

KOD One™ PCR Master Mix KOD One™ PCR Master Mix -Blue-

KMM-101NV-5X1ML KMM-101NVS-0.25ML KMM-201NV-5X1ML KMM-201NVS-0.25ML

Store at -20°C

Contents

i	- 4	-	-	•					-					
	[1	- 1		ľ	1	ŧ١	r	n	М	11	C	tı	^	n
		- 1				u		.,	u	·		LI	٠,	

- [2] Components
- [3] Quality testing
- [4] **Primer design**
- [5] **Protocol**
 - 1. Standard reaction setup
 - 2. Cycling conditions
- [6] **Templates**
- [7] PCR products Cleanup
- [8] Cloning of PCR products
- [9] Examples
- [10] Troubleshooting

CAUTION

All reagents in this kit are intended for research purposes. Do not use for diagnosis or clinical purposes. Please observe general laboratory precautions and safety while using this kit. All trademarks, trade names, or company names referenced herein are used for identification only and are the property of their respective owners.

JAPAN

TOYOBO CO., LTD. Tel(+81)-6-6348-3846 www.toyobo.co.jp/e/bio Bio_overseas@toyobo.jp **CHINA**

TOYOBO (SHANGHAI) BIOTECH, CO., LTD. Tel(+86)-21-587949



[1] Introduction

Description

KOD OneTM PCR master Mix and KOD OneTM PCR Master Mix -Blue- are 2 x PCR master mixes based on genetically modified KOD DNA polymerase (UKOD). KOD OneTM series enables fast PCR, which has an extension time of 5 sec./ kb by applying UKOD and a new Elongation Accelerator. In addition, these master mixes provide greater efficiency and elongation capabilities than conventional PCR enzymes. In particular, these show greater amplification success from crude specimens. Furthermore, these master mixes can be applied to amplify from templates containing uracils (dU) or using primers containing inosines (dI) and uracils (dU).

KOD OneTM series contains two types of anti-KOD DNA polymerase antibodies that inhibit the polymerase and 3'→5' exonuclease activities, thus allowing for Hot Start PCR. These master mixes generate blunt-end PCR products because of 3'→5' exonuclease (proof-reading) activity of KOD DNA polymerase.

Features

- Fast

KOD OneTM series can amplify the targets using the following very short conditions:

≤ 1 kb: 1 sec. 1~10 kb: 5 sec./ kb 10 kb~: 10 sec./ kb

The cycling conditions can be set flexibly when various targets having different sizes are amplified.

- Easy to Use

KOD OneTM series contains all reaction components except primers and templates and provide high reproducibility by reducing operations. In addition, KOD OneTM PCR Master Mix -Blue- includes a loading dye (BPB) to allow direct loading onto agarose gels.

- High Fidelity

KOD OneTM series exhibits approximately 80-fold higher fidelity than Taq DNA polymerase. These mixes can be used for various purposes where this would be an advantage (e.g., in the preparation of long target amplicons for sequencing).

- High Efficiency

KOD One^{TM} series is effective for amplification from crude samples (*e.g.*, biological samples, foodstuffs, soil extract, etc.). Various samples or lysates can be used directly as templates.

- Primers or Templates Containing Inosines (dI) or Uracils (dU) Can Be Used

KOD OneTM series can use primers or templates containing inosines (dI) or uracils (dU), whereas conventional high-fidelity PCR enzymes cannot.

- Low bias (Compatible for NGS library amplification)

KOD OneTM series enables uniform amplification even for GC-rich and AT-rich targets. Library amplification bias affects coverage in next-generation sequencing. With its low bias and high-speed amplification capabilities, KOD OneTM series is well-suited for amplifying both short-read and long-read libraries.





[2] Components

KOD One^{TM} series includes the following components for 200 reactions, 50 μl total reaction volume.

<KMM-101NV>

KOD OneTM PCR master Mix 5X1ML

<KMM-201NV>

KOD OneTM PCR master Mix -Blue- 5X1ML

Note:

The reagents can be stored at 4° C for a month. For longer storage, the reagents should be kept at -20° C.

