

日本語データシート



<u>Product Name</u>: DynaMarker® Prestain Marker for RNA High

Code No.: DM260

Range: 200-8,000 bases

Size: 180 µl (30 loadings)

Storage: store at -80 °C

Description:

The ^{DynaMarker®} Prestain Marker for RNA High is a visible molecular weight marker for ssRNA, consisting of six colored (blue and purple) nucleic acids. The six colored bands (apparent molecular weights are 200, 500, 1,000, 2,000, 4,000 and 8,000 bases) are suitable for monitoring denaturing agarose gel electrophoresis and blotting onto membranes. The ^{DynaMarker®} Prestain Marker for RNA High shows the same mobility as that of the ^{DynaMarker®} RNA High (code # DM160) on denaturing agarose gel electrophoresis (>90 % accuracy, see table 1 and figure 2). The ^{DynaMarker®} Prestain Marker for RNA High is supplied in a ready-to-use mixture without requiring heating or the addition of a denaturing agent before use.



40 mM MOPS (pH 7.0), 10 mM AcONa, 1 mM EDTA • 2Na, 10 % Glycerol

Quality Control:

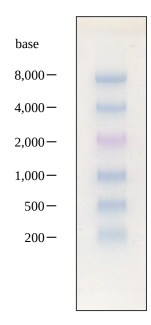
After 24 hrs incubation of the $^{DynaMarker \circledast}$ Prestain Marker for RNA High at 37 $^{\circ}$ C, no visible degradation of the marker is observed in 1 % agarose -2.2 M formaldehyde gel electrophoresis.

Recommended loading volumes:

Comb	Load volume	
4 ~ 6 mm	4 ~ 6 μl	
>6 mm	>6 µl	

Note:

- For accurate electrophoretic determination of molecular weights, the ^{DynaMarker®} RNA High (code # DM160) or ^{DynaMarker®} RNA Easy Measurement N (code # DM170) should be used.
- The migration of the ^{DynaMarker®} Prestain Marker for RNA High has been optimized to use 0.8 1.5 % of agarose gel concentration (see table 1).
- This product is not for acrylamide gel electrophoresis.
- Particularly avoid freeze thaw cycle.



 $^{DynaMarker \circledast}$ Prestain Marker for RNA High

Figure 1: Electrophoresis profile of DynaMarker® Prestain

Marker for RNA High (6 µl) on 1 %

agarose – 2.2 M forumaldehyde gel / 1 ×

MOPS buffer[®] as running buffer.

Same as MESA Buffer

Ver.1.6

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concentration of agarose

_		0.8 %	1.0 %	1.5 %
-	8000 base	96.5	94.8	90.4
	4000	98.5	100.0	92.2
gh	2000	104.4	102.8	101.6
RNA High	1000	103.4	103.2	101.4
×	500	106.4	102.0	103.1
щ	200	102.7	100.0	108.7

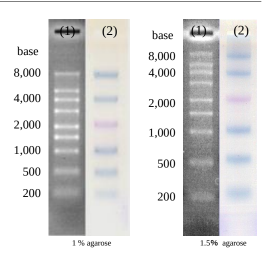


Figure 2: Electrophoresis profile of ^{DynaMarker®} RNA High (1) and ^{DynaMarker®} Prestain Marker for RNA High (2) on 1 % and 1.5 % agarose.

Recommended usage:

The DynaMarker® Prestain Marker for RNA High is suitable for monitoring denaturing agarose gel electrophoresis and blotting onto membrane. One example is shown below:

•Electrophoresis and blotting of ^{DynaMarker®} Prestain Marker for RNA High

1) Preparation of 0.8 % agarose – 2.2 M formaldehyde gel

Agarose	0.8 g
$10 \times MOPS$ (= MESA Buffer)	10 ml
deionized formaldehyde	18 ml
RNase free water	72 ml
total	100 ml

Dissolve the agarose by boiling in a microwave oven. Cool the solution to 55 $^{\circ}$ C and add 10 ml of $10 \times MOPS$ buffer and 18 ml of deionized formaldehyde. In a fumehood, cast an agarose gel with slots formed by a 4~6 mm comb. Remove the comb, place the gel in the gel tank, and add sufficient $1 \times MOPS$ running buffer to cover to a depth of \sim 1 mm.

