



For research use only

ISO9001

1Kbp plus 100bp DNA Ladder Marker, Concentrated

Product	Conc.	Cat. No.	Remarks
1Kbp plus 100bp DNA Ladder Marker, Ready-to-use	500 µl (83 µg)	EBM-1003C	Concentrated type in TE buffer

Description

The 1Kbp plus 100bp DNA ladder marker is a mixture of double-stranded DNA fragments for broad range determination of the exact size of DNA fragments. The 1Kbp plus 100bp DNA ladder marker consists of total 18 DNA fragments ranging in size from 100 to 10,000 bp. Ruler bands (500, 1,000, 1,500, 5,000 bp) are more brighter than the other bands and helpful for easy size discrimination on the gel electrophoresis. This ladder marker can be stained by ethidium bromide or any other known DNA staining methods.

The 83 µg of marker DNA is contained in 500 µl TE buffer. Before start, optimal concentration and loading volume should be empirically determined by users to get the best result. This product is supplied with 1 ml of 6x gel loading buffer to dilute marker DNA.

Storage Buffer

- Marker DNA : 83 µg in 500 µl TE buffer
- 6x Gel loading buffer : 60 mM Tris-HCl, pH8.0, 6 mM EDTA, 30% Glycerol, 0.03% Bromophenol Blue, and 0.03% Xylene Cyanol

Recommended Storage Condition

- -20°C for 2 year
- 4°C for 6 months
- Room temperature (20-25°C) for 2 months

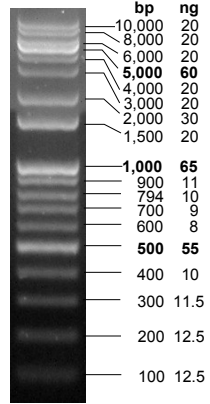
Usage Information

- Recommended loading : 200-500 ng/well according to the gel size, well size, and running length.
- Range : 100 – 10,000 bp
- Number of bands : 18

Cautions

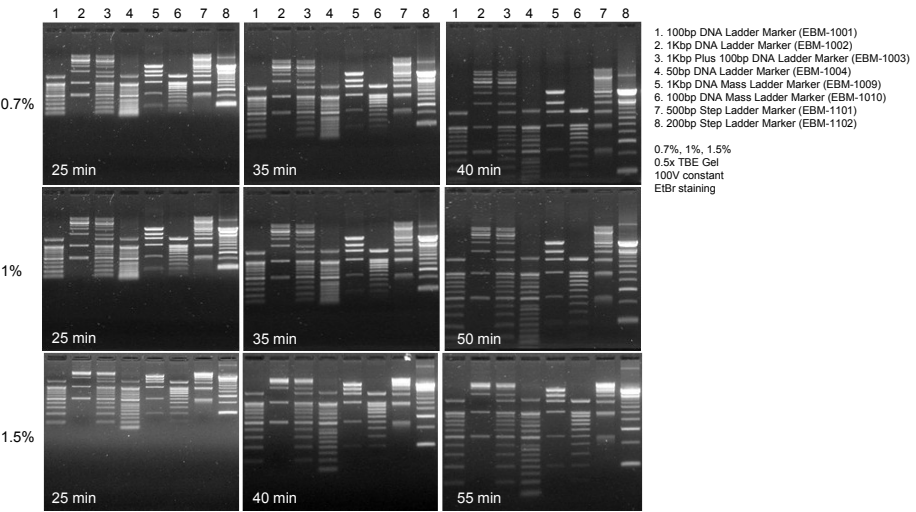
- Always use the fresh tip to take out marker solution.
(If you do not, trace amount of contaminated DNases from buffer tank may degrade marker DNA rapidly)
- Don't boil the product.
- Use appropriate % of gels for separation of 100 to 10,000 bp sizes
(0.7 to 1.5% agarose gel is recommended)
- Confirm that the concentration of DNA staining dye is optimal before use.
(Breakage or suboptimal concentration of ethidium bromide in gel is a main cause of low estimation of marker concentration or your DNA. 5 ng of DNA should be seen in normal condition)
- Loading volume and concentration should be optimized by gel size, well size, and running length.
- Low sized DNA bands can be gradually disappeared as running is progressing.
(This is because some DNA is getting out from gel to buffer during horizontal electrophoresis, not because the DNA concentration is incorrect. This will be the same for your DNA)

1Kbp plus 100bp DNA Ladder Marker



415 ng/lane ;
1.5% agarose in 0.5x TAE, stained with ethidium bromide

Migration Patterns in Different % of Agarose Gels



Recommended Gel Percentages for Separation of Linear DNA

Agarose Gel, %	Range of Separation, bp	Polyacrylamide Gel, %	Range of Separation, bp
0.5	1,000 - 30,000	3.5	100 - 1,000
0.7	800 - 12,000	5	80 - 500
1	500 - 10,000	8	60 - 400
1.2	400 - 7,000	12	40 - 200
1.4	200 - 4,000	20	5 - 100
2	50 - 2,000		

DNA Size Migration with Sample Loading Dyes

Agarose Concentration, %	Xylene cyanol FF	Bromophenol blue	Orange G
0.7 - 1.7	~4000 bp	~300 bp	~50 bp
2.5 - 3.0	~800 bp	~100 bp	~30 bp

Composition of Gel Electrophoresis Buffers

Buffer	Working Concentration	Stock Concentration (per Liter)		
Tris-acetate (TAE)	1x	20 mM Tris-acetate	Tris base	96.9 g
		1 mM EDTA	Glacial acetic acid	22.84 ml
			0.5 M EDTA (pH8.0)	40 ml
Tris-borate (TBE)	0.5x	45 mM Tris-borate	Tris base	108 g
		1 mM EDTA	Boric acid	55 g
			0.5 M EDTA (pH8.0)	40 ml