

Restriction Endonuclease Mlu NI (Bal I)

From Micrococcus luteus N

Cat. No. 11 526 430 001

200 units (10 U/μl)



(Ii Version 07)
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Store at -15 to -25°C

Stability/Storage

The undiluted enzyme solution is stable when stored at -15 to -25° C until the control date printed on the label. Do not store below -25° C to avoid freezing.

Sequence specificity

Mlu NI recognizes the sequence TGG/CCA and generates fragments with blunt ends (1).

Compatible ends

The enzyme generates compatible ends to any blunt

Isoschizomers

Mlu NI is an isoschizomer to Bal I and Msc I.

Methylation sensitivity

Mlu NI is inhibited by methylation of either of two C residues

Storage buffer

20 mM Tris-HCl, 300 mM KCl, 1 mM EDTA, 10 mM 2-Mercaptoethanol, 0.05 polydocanol, 50% Glycerol (v/v), pH approx. 8.0 (at 4° C).

Suppl. Incubation buffer, 10x

330 mM Tris-acetate, 660 mM K-acetate, 100 mM Mg-acetate, 5 mM Dithiothreitol, pH 7.9 (at 37° C), (= SuRE/Cut Buffer **A)**

Activity in SuRE/Cut Buffer

Bold face printed buffer indicates the recommended buffer for optimal activity:

Α	В	L	M	Н
100%	0-10%	10-25%	10-25%	0-10%

Incubation temperature

System

37°C

Unit definition

One unit is the enzyme activity that completely cleaves 1 μ g λ $dam^ dcm^-$ DNA in 1 h at **37° C** in a total volume of 25 μ l SuRE/Cut buffer **A**.

Typical experiment

Component	Final concentration
DNA	1 μg
10 × SuRE/Cut Buffer A	2.5 μl
Repurified water	Up to a total volume of 25 μl
Restriction enzyme	1 unit

Incubate at 37°C for 1 h.

Heat inactivation

Mlu NI is completely inactivated by incubation at 65° C for 10 min.

Number of cleavage sites on different DNAs (2):

λ	Ad2	SV40	Φ X174	M13mp7	pBR322	pBR328
18	17	0	0	1	1	1

PFGE tested

Mlu NI has been tested in Pulsed-Field-Gel-Electrophoresis (test system bacterial chromosomes). For cleavage of genomic DNA (E. coli dam dcm), embedded in agarose for PFGE analysis, 10 units of enzyme/ μg DNA and 4 h incubation time at 37° C are recommended.

Ligation and recutting assay

 $\it Mlu$ NI fragments obtained by complete digestion of 1 $\mu g \; \lambda \; dam^- dcm^-$ DNA are ligated with 1 U T4-DNA ligase in a volume of 10 μl by incubation for 16 h at 4° C in 66 mM Tris-HCl, 5 mM MgCl $_2$, 5 mM Dithiothreitol, 1 mM ATP, pH 75 (at 20° C) resulting in >85 % recovery of $\lambda \; dam^- dcm^-$ DNA.

Subsequent re-cutting with \emph{Mlu} NI yields > 95% of the typical pattern of λ $\emph{dam}^-\emph{dcm}^-$ DNA \times \emph{Mlu} NI fragments.

Troubleshooting

A critical component is the DNA substrate. Many compounds used in the isolation of DNA such as phenol, chloroform, ethanol, SDS, high levels of NaCl, metal ions (e.g., Hg²+, Mn²+) inhibit or alter recognition specificity of many restriction enzymes. Such compounds should be removed by ethanol precipitation followed by drying, before the DNA is added to the restriction digest reaction. Appropriate mixing of the enzyme is recommended.

Quality control

Lot-specific certificates of analysis are available at www.roche-applied-science.com/certificates.

Absence of unspecific endonuclease activities

1 μ g λ dam^-dcm^- DNA is incubated for 16 h in 50 μ l SuRE/Cut buffer A with excess of Mlu NI. The number of enzyme units which do not change the enzyme-specific pattern is stated in the certificate of analysis.

Absence of exonuclease activity

Approx. $5 \mu g$ [3 H] labeled calf thymus DNA are incubated with $3 \mu l$ Mlu NI for 4 h at 37° C in a total volume of 100 μl 50 mM Tris-HCl, 10 mM MgCl₂, 1 mM Dithioerythritol, pH approx. 7.5. Under these conditions, no release of radioactivity is detectable, as stated in the certificate of analysis.

References

- 1 Roberts, R. J. (1990) Nucleic Acids Res. 18, Supplement, 2331.
- 2 Rebase The Restriction Enzyme Database:
- http://rebase.neb.com
- 3 Benchmate: http://www.roche-applied-science.com/benchmate

Ordering Information

Roche Applied Science offers a large selection of reagents and systems for life science research. For a complete overview of related products and manuals, please visit and bookmark our home page, www.roche-applied-science.com, and our Special Interest Sites, including "Mapping & Cloning": http://www.restriction-enzymes.com.

The convenient RE Finder Program located on our Bench Mate website, http://www.roche-applied-science.com/benchmate helps you identify the enzymes that will cut your DNA sequence, and displays the names and recognition sequences of enzymes and isoschizomers as well as links to detailed information (e.g. package insert) of the selected restriction enzyme.