[3] Quality Testing

Quality testing is performed by amplification of a 10 kb target using fast PCR, which has an extension time of 50 sec.

Technical information (data) contained in this instructional manual are for description representation and application of the product and does not constitute any guarantee.

[4] Primer Design

- -Primers should be 22–35 bases with a melting temperature (T_m) over 63°C.
- -Optimal GC content of primers is 45%–60%. The ideal GC contents of the 5' half and the 3' half are 60%–70% and 45%–50%, respectively.
- -The priming efficiency of primers can be promoted by anchoring the 3'end of primers with G or C.
- -Primers should be designed so as not to generate intermolecular secondary structures or primer dimers.
- -Primers for long target amplification should be 25–35 bases with Tm over 65°C.
- -Primers containing inosine (dI) or uracil (dU) can be used and are applicable for many purposes such as metagenomics analyses. The design of primers can be facilitated by use of specific software that allow replacement of thymine (dT) with uracil (dU) or inosine (dI).

The following online tool is recommended for amplification after bisulfate.

MethPrimer, http://www.urogene.org/methprimer/index.html

-The Tm of primers should be calculated using the Nearest Neighbor method. The Tm values in this manual were calculated using this method with the following parameters.

Na⁺ concentration: 50 mM

Oligonucleotide concentration: 0.5 µM



[5] Protocol

1. Standard reaction setup

Before preparing the mixture, all components should be completely thawed, except for the enzyme solution.

Component	Volume	Final Concentration
PCR grade water	Xμl	
KOD One TM PCR Master Mix	25 μl	1×
10 pmol /μl Primer #1	1.5 µl	0.3 μΜ
10 pmol /μl Primer #2	1.5 µl	0.3 μΜ
		Genomic DNA ≤200 ng / 50 μl
Tamplete DNA	Yμl	Genomic DNA ≤200 ng / 50 μl Plasmid DNA ≤50 ng / 50 μl
Template DNA	~	cDNA \leq 750 ng (RNA equiv.) / 50 µl Crude sample \leq 5 µl/50 µl
		Crude sample ≤5 μl/50 μl
Total reaction volume	50 μl	

Notes:

- -Optimal primer concentration is 0.3 μ M. In the case of long targets (\geq 10 kb), reduced primers concentration (0.15 μ M) may give more effective amplification.
- -When PCR yield is low, increased primers concentration (0.5 μM) may give more effective amplification.
- -For PCR reactions, thin-walled tubes are recommended.

2. Cycling conditions

The following cycle is recommended.

3-step cycle	Amplicon size	Amplicon size	Amplicon size	
	≤ 1 kb	1 ~10 kb	≥ 10 kb	<u></u>
Denaturation:	98°C, 10 sec.	98°C, 10 sec.	98°C, 10 sec.	_ - 1
Annealing:	(Tm -5) °C, 5 sec.	(Tm -5) °C, 5 sec.	(Tm -5) °C, 5 sec.	
Extension:	68°C, 1 sec.	68°C, 5 sec. /kb	68°C, 10 sec. /kb	→ 25–45 cycles

Notes:

- -Longer extension time may enhance efficiency. For amplification from a low copy DNA or crude sample, the extension time should be 10 sec. /kb.
- -The extension temperature should be adjusted in accordance with the Tm of primers. If the Tm value of the primer is over 73°C, the extension temperature should be set at 68°C.
- -Poor amplification may be improved by changing the denaturation step to 94°C, 15 sec.



<Other cycles>

When non-specific or smeared bands are observed, the following cycles is recommended.

