

# **TOOLSite-Directed Multi-Mutagenesis Kit**

For rapid site-directed mutation at multiple sites



## Contents

Introduction	3
Kit Contents	4
Work Flow	5
Protocol	6
Troubleshooting	12

## Introduction

TOOLSite-Directed Multi-Mutagenesis Kit is designed for rapid site-directed mutagenesis based on the new rapid cloning technology. This kit can perform three to five separate site-directed mutagenesis in one reaction. The amplified target fragments are digested by DpnI, cyclized by Multi-Toolsitase/buffer, and then transformed to *Escherichia coli* to complete the site-directed mutagenesis. The kit consists of two modules: TOOLS Ultra High Fidelity DNA Polymerase amplification module and rapid cloning module. TOOLS Ultra High Fidelity DNA Polymerase can significantly reduce the possibility of unexpected mutations during amplification of up to 20 kb. The rapid cloning module replaces conventional annealing ring-forming reactions with homologous recombination reactions, requiring considerably less template and having a more flexible primer design strategy. The kit is equipped with Multi-Toolsitase, a recombinant enzyme specifically optimized for multi-site-directed mutations. DpnI digestion of specific amplification products can be directly applied in a recombination reaction without purification. Highly optimized reaction buffer, fast-operating procedures, and efficient site-directed mutagenesis make TOOLSite-Directed Mutagenesis Kit the preferred choice for DNA multi-site mutation.

#### Features

- 1. TOOLS Ultra High Fidelity DNA Polymerase provides high-fidelity polymerase chain reaction (PCR) with the lowest mutation rate.
- 2. TOOLS Ultra High Fidelity DNA Polymerase with excellent long fragment amplification capability can be widely used for any plasmid amplification within 20 kb.
- 3. Amplification is conducted exponentially, and the template usage is extremely low, which is beneficial for completely degrading the original methylation template.
- 4. DpnI eliminates original template contamination.
- 5. The rapid Multi-Toolsitase cloning system can efficiently cyclize PCR products.
- 6. Amplified products can be directly used for recombinant reaction after digestion with DpnI.
- 7. Mutations of three to five discontinuous sites (>50 bp apart) can be achieved in one reaction.

Note: Please use methylase nondefective host strains (e.g., Top10, DH5 $\alpha$ , and JM109) to extract the original plasmid when using this kit for site-directed mutagenesis of plasmids.

## **Kit Contents**

Contents	TTC-CB15 (10rxn)
2X Multi-Buffer I	1.25 ml
dNTP (10 mM)	50 µl
TOOLS Ultra High Fidelity DNA Polymerase	50 µl
Dpnl (10U/µl)	50 µl
5x Multi-Buffer II	40 µl
Multi-Toolsitase	20 μl

#### Storage

All components of the kit can be stored at -20°C for 1 year. Repeated freeze-thaw cycles should be avoided.

## **Work Flow**

Separate double-base site-directed mutations (The distance between two mutation sites is >50 bp).



Figure 1: Overview of separate three-base site-directed mutations with this kit

#### Quick guide

Choose the mutation sites A, B, and C as the boundary to divide the vector into fragments AB, BC, and CA. The three mutation sites contain design reverse complementary primers. Amplify fragments AB (forward primer A and reverse primer B), BC (forward primer B and reverse primer C), and CA (forward primer C and reverse primer A) with the original plasmid as the template (Figure 1, I). The amplification product was digested by DpnI (Figure 1, I), and then, the digestion product was directly used in the recombination reaction (Figure 1, II). The recombination product was transformed directly to complete multiple-base site-directed mutagenesis (Figure 1, III).

### Protocol

1. Primer design

To introduce the site-directed mutations of three separate bases in a plasmid, three pairs of primers are needed to amplify the plasmid in three parts. The basic principles for primer design: The 5' ends of reverse and forward primers comprise a 15–21-bp reverse complementary region. The mutation sites can be in the complementary region (the mutation should be introduced in both primers) or in the noncomplementary region of any one of the primers. The mutation site should NOT be at the end of the primer. Figure 2 shows the details of the primer design, illustrated with introduction of three base mutations into vector pUC18.