2) Loading and electrophoresis.

Thaw the $^{DynaMarker@}$ Prestain Marker for RNA High completely before use. Load your denatured RNA sample and 6 μl of $^{DynaMarker@}$ Prestain Marker for RNA High (use a 4~6 mm comb) on to a well and run the gel using $1\times MOPS$ electrophoresis buffer at 4~5 V / cm.

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- 3) Transfer the ^{DynaMarker®} Prestain Marker for RNA High and RNA from the gel to positively charged nylon membranes. (see figure 3 and 4)
 - 3-1) Place the gel in RNase-free glass dish, and rinse (for formaldehyde removal) with several changes of sufficient deionized water to cover the gel.
 - 3-2) Add ~10 gel volumes of 3 M NaCl / 10 mM NaOH (transfer buffer) to the dish and soak for 30 min.
 - 3-3) Cut a piece of nylon membrane slightly larger than the gel. Soak the membrane and two sheets of blotting paper of appropriate size in 10× SSC for at least 5 min.
 - 3-4) Place the support (e.g. oblong sponge) in a glass or a plastic dish. Fill the dish with enough transfer buffer (soak the support about half-submerged in buffer.).
 - 3-5) Place the gel on the support in inverted position so that it is centered on the wet blotting paper.
 - 3-6) Place the wet nylon membrane on top of the gel. (! notice: Remove air bubbles.)
 - 3-7) Place the wet blotting paper on the top of the wet nylon membrane. (! notice: Remove air bubbles.)
 - 3-8) Cut a stack of paper towels (5~8 cm high), and place the towels on the blotting papers.
 - 3-9) Put a glass plate on the top of the stack and weight it down with an about 400 g weight.
 - 3-10) Allow upward transfer of RNA to occur for 1hr.
 - 3-11) Transfer the membrane to a glass tray containing 6× SSC, and rinse for 5 min.
 - 3-12) Remove the membrane from the 6× SSC and allow excess fluid to drain away. Then dry the membrane on blotting paper for a few minutes.
 - 3-13) Fix the RNA to the membrane with a UV-crosslinker.
 - 3-14) Cut off the marker lane.
 - 3-15) Carry out northern hybridization.

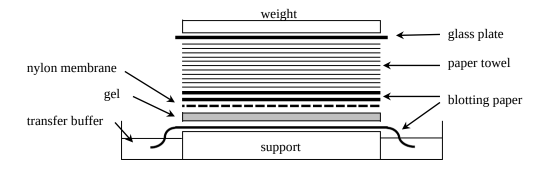


Figure 3: Upward Capillary Transfer.

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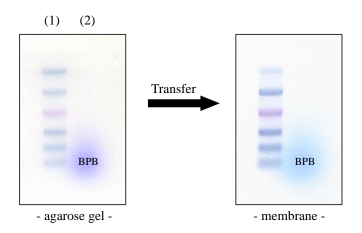


Figure 4: Left : Electrophoresis profile of $^{DynaMarker@}$ Prestain Marker for RNA High (1) and RNA sample (2) on 0.8 % agarose - 2.2 M formaldehyde gel / 1 \times MOPS buffer.**

Right: Blotting of (1) and (2) onto nylon membrane.

XSame as MESA Buffer

References:

- Joseph Sambrook, and David W. Russell (2001) Molecular Cloning: A Laboratory Manual, 3rd ed.,
 Cold Spring Harbor Laboratory Press.
- Frederick M. Ausubel, Roger Brent, Robert E. Kingston, David D. Moore, J. G. Seidman, John A. Smith, and Kevin Struhl (1994—) Current Protocols in Molecular Biology, John Wiley & Sons, Inc.

Related Products:

DM253	DynaMarker® Prestain Marker for Small RNA Plus Prestained marker for RNA(20-100 bases)
DM192	DynaMarker® Small RNA II RNA marker (20-100 bases)
DM152	DynaMarker® RNA Low II RNA marker (20-500 bases)
DM170	DynaMarker® RNA High for Easy Electrophoresis RNA marker (200-8,000 bases) & RNA Loading Buffer. RNA sample can be electrophoresed on non-denaturing agarose gel as well as on denaturing agarose gel with this Loading Buffer.
DM660	DynaMarker® Protein MultiColor Stable II Prestained protein marker. Stable at 4°C.