3-step cycle	Amplicon size	Amplicon size	•
	< 10 kb	≥ 10 kb	
Denaturation:	98°C, 10 sec.	98°C, 10 sec.	25 451
Extension:	68°C, 5 sec. /kb	68°C, 10 sec. /kb	25–45 cycles

Step-down cycle	Amplicon size	Amplicon size	
	< 10 kb	≥ 10 kb	
Denaturation:	98°C, 10 sec.	98°C, 10 sec.	5 cycles
Extension:	74°C, 5 sec./kb	74°C, 10 sec./kb	5 cycles
Denaturation:	98°C, 10 sec.	98°C, 10 sec.	5 cycles
Extension:	72°C, 5 sec./kb	72°C, 10 sec./kb	5 cycles
Denaturation:	98°C, 10 sec.	98°C, 10 sec.	5 cycles
Extension:	70°C, 5 sec./kb	70°C, 10 sec./kb	5 cycles
Denaturation:	98°C, 10 sec.	98°C, 10 sec.	15–30 cycles
Extension:	68°C, 5 sec./kb	68°C, 10 sec./kb	

[6] Template

a. Purified DNA or cDNA

Appropriate template amounts for a 50 µl reaction are summarized in the following table.

		Approved range	Recommended
Genomic DNA	Eukaryotic DNA	1–200 ng	50 ng
	Prokaryotic DNA	0.1–200 ng	10 ng
Plasmid DNA		1 pg-50 ng	10 ng
cDNA		< 750 ng (RNA equiv.)	50 ng (RNA equiv.)
Lambda phage DNA		0.01–10 ng	1 ng

-Contaminating RNA in cDNA inhibits the PCR reaction. PCR should be performed using template DNA containing <750 ng of RNA in a 50 μ l reaction.

-Quality of the template DNA should be checked by electrophoresis. The length and purity of the template DNA affects amplification results.

b. Tissues and cells

When adding biological samples directly to the PCR reaction solution, the following samples can be applied to the 50 μl reaction.



www.toyobo.co.jp/e/bio

Sample	Appropriate template amount	Remarks
E. coli	Picked small amount of cells from colonies	
Yeast	Picked small amount of cells from colonies	
Fungus	Picked small amount of cells from colonies	When reproducibility is not good, suspended cells in TE buffer should be added (2–5 μl)
Cultured c	ells 10 ¹ –10 ⁵ cells/2 μl medium or PBS	
Whole Blo	ood 1–2 µl	
Nail	1 × 1 mm	As the concentration of extracted
Hair root	1–2 cm ►	DNA is low, 35–40 cycles are needed.
Leaf	$2 \times 2 \text{ mm}$	
Milled rice	$0.5 \times 0.5 \text{ mm}$	
Mouse tail	$1 \times 1 \text{ mm}$	On an agarose gel assay, a portion
		of amplicon may remain in the slots.

^{*}In the case of the direct amplification from animal tissues, such as mouse tail, a portion of the amplification product may remain in the gel slot on an agarose gel assay. Adding 10 μ l of 20 mg/ml proteinase K to 50 μ l PCR products prior to the electrophoresis is effective for dissociating the aggregates.

c. Lysate

To make the lysate for PCR, the following methods are recommended. The lysates can be stored at 4° C for several weeks. For long-term storage, the lysates should be stored at -20° C.

<Alkaline lysis method>

The following "Alkaline lysis method" is recommended for rapid preparation of mouse tails or nail lysates suitable for amplification.



^{*}Mouse tail cannot be dissolved completely.



[96-well PCR plate protocol by alkaline lysis method]

The following protocol is useful for the preparation of lysates from a large number of mouse tail samples.

- 1. Transfer mouse tails (ca 3mm) to a 96-well PCR plate.
- 2. Add 180 µl of 50 mM NaOH and vortex.
- 3. Spin down.
- 4. Incubate at 95°C for 10 min using a thermal cycler.
- 5. Add 20 µl of 1M Tris-HCl (pH 8.0) and vortex.
- 6. Spin down.

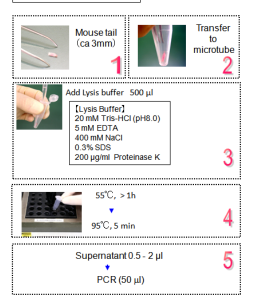
Supernatant 0.5–2 μ l \rightarrow PCR (50 μ l)

<Proteinase K method>

The following "Proteinase K method" is recommended for efficient preparation of mouse tails or nail lysates suitable for amplification. This protocol can also be applied to the following samples.

