0.2 ml thin-wall 8-tube strips	vali	50 µl/rxn	K-2603
with attached cap	480 tubes	20 µl/rxn	K-2602
		50 µl/rxn	K-2604

Notice

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Explanation of Symbols



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1

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Experimental Procedures

	Steps	Procedure Details					
		 Add template DNA, primers, and nuclease-free water into <i>AccuPower</i>[®] Taq PCR PreMix tubes to make a total volume of 20 μl or 50 μl. Do not include the dried pellet. Amount of template 					
		Template DNA	Amount of template				
		-	20 µl reaction		50 µl reaction		
		Bacteriophage λ , Plasmid DNA			100 fg-500 ng		
1		Total genomic DNA	1-500	ng	1 ng-1 µg		
		Preparation of reaction mixture Components 20 µl reaction 50 µl re					
	Preparation of	Components Template DNA			50 µl reaction		
	reaction mixture	Forward primer (10 pmol/µl)	Variable (1	. ,	Variable (1-25 µl) 1-5 µl		
		,	0.5-2	-	-		
		Reverse primer (10 pmol/µl)			1-5 µl		
	Nuclease-free water		Variable		Variable		
		Total volume	20 µl		50 µl		
		2. Dissolve the vacuum-dried blue pellet by pipetting or vortexing, and briefly spin down					
		3. Perform the reaction under the following conditions.					
		Step Te	emperature	Time	Cycles		
	Loo of some	Pre-denaturation	95°C	1-5 min*	1 cycle		
2		Denaturation	95°C 30 sec				
2	ARC	Annealing	45-65°C† 30 sec		25-35 cycles		
	Incubate reactions in a	Extension	72°C	0.5-1 min/kb			
	thermal cycler	Final extension	72°C 3-5		1 cycle		
		* When using genomic DNA as template DNA, set it to 5 min. † The optimal annealing temperature depends on the melting temperature of the primers.					
3		4. After the reaction, maintain the reaction mixture at 4-8°C. The samples can be s at -20°C until use.					
J	Analyze with gel electrophoresis	5. Load 5 μl of samples on agarose gel without adding a loading-dye mixture, and perform gel electrophoresis for analysis.					
		 If primer's Tm value is more than 65°C or PCR product size is more than 5 kb, fol the conditions as below. 					
			emperature	Time	Cycles		
			emperature 95°C	Time 1-5 min	Cycles 1 cycle		
	(!)	Step T	-		1 cycle		
	Option	StepTPre-denaturation	95°C	1-5 min	-		

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