Note: Primer designs at mutation sites B and C are the same as that of site A. Calculation of the Tm value of primers should be based on the region between the mutation site and the 3' end of the primer. The Tm value should exceed 60°C through adjusting the primer length. Please note that the region

between the mutation site and the 5' end of the primer should not be included for calculating the Tm value.

2. Target plasmid amplification

The plasmid is divided into fragments AB, BC, and CA through mutation sites A, B, and C. Phanta Max Super-Fidelity DNA Polymerase is used to amplify the three fragments. Primers required to amplify AB are the forward primer of mutation site A and the reverse primer of mutation site B. Primers required to amplify BC are the forward primer of mutation site B and the reverse primer of mutation site C. Furthermore, primers required to amplify CA are the forward primer of mutation site C and the reverse primer of mutation site A.

Use TOOLS Ultra High Fidelity DNA Polymerase to amplify the target plasmid. Each component should be mixed well after thawing and placed back at -20°C after use. The recommended reaction system is as follows:

Component	Volume
ddH2O	Up to 50 µl
2X Multi-Buffer I	25 μl
dNTP (10 mM) <sup>a</sup>	1 μl
Template DNA <sup>b</sup>	Optional
Primer1 (10 µM)	2 µl
Primer2 (10 µM)	2 µl
TOOLS Ultra High Fidelity DNA Polymerase <sup>c</sup>	1 μl

- a. Do not use dUTP or any primer and template containing uracil.
- b. Given the normal amplification of the plasmid, use the least possible number of templates. Less than 1 ng of freshly extracted plasmid is recommended.
- c. The recommended final concentration of enzyme is 1 U/50  $\mu$ L. The optimal concentration of TOOLS Ultra High Fidelity DNA Polymerase is 0.5–2 U per 50  $\mu$ L. No more than 2 U per 50  $\mu$ L is recommended, especially when the amplicon is >5 kb.

After all components are mixed, the recommended PCR program is as follows:

Steps	Temperature	Time	Cycle number	
Predenaturation <sup>a</sup>	95°C	30 sec	1	
Denaturation <sup>a</sup>	95°C	15 sec		
Annealing <sup>b</sup>	60°C -72°C	15 sec	30 <sup>d</sup>	
Extension <sup>c</sup>	72°C	30-60 sec/kb	ec/kb	
Complete extension	72°C	5 min	1	

a. For most plasmids, the appropriate denaturation temperature is 95°C.

- b. TOOLS Ultra High Fidelity DNA Polymerase can promote the annealing between the template and primers efficiently. In general, the annealing temperature is the Tm of primers. If required, the temperature gradient can be established to determine the optimal temperature for binding the primer to the template. Long annealing time may disperse amplification products.
- c. Long extension time can improve the yield of amplification products.
- d. To prevent introducing nontarget mutations, we recommend that the amplification cycle be <35. If the amplification efficiency is good, we recommend the amplification cycle to be <30.

After the PCR reaction, a small amount of the amplification product is subjected to gel electrophoresis. If the target plasmid is correctly amplified, please continue with the next step.

3. Amplification products are digested by DpnI to remove the methylated plasmid.

The amplification product of step 2 includes the original template plasmid; thus, we must digest the product with DpnI before recombination cyclization to prevent false-positive transformants after transformation. The recommended reaction system is as follows:

Component	Volume
DpnI	1 μl
Amplification product	40-50 µl

Place the reaction mixture at 37°C for 1–2 hours. If the amplification product of step 2 is a single band, DpnI-digested products can be used in subsequent recombination reactions without purification. If the product is not a single band, gel extraction purification should be performed before the next step.

4. Recombination reaction

The 5' ends of forward and reverse primers share a complete reverse complementary sequence; thus, homologous recombination can occur between the 5' and 3' ends of the amplification product catalyzed using Multi-Toolsitase to complete amplification product cyclization. The following components are added sequentially to the bottom of 1.5-mL sterile Eppendorf tubes or PCR tube on ice-water bath. If liquid sticks on the tube walls, please collect the liquid to the bottom of the tube through brief centrifugation.