Mouse tail: 3 mm
Nail: 3 mm
Leaf: 3×3 mm
A rice grain

Proteinase K method





<One step method>

The following "One-step method" is recommended for rapid preparation of a plant tissue lysate suitable for amplification.



*BioTechniques, 19: 394 (1995)

<Homogenization method>

The following "Homogenization method" is recommended for effective preparation of a plant tissue lysate suitable for amplification. This method is effective for amplification of genomic DNA targets.



JAPAN TOYOBO CO., LTD. Tel(+81)-6-6348-3846 www.toyobo.co.jp/e/bio Bio_overseas@toyobo.jp

TOYOBO (SHANGHAI) BIOTECH, CO., LTD. Tel(+86)-21-587949

www.toyobo.co.jp/e/bio



[7] PCR products Cleanup

-PCR products cleanup may be required to remove enzyme, dNTPs, and reaction buffer prior to sensitive downstream applications. When purifying PCR products of KOD OneTM PCR Master Mix - Blue- using magnetic beads, abnormal pelleting of the beads may be observed. The following treatment before purification can be used to improve bead cleanup if this occurs.

<Proteinase K treatment>

Add 1 μ L 10-20mg/mL Proteinase K to 50 μ L of PCR product, mix well and incubate at room temperature for at least 1 min.

<Tween 20 treatment>

Add 1 µL 10% Tween 20 to 50 µL of PCR product.

[8] Cloning of PCR products

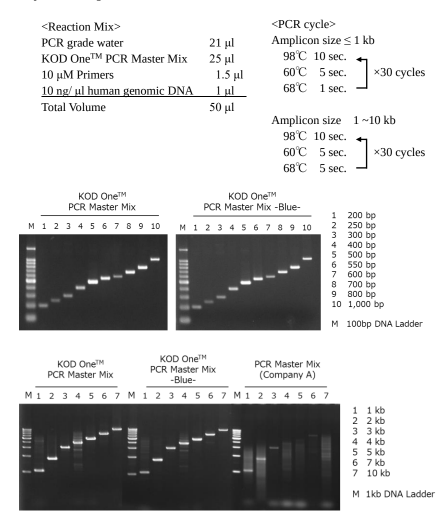
- -KOD OneTM PCR Master Mix / KOD OneTM PCR Master Mix -Blue- generate blunt-end PCR products because of 3'→5' exonuclease (proof-reading) activity. Therefore, PCR products can be cloned using blunt-end cloning methods.
- -PCR products of KOD OneTM PCR Master Mix / KOD OneTM PCR Master Mix -Blue-should be purified prior to restriction enzyme treatments in cloning steps. The 3'→5' exonuclease activity of KOD DNA polymerase remains at the end of the PCR reaction.



[9] Examples

Example 1. Fast PCR

Various targets were amplified with KOD OneTM PCR Master Mix and KOD OneTM PCR Master Mix -Blue- using the fast cycling conditions. KOD OneTM series successfully amplified all targets.

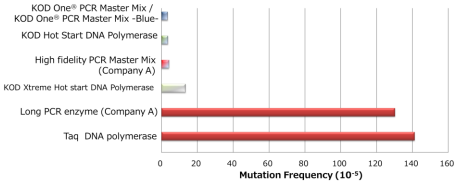




Example 2. PCR error ratio

The error ratio of various PCR enzymes were compared by determining the sequences of the amplicons from human β -globin gene.

KOD One^{TM} PCR Master Mix and KOD One^{TM} PCR Master Mix -Blue- showed excellent fidelity and the mutation frequency was approximately 80 times lower than that of rTaq DNA polymerase.