Product	Application	Packsize	Cat. No.
Restriction Enzymes	DNA restriction digestion	Please refer to websi	te or catalogue
Rapid DNA Liga- tion Kit	Ligation of sticky- or blunt-ended DNA fragments in just 5 min at 15 - 25 °C.	Kit (40 DNA ligations)	11 635 379 001
T4 DNA Ligase	Ligation of sticky- and blunt- ended DNA fragments.	100 U 500 units (1 U/μl)	10 481 220 001 10 716 359 001
rAPid Phosphatase	Dephosphorylation of 5´-phosphate residues from nucleic acids	1000 U 5000 U	04 898 133 001 04 898 141 001
rAPid Dephos and Ligation Kit	Dephosphorylation of nucleic acids.	40 reactions 160 reactions	04 898 117 001 04 898 125 001
Alkaline Phospha- tase (AP), special quality for molecu- lar biology	Dephosphorylation of 5'-phosphate residues from nucleic acids.	1000 U (20 U/μl)	11 097 075 001
Agarose MP	Multipurpose agarose for analytical and prepara- tive electrophoresis of nucleic acids	100 g 500 g	11 388 983 001 11 388 991 001
Agarose LE	Separation of nucleic acids in the range 0.2 - 1.5 kbp	100 g 500 g	11 685 660 001 11 685 678 001
Agarose Gel DNA Extraction Kit	For the elution of DNA fragments from agarose gels.	1 Kit (max. 100 reactions)	11 696 505 001
High Pure PCR Product Purifica- tion Kit	Purification of PCR or enzymatic modification reaction (e.g. restriction digest)	50 purifications 250 purifications	11 732 668 001 11 732 676 001
SuRE/Cut Buffer Set for Restriction Enzymes	Incubation buffers A, B, L, M and H for restriction enzymes	1 ml each (10× conc. solutions)	11 082 035 001
SuRE/Cut Buffer A	Restriction enzyme incubation	5 × 1 ml (10× conc. solution)	11 417 959 001
SuRE/Cut Buffer B	Restriction enzyme incubation	5×1 ml ($10 \times$ conc. solution)	11 417 967 001
SuRE/Cut Buffer H	Restriction enzyme incubation	5×1 ml ($10 \times$ conc. solution)	11 417 991 001
SuRE/Cut Buffer L	Restriction enzyme incubation	5 × 1 ml (10× conc. solution)	11 417 975 001
SuRE/Cut Buffer M	Restriction enzyme incubation	5×1 ml ($10 \times$ conc. solution)	11 417 983 001
Water, PCR Grade	Specially purified, double-distilled, deionized, and	100 ml (4 vials of 25 ml) 25 ml	03 315 843 001 03 315 932 001
	autoclaved	(25 vials of 1 ml) 25 ml (1 vial of 25 ml)	03 315 959 001
BSA, special qual- ity for molecular biology	Maintaining enzyme stability	20 mg (1 ml)	10 711 454 001

Printed Materials

You can view the following manuals on our website:
Lab FAQS "Find a Quick Solution"
Restriction Enzyme Ordering Guide
Molecular Weight Markers for Nucleic Acids

Changes to previous version

Update of quality control.

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Regulatory Disclaimer

For life science research only. Not for use in diagnostic procedures.

Commonly used bacterial strains

Strain	Genotype
BL21	<i>E.</i> coli BF^- dcm omp T hsd $S(r_{B^-}m_{B^-})$ gal (Studier, F.W. et al (1986) <i>J. Mol. Biol.</i> , 189 , 113.)
C600 ^e	supE44 hsdR2 thi-1 thr-1 leuB6 lacY1 tonA21; (Hanahan, D. (1983) J. Mol. Biol. 166 , 557.)
DH5α	supE44 Δ(lacU169 (φ80dlacZΔM15) hsdR17 recA1 endA1 gyrA96 thi-1 relA1; (Hanahan, D. (1983) J. Mol. Biol. 166 , 557.)
HB101	supE44 hsdS20 recA13 ara-14 proA2 lacY1 galK2 rpsL20 xyl-5 mtl-1; (Hanahan, D., (1983) J. Mol. Biol. 166 , 557.)
JM108	recA1 supE44 endA1 hsdR17 gyrA96 relA1 thi Δ (lac-proAB); (Yanisch- Perron, C. et al., (1985) Gene 33 , 103.)
JM109	recA1 supE44 endA1 hsdR17 gyrA96 relA1 thi Δ (lac-proAB) F'[traD36proAB ⁺ , lacl ^q lacZ Δ M15]; (Yanisch- Perron, C. et al., (1985) Gene 33 , 103.)
JM110	rpsL (Str ^f) thr leu thi-I lacY galK galT ara tonA tsx dam dcm supE44 Δ (lac-proAB) F[traD36proAB ⁺ , lacf ^q lacZ Δ M15]; (Yanisch- Perron, C. et al., (1985) Gene 33 , 103.)
K802	supE hsdR gal metB; (Raleigh, E. et al., (1986) Proc.Natl. Acad.Sci USA, 83, 9070.; Wood, W.B. (1966) J. Mol. Biol., 16 , 118.)
SURE ^r	recB recJ sbc C201 uvrC umuC::Tn5(kan ^f) lac , Δ(hsdRMS) endA1 gyrA96 thi relA1 supE44 F'[proAB ⁺ lacI ^q lacZΔM15 Tn10 (tet'); (Greener, A. (1990) Stratagies, 3 , 5.)
TG1	supE hsd $\Delta 5$ thi Δ (lac-proAB) F[traD36proAB ⁺ , lacl ^q lacZ Δ M15]; (Gibson, T.J. (1984) PhD Theses. Cambridge University, U.K.)
XL1-Blue ^r	supE44 hsdR17 recA1 endA1 gyrA46 thi relA1 lac F'[proAB ⁺ , lacl ^q lacZ∆M15 Tn10 (tet ^f); (Bullock et al., (1987) BioTechniques, 5, 376.)

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