Component	Volume
ddH <sub>2</sub> O	Up to 20 µl
5x Buffer II	4 μl
DpnI digested fragment AB	x ng
DpnI digested fragment BC	x ng
DpnI digested fragment CA	x ng
Multi-Toolsitase	2 µl

Optimal DNA fragments in the Multi-Toolsitase double-base mutation recombination reaction system

are as follows: a long fragment of 0.03 pmol and a short fragment of 0.06 pmol. The corresponding mass of DNA fragments can be calculated roughly by using the following formula:

Optimal amount of product digested by DpnI =  $[0.02 \times$  the number of target plasmid base] ng (0.03 pmol). For example, if fragments AB, BC, and CA are 1, 2, and 5 kb, respectively, the optimal amounts of fragments AB, BC, and CA digested by DpnI are 20 ( $0.02 \times 1,000 = 20$ ), 40 ( $0.02 \times 2,000 = 40$ ), and 100 ( $0.02 \times 5,000 = 100$ ) ng, respectively.

Note: Too much or too little DNA will reduce cyclization efficiency. Confirm the DNA concentration through gel electrophoresis in advance, and mix the components strictly in accordance with the recommended amount. When the calculated optimal amount is <10 ng or >200 ng, add 10 or 200 ng. When the product digested by DpnI is used directly in the recombination reaction, the product volume should be less than one-fifth of the total volume.

After adding all the components, please mix them through gently pipetting up and down several times and avoid air bubbles (please do not vortex or shake vigorously). Incubate the tube at  $37^{\circ}$ C for 30 min. After the reaction, please place the tube in an ice bath for 5 min. The product can be transformed directly or stored at  $-20^{\circ}$ C.

5. Transformation, plating, and identification of clone

In 20  $\mu$ L of a cooled reaction mixture, add 200  $\mu$ L of competent cells. Mix gently through flicking the tube, and place the tube on ice for 30 min. Incubate it at 42°C for 45–90 s for heat shock. Then, place it in ice-water bath for 2 min. Add 900  $\mu$ L of SOC or LB medium. Again, incubate at 37°C for 10 min and shake (150 rpm) for 45 min at 37°C. Plate 100  $\mu$ L of the bacterial culture on a selective plate. Then, incubate overnight at 37°C.

Note: We recommend using competent cells with a transformation efficiency of  $>10^8$  cfu/µg. Otherwise, centrifuge your bacterial culture at 5,000 rpm for 3 min to collect bacteria, resuspend the bacteria with 100 µL of LB medium, and then plate all the bacterial cells.

0.1	. •
()thar	augaadiana
	SUPPESHOUS

Experiment procedure	Should do	Should NOT do
Selection of the reverse	Try to choose a region containing a	Select the region with a
complementary region	nonrepetitive sequence with evenly	repetitive sequence with high
of primers	distributed GC. When the GC content	GC or AT content.
	of the selected region is 40%-60%,	
	recombination cyclization will reach	
	the maximum efficiency.	

Primer design	Design as shown in Figure 2 or 4.	The reverse complementary
		region is shorter than the
		recommended length or adds
		the wrong sequence.
Experiment scheme	For two mutation sites, if the distance	Ignore the distance of the two
selection	is >50 bp, choose protocol 2, and if	mutation sites, and choose
	the distance is <50 bp, choose	protocol 2.
	protocol 1.	
Plasmid amplification	Please perform highly specific	Amplification product is not
	amplification.	specific with many nonspecific
		products.
Number of templates	Use the minimum number of	Use too many templates.
	templates if it does not affect the	
	amplification yield.	
PCR template should be	Use the host strain with methylase	Use the host strain without
a methylated plasmid	(e.g., Top10, DH5α, and JM109) to	methylase to extract the
	extract the original plasmid.	original plasmid.
Template quality	Long-term storage and repeated	Use plasmids that were stored
	freezing and thawing may cause	for long term and templates
	breakage, open loop, or degradation	that were repeatedly frozen and
	of plasmids. Hence, we recommend	thawed.
	using freshly prepared plasmids as	
	template.	
Purification of DpnI-	If the product is not a single band, gel	Gel extraction operation is not
digested products	extraction operation is required.	performed when the product is
		not a single band.
DNA quantification of	Quantify through agarose gel	Quantify through the
DpnI-digested products	electrophoresis.	absorbance assay.
Preparation of	Prepare reaction in ice-water bath.	Prepare the reaction at room
recombination reaction	Use the recommended optimal	temperature. Use a random
	amount of DNA at an optimal ratio.	amount of DNA. When the
	When the DpnI-digested product is	DpnI-digested product is used
	used directly in the recombination	directly in the recombination
	reaction, the product volume should	reaction, the product volume is