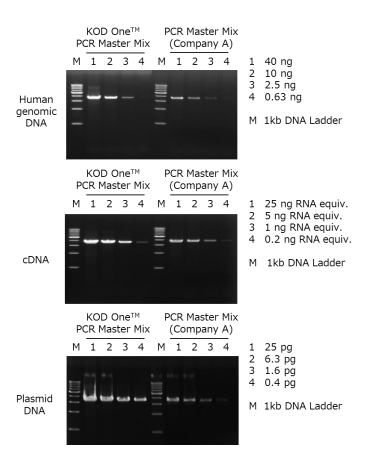
Number of misincorporated beses /100,000 bases



Example 3. Amplification efficiency and sensitivity

The 3.5 kb fragments were amplified from human genomic DNA, cDNA and plasmid DNA. Each PCR reaction was performed according to the recommended conditions. KOD OneTM PCR Master Mix showed higher sensitivity using 5 sec./kb extension time.

<reaction mix=""></reaction>		<pcr cycle=""></pcr>	
PCR grade water	21 μl	98℃ 10 sec.	← ⊓
KOD One TM PCR Master Mix	25 μl	60° C 5 sec.	×30 cycles
10 μM Primers	1.5 µl	68°C 18 sec.	_
Each template DNA	1 μl		
Total Volume	50 µl		

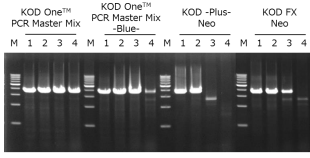




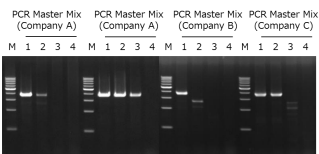
Example 4. Amplification from cDNA

The inhibitory effect of RNA in cDNA was compared using various PCR enzymes. KOD OneTM Master Mix was not susceptible to RNA inhibition, and it was able to amplify targets under high concentrations of cDNA.

<reaction mix=""></reaction>		<pcr cycle=""></pcr>	
PCR grade water	17 μl	98℃ 10 sec.	← 1
KOD One TM PCR Master Mix	25 μl	60° C 5 sec.	×30 cycles
10 μM Primers	1.5 µl	68℃ 18 sec.	
Each cDNA	5 μl		
Total Volume	50 ul		



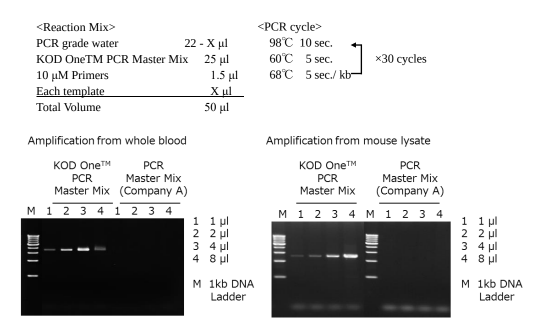
- 125 ng RNA equiv. 2 250 ng RNA equiv. 500 ng RNA equiv.
- 1000 ng RNA equiv.
- M 1kb DNA Ladder





Example 5. Amplification from crude samples

Amplification from whole blood and that from mouse lysate were compared. KOD OneTM PCR Master Mix amplified the targets efficiently.



Example 6. Amplification using degenerate primers containing inosine.

The 2.8 kb fragments were amplified using degenerate primers containing inosine. KOD OneTM PCR Master Mix was able to amplified, whereas conventional high-fidelity PCR enzymes were not.

```
<Reaction Mix>
                                          <PCR cycle>
 PCR grade water
                                            98°C 10 sec.
                                21 µl
 KOD One<sup>TM</sup> PCR Master Mix
                                                             ×30 cycles
                                25 µl
                                            60°C
                                                 5 sec.
 10 μM Primers*
                                            68°C 15 sec.
                                 1.5 µl
 50 ng/μl E.coli genomic DNA
                                 1 μl
 Total Volume
                                50 μl
                                          <Primer sequence>
                                           Fwd: ATGGTICARATHCCICARAAY
                                           Rev: RTGIGCYTGRTCCCARTTYTC
M 1 2 3
                  KOD One™ PCR Master Mix
                   KOD One™ PCR Master Mix -Blue-
                  KOD -Plus- Neo
               M 1kb DNA Ladder
                    *In cases where degenerate primers are used, the primer
                     concentrations should be increased depending on the degree
                     of degeneracy.
```

