

# LightNing® Dpnl

REF: EG15585S





Isoschizomers\*: Mall

\*Isoschizomers may have different methylation sensitivities.

### **Storage Condition**

**-20°**C

### Components

Components	Amount
LightNing® DpnI (20 U/µI)	50 µl (1000 U)
10× CutOne® Buffer	1 ml
10× CutOne® Color Buffer	1 ml

### **Description**

LightNing® enzymes are a series of engineered restriction enzymes that are capable of fast DNA digestion. All LightNing® enzymes show superior activity in the universal CutOne® and CutOne® Color Buffer, and are able to digest DNA in 5~15 minutes. This enables any combination of restriction enzymes to work simultaneously in one reaction tube and eliminates the need for sequential digestions. LightNing® enzymes have passed multiple strict quality controls, and can be used to digest plasmid, genomic and viral DNA as well as PCR products.

CutOne® Color Buffer includes a density reagent along with red and yellow tracking dyes that allow for direct loading of the reaction mixtures on a gel. The red dye of the CutOne® Color Buffer migrates with 2.5 kb double-strand DNA fragments in a 1% agarose gel, and the yellow dye migrates with 10 bp double-strand DNA fragments in a 1% agarose gel.

#### **Recommended Reaction Conditions**

1× CutOne® Buffer; Incubate at 37°C:

Refer to "Protocol for Fast DNA Digestion" for reaction setup.

#### **Heat Inactivation**

Incubation at 80°C for 20 minutes.

### **Quality Control**

#### **Functional Test**

A 20  $\mu$ l reaction in CutOne® Buffer containing 1  $\mu$ g of pUC19 DNA and 1  $\mu$ l of LightNing® DpnI incubated for 15 minutes at 37°C results in complete digestion as determined by agarose gel electrophoresis.

#### Prolonged Incubation / Star Activity Assay

A 20  $\mu$ I reaction in CutOne® Buffer containing 1  $\mu$ g of pUC19 DNA and 1  $\mu$ I of LightNing® DpnI incubated for 3 hours at 37 °C results in a DNA pattern free of detectable nuclease degradation as determined by agarose gel electrophoresis. Longer incubation may result in star activity.

#### Non-specific Endonuclease Activity

A 20  $\mu$ I reaction in CutOne® Buffer containing 1  $\mu$ g of supercoiled plasmid and 1  $\mu$ I of LightNing® DpnI incubated for 4 hours at 37°C results in <10% conversion to the nicked or linearized form as determined by agarose gel electrophoresis.

### **Icon Descriptions**

- This enzyme will digest unit substrate in 5~15 minutes under recommended reaction conditions.
- $\boxed{37}$  The enzyme's optimum reaction temperature is  $37^{\circ}\text{C}$ .
- ©pG Cleavage with this restriction enzyme may be blocked or impaired when the substrate DNA is methylated by the CpG methylase.
- EB Cleavage with this restriction enzyme may be blocked or impaired when the substrate DNA is methylated by the EcoBI methylase.
- The enzyme can be heat inactivated at by incubation 80°C for 20 minutes.
- 3 hours incubation do not show star activity, but longer incubation may result in star activity.



### Method of application

#### 1. Protocol for Fast DNA Digestion

① Combine the following reaction components on ice in the order indicated:

	Plasmid DNA	Genomic DNA
$ddH_2O$	15 µl	30 µl
10× CutOne® Buffer or 10× CutOne® Color Buffer	2 μΙ	5 µl
DNA	2 μl (up to 1 μg)	10 µI (5 µg)
LightNing <sup>®</sup> DpnI	1 µl	5 µl
Total	20 μΙ	50 μΙ

- 2 Mix gently and spin down;
- 3 Incubate at 37°C for 15 minutes (plasmid DNA) or for 15~30 minutes (PCR product) or for 30~60 minutes (genomic DNA);
- ④ Optional: Inactivate the enzyme by heating for 20 minutes at 80°C;
- ⑤ If the CutOne® Color Buffer was used in the reaction, load an aliquot of the reaction mixture directly on a gel.

#### 2. Double and Multiple Digestion of DNA

- $\ensuremath{\textcircled{1}}$  Use 1  $\mu l$  of each enzyme and scale up the reaction conditions appropriately;
- 2 The combined volume of the enzymes in the reaction mixture should not exceed 1/10 of the total reaction volume;
- ③ If the enzymes require different reaction temperatures, start with the enzyme that requires a lower temperature, then add the second enzyme and incubate at the higher temperature.

#### 3. Scaling up Plasmid DNA Digestion Reaction

DNA	1 µg	2 µg	3 µg	4 µg	5 µg
LightNing® DpnI	1 μΙ	2 μΙ	3 μΙ	4 μΙ	5 μΙ
10× CutOne® Buffer or 10× CutOne® Color Buffer	2 µl	2 μΙ	3 μΙ	4 μΙ	5 μΙ
Total	20 μΙ	20 μΙ	30 µl	40 μΙ	50 µl

Note: Increase the incubation time if the total reaction volume exceeds 20  $\mu$ l.

#### 4. Digesting Plasmid Templates in PCR Products

Add 1  $\mu$ I of LightNing<sup>®</sup> DpnI to 50  $\mu$ I of PCR products, mix well, incubate at 37°C for 60 minutes, and then heat-inactivate at 80°C for 20 minutes. The resulting product can be used for downstream transformation experiments.

### **Number of Recognition Sites in DNA**

λDNA	ФХ174	pBR322	pUC57	pUC18/19	SV40	M13mp18/19	Adeno2
116	0	22	15	15	8	7	87

### **Methylation Effects on Digestion**

Dam	Dcm	CpG	EcoKI	EcoBI
No effect	No effect	Impaired	No effect	Impaired

### **Activity in Different Buffers\***

CutOne® Buffer		Thermo Scientific	NEB	Takara
		FastDigest Buffer	rCutSmart™ Buffer	QuickCut™ Buffer
Activity	100%	100%	100%	100%

<sup>\*</sup>The activity data come from the functional test described above.

## Activity of DNA Modifying Enzymes in CutOne® and CutOne® Color Buffers

EG15208S Alkaline Phosphatase (Fast)	100%
EG15205S T4 DNA Ligase (Fast)*	100%

<sup>\*</sup>ATP is required for T4 DNA Ligase activity.