	be less than one-fifth of the total	more than one-fifth of the total
	volume.	volume.
Recombination reaction	Incubate tubes at 37°C for 30 min in a	The reaction temperature is
	PCR instrument or water bath with	higher or lower than 37°C. The
	precise temperature control.	reaction time is more or less
		than 30 min.
Termination of	The tubes should be cooled down in	Tubes at room temperature
recombination reaction	ice-water bath for 5 min immediately	after the reaction.
	after the reaction.	
Transformation	The cooled product should be	The cooled products are placed
	transformed within an hour. The	at room temperature for a long
	product should be kept in an ice-water	time before transformation. For
	bath before transformation. For long-	long-term storage, keep the
	term storage, keep the product at	product at 4°C.
	-20°C.	

## Troubleshooting

Plasmids cannot be amplified:

- 1. Primer design is wrong: please check the primer design.
- 2. The amplification reaction mixture is not correctly prepared: please perform the experiment again.
- 3. The amplification reaction is not optimized: Mg<sup>2+</sup> concentration, enzyme amount, and amplification program can be optimized.
- 4. Template quality is not good: long-term storage and repeated freezing and thawing can cause breakage, open loop, or degradation of plasmids. Please use freshly prepared plasmids as templates. No or few colonies on the plate:
- 1. Competent cells have low efficiency. Use competent cells that are freshly prepared or stored properly, and make sure the transformation efficiency of the competent cells is  $>10^7$  cfu/µg. Please set up a group transformed with plasmid as control to detect the transformation efficiency of competent cells.
- 2. The amount of DNA is insufficient for the recombination reaction or the fragment ratio is inappropriate. Please add the recommended amount of DNA. Please check the concentration of the product digested by DpnI. The absorbance assay method is highly vulnerable to the purity of DNA and the pH of DNA dilutions. The measured values often deviate from the actual DNA concentration. Hence, we strongly recommend measuring the DNA concentration through agarose gel electrophoresis.
- 3. DNA in recombination cyclization is impure, inhibiting the reaction, or the volume of unpurified product digested by DpnI is more than one-fifth of the total volume. Try performing gel extraction of DpnI-digestion products. Avoid using complexing agents (e.g., EDTA) in recombination reaction. Therefore, we recommend dissolving purified DNA in ddH<sub>2</sub>O of pH 8.0 instead of TE buffer.
- 4. Add excessive amount of DNA in competent cells: the DNA volume should not exceed one-tenth the volume of competent cells; otherwise, it will reduce the transformation efficiency.
- 5. Transformation inhibitory effect occurs: High concentration of input DNA can inhibit the transformation. In this case, one-fifth of the DNA should be used for transformation.

The site-directed mutation is incorrectly introduced:

- 1. Primers are not designed correctly. Check the primer design.
- 2. Template plasmids are not methylated. DpnI can recognize only methylated DNA. Please purify the template plasmids from host strains with functional methylases.
- Use excessive plasmids as template. For most plasmids, 1 ng of DNA is sufficient as template for the PCR reaction. Excessive plasmids will lead to incomplete digestion by DpnI, which reduces the successful rate of mutation introduction.

Mutations at nontarget site:

- 1. The template plasmid carries some unknown mutations: confirm the template sequence.
- 2. Too many amplification cycles: to prevent nontarget mutations during amplification, amplification cycles should not exceed 30 when amplification efficiency is good.

Special Remarks

- When choosing the reverse complementary region of primers, please avoid repetitive sequences. When the GC content is 40%–60%, cyclization recombination efficiency is maximized. When the GC content is >70% or <30%, the cyclization efficiency is significantly inhibited.</li>
- 2. The double-base mutation strategy can be used for single-base mutations (one of the two sites do not undergo base modification). Therefore, if amplification cannot be performed in a single-base mutation, use the double-base mutation strategy.

For research use only. Not for diagnostic or clinical use.

BIOTOOLS CO., LTD biotech.comwww.tools-biotech.com +886-2-2697-2697 info@tools-