JAPAN
TOYOBO CO., LTD.
Tel(+81)-6-6348-3846
www.toyobo.co.jp/e/bio
Bio_overseas@toyobo.jp

CHINA

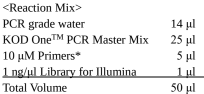
TOYOBO (SHANGHAI) BIOTECH, CO., LTD. Tel(+86)-21-587949



Example 7. Amplification of NGS library

NGS library of Thermus thermophilus genome (GC content 70%) was amplified using KOD OneTM PCR Master Mix and other companies' product, and the amplified libraries were sequenced on Illumina MiSeq. This is the result of examining the GC bias of amplified libraries. The Normalized Coverages (O) close to 1 indicate low bias. KOD OneTM PCR Master Mix showed lower GC bias than PCR Reagent of company B. And KOD OneTM PCR Master Mix shows a high coverage rate. It can be used for sequence analysis with a small number of reads.

The optimal number of cycles may be 1-3 cycles higher or lower depending on the sample type and size distribution of the input DNA. For details of cycle numbers, please refer to the following table.



Comparison of GC bias

KOD One™ PCR Master Mix

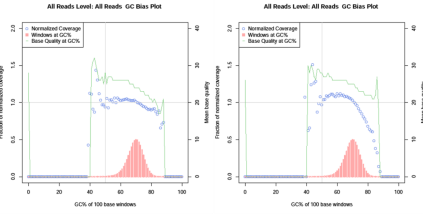
98℃ 10 sec. 60°C 5 sec. ×14 cycles 68°C 1 sec.

<PCR cycle>

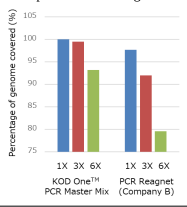
<Primer sequence for TruSeq adapter> Fwd: AATGATACGGCGACCACCGAGATC Rev: CAAGCAGAAGACGGCATACGAG

PCR Reagent (Company B)

All Reads Level: All Reads GC Bias Plot



Comparison of coverage



Recommended cycle numbers

Input DNA	Cycle numbers
1 μg	0-1
500 ng	1-2
100 ng	4-5
50 ng	5-6
10 ng	8-10
1 ng	13-15
0.25 ng	16-18

JAPAN TOYOBO CO., LTD. Tel(+81)-6-6348-3846 www.toyobo.co.jp/e/bio Bio_overseas@toyobo.jp

CHINA

TOYOBO (SHANGHAI) BIOTECH, CO., LTD. Tel(+86)-21-587949



[10] Troubleshooting

Symptom	Cause	Solution
	Cycling condition is	Increase the extension time up to 10 ~30 sec./ kb.
	unsuitable.	Increase the number of cycles by 2–5 cycles.
		Lower annealing temperature in the 3-step cycle
		decrements up to Tm-7–10°C.
	Template DNA is not good	Increase the amount of template DNA.
	in quality and /or quantity.	Decrease the amount of template DNA to reduce
		the contaminated PCR inhibitors.
		Use purified templates.
No PCR product / low yield		Degrade or eliminate RNA in the DNA sample.
	Primer is not good.	Decrease the primer concentration from 0.3 µM to
		0.15 μM. This solution may be especially effective
		for the amplification of targets over 10 kb.
		Increase the primer concentration from 0.3 µM to
		0.5 μM. This solution may be especially effective
		for amplification from low copies template.
		Use fresh primers.
		Redesign primers.
	Cycling conditions are	Change from 3-step cycling to 2-step cycling.
	unsuitable.	Change from 2-step cycling to step-down cycling.
		Decrease the number of cycles by 2–5 cycles.
	Too much template DNA.	Reduce the amount of template DNA.
Smearing / Extra band	Quality of primers is not	Use fresh primers.
	sufficient.	Redesign primers.
		(Longer primers (25-35 mer) may eliminate
		smearing or extra bands)
	Too much enzyme	Reduce enzyme to 0.5–0.8U/50 µl reaction.
Poor TA cloning efficiency	PCR products have blunt	Clone the PCR products according to general
	ends.	blunt-end cloning guidelines.