

## 2× Taq PCR Mix

Cat. no. 4992229/4992230/4992255/4992256

### Storage

For long term storage, store at -20°C, and *Taq* PCR MasterMix retain full activity for repeated freezing and thawing. For regular use, please store at 2-8°C.

### Contents

Contents	4992229	4992230	4992255	4992256
2× <i>Taq</i> PCR Mix	1 ml	5 × 1 ml	1 ml	5 × 1 ml
ddH <sub>2</sub> O	1 ml	5 ml	1 ml	5 ml
Loading dye in PCR Mix	Yes	Yes	No	No

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The product is used for research only, neither intended for the diagnosis, or treatment of a disease, nor for the food, or cosmetics etc.

### Introduction

*Taq* PCR Mix is a 2× concentrated, optimized mixture composed of *Taq* DNA polymerase, dNTPs, MgCl<sub>2</sub>, reaction buffer, PCR reaction enhancer, optimizer and stabilizer. The advantages of 2× *Taq* PCR MasterMix include high convenience, sensitivity, specificity and stability. It minimizes man-made errors during PCR operating process. *Taq* PCR Mix is suitable for routine PCR reaction, amplification of complex templates such as GC rich templates (>60%) and templates with secondary structure, and large-scale gene detection.

### Mix Components (2×)

0.1 U/μl *Taq* Polymerase  
 500 μM dNTP each  
 20 mM Tris-HCl (pH8.3)  
 100 mM KCl  
 3 mM MgCl<sub>2</sub>  
 Stabilizer and enhancer

### Description

2 × *Taq* PCR Mix is designed for quick and easy preparation of reaction mixture, which minimizes the contamination during PCR operating process. For PCR reaction set-up, users only need to pipet an aliquot part of 2× *Taq* PCR Mix and dilute the Mix to 1× by adding templates, primers and water up to the reaction volume. There are two types of this product: Mix with loading dye (blue) and Mix without loading dye (colorless); PCR products produced by using Mix with loading dye can be loaded directly without extra loading buffer.

### Application

- Gene detection: 2 × *Taq* PCR Mix is especially suitable for large-scale gene detection, semi-quantitative PCR, detection of tiny amount of DNA, etc.
- Amplification of DNA and complex templates such as GC rich templates (>60%) and templates with complex secondary structure. The enzyme of *Taq* PCR Mix generates PCR products with A-tailing, suitable for TA cloning.

### Example

**Note: The following example only for reference, user must set up optimal reaction system according to different reaction conditions such as different templates or primers etc.**

- To 25 μl PCR reaction system: 1 kb fragment of human genomic DNA was amplified by using 2× *Taq* PCR Mix (If use different reaction system, please proportionally increase or decrease the amount of reaction components referring to this system).

Template	< 1 μg
Primer 1(10 μM)	1 μl
Primer 2(10 μM)	1 μl
2× <i>Taq</i> PCR Mix	12.5 μl
ddH <sub>2</sub> O	up to 25 μl

- PCR cycle set-up

94°C	3 min	} 30 cycles
94°C	30 sec	
55°C	30 sec	
72°C	1 min	
72°C	5 min	

- Result detection: Load 5 μl PCR products to agarose gel for PCR detection.