

Super FCR Masterializ ZA

< For Research Use Only >

Cat No :YT1553 Size: 1ml

Storage: Store at -20°C.

Introduction

The Enzyme used in this mastermix is Taqplus. Taq Plus DNA polymerase is a mixture of Taq DNA polymerase and an enzyme containing $3' \rightarrow 5'$ exonuclease activity.

Its fidelity is 6 times greater than that of Taq DNA Polymerase. Compared with *Taq* DNA Polymerase, *Taq* Plus DNA polymerase has stronger amplification performance, higher sensitivity, and is more tolerant of impurities within 5kb amplifying range.

2X super PCR MasterMix contains everything required for PCR, except primers and template, thereby easing PCR setup and improving reproducibility. It can amplify up to <u>10 kb</u> from human genomic DNA or up to <u>15 kb</u> from λ DNA. Protective agents in the 2X super PCR MAsterMix enable the resistance to repeated freeze-thaw cycles.

Attention: if you have more than 20 freeze-thaw cycles, then first aliqoat the mastermix.

Dyes contained in 2X super PCR MAsterMix enable direct loading PCR products onto agarose gels. The obtained PCR

PCR products contain A at the 3'ends and can be directly cloned into T-Vectors.

Package Information

Components

2X super PCR MAsterMix (Dye Plus) Cat	No: YT1553 Size:1ml
---------------------------------------	---------------------

Unit Definition

One unit (U) is defined as the amount of enzyme that incorporates 10 nmol of dNTPs into acid-insoluble products in 30

minutes at 74°C with activated salmon sperm DNA as the template/primer.

Protocol

General reaction Mixture for PCR	50µl reaction	25μl reaction	
ddH2O	Το 50 μl	To 25 μl	
Super PCR mastermix 2X	25 μl	12.5 μl	
Template DNA*	Optional	Optional	
Primer1 (10μM)	2 μl	1 μl	
Primer 2 (10μM)	2 μl	1 μl	

تلفن :www.yektatajhiz.com •۲۱-۷۷۷۳۶۰۳۶

info@yektatajhiz.com



*Recommended amount of DNA template for a 50 µl reaction system is as follow:

Human genomic DNA	0.1∼1 μg	
Bacterial Genomic DNA	10 \sim 100ng	
λ DNA	$0.5{\sim}5$ ng	
Plasmid DNA	0.1~10ng	

فکس: ۲۱-۷۷۷۲۷۸۰۵

Thermocycling Condition for a routine PCR

94∘C	5 min (pre-denaturation)
94∘C	30 sec
55∘C *	30 sec 🗧 35 Cycles
72∘C	60 sec/kb 🔄
72∘C	7 min (final extention)
4∘C	Hold

*The optional annealing temperature should be $1-2 \circ C$ lower than the T_m of primers used.

Primers Designing Notes

1. Choose C or G as the last base of the 3' end of the primer;

- 2. Avoid continuous mismatching at the last 8 bases of the 3' end of the primer;
- 3. Avoid hairpin structure at the 3' end of the primer;
- 4. Tm of the primers should be within the range of $55^{\circ}C \sim 65^{\circ}C$;
- 5. Additional sequence should not be included when calculating Tm of the primers;
- 6. GC content of the primers should be within the range of 40% \sim 60%;
- 7. Tm and GC content of forward and reverse primes should be as similar as possible.