

 Product Name :
 DynaMarker
 Prestain Marker for Small RNA Plus

 Code No. :
 DM253S

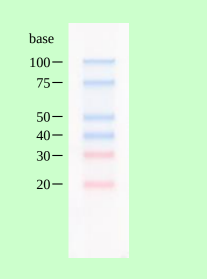
 Range :
 20 - 100 bases

 Size :
 75 μl (15 loadings)

 Storage :
 store at -20 °C

Description:

The ^{DynaMarker} Prestain Marker for Small RNA Plus consists of six prestained single-strand (blue and red) nucleic acids (apparent molecular weights are 20, 30, 40, 50, 75 and 100 bases) and it is visible during electrophoresis. The ^{DynaMarker} Prestain Marker for Small RNA Plus is suitable for monitoring denaturing polyacrylamide gel electrophoresis and blotting onto membranes. The apparent sizes of bands in ^{DynaMarker} Prestain Marker for Small RNA Plus are in excellent agreement with sizes of non-stained RNAs, 20, 30, 40, 50, 75 and 100 bases in length (about 95 % accuracy, see table 1 and figure 2). The ^{DynaMarker} Prestain Marker for Small RNA Plus is supplied in a ready-to-use mixture and doesn't require heating or addition of a denaturing agent before use.



DynaMarker® Prestain Marker for Small RNA Plus

Figure 1. Electrophoresis profile of ^{DynaMarker} Prestain Marker for Small RNA Plus (5 μl) on 10 % polyacrylamide – 7.5 M urea gel / 1 × TBE buffer as running buffer.

Storage buffer :

2 mM Tris-HCl (pH 8.0), 8 mM EDTA, 78 % Formamide

Quality Control:

After 24-hrs incubation of the ^{DynaMarker} Prestain Marker for Small RNA Plus at 37 $^{\circ}$ C, no visible degradation of the marker is observed in 10 % polyacrylamide – 7.5 M urea gel electrophoresis.

Recommended loading volumes :

Comb	Load volume		
4-10 mm	5-10 µl		
>10 mm	>10 µl		

Note :

- For accurate electrophoretic determination of molecular weights, the ^{DynaMarker} Small RNA II (code # DM192) or ^{DynaMarker} Small RNA II Easy Load (code # DM197) should be used.
- A migration of the ^{DynaMarker} Prestain Marker for Small RNA Plus is optimized to use 10 15 % acrylamide gel electrophoresis (see table 1).
- This product is not for agarose gel electrophoresis.

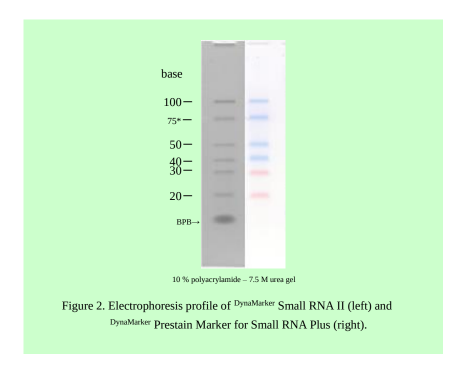


				acrylam	ide concentratio	on (cond	(condition: acrylamide:bis = 29:1, 1× TBE)		
_		5.0 %	7.5 %	10 %	12.5 %	15 %	17.5 %	20 %	
^{larker} Small RNA II + 75 base RNA	100 base	105.6 %	105.6	101.6	98.4	97.2	93.6	92.6	
	75*	106.2	104.7	103.5	99.5	98.5	94.7	92.4	
	50	101.4	101.4	101.1	98.7	97.5	95.0	92.2	
	40	103.1	102.0	103.2	100.8	100.0	97.4	93.9	
	30	91.0	96.9	98.2	98.9	99.2	99.5	98.8	
DynaMarker	20	89.8	95.8	98.2	100.3	101.6	101.4	101.4	

Table 1. This shows apparent molecular weights compared with the ^{DynaMarker} Small RNA II, and suitable acrylamide concentrations for electrophoresis of the ^{DynaMarker} Prestain Marker for Small RNA Plus.

: Recommend : Possible

(* 75 base RNA is from a newly synthesized RNA. A 75 base RNA is not included in ^{DynaMarker} Small RNA II.)



Recommended usage :

The ^{DynaMarker} Prestain Marker for Small RNA Plus is suitable for monitoring denaturing acrylamide gel electrophoresis and blotting onto membrane. One example is shown below:

• Electrophoresis and blotting of DynaMarker Prestain Marker for Small RNA Plus

1) Preparation of 10 % polyacrylamide - 7.5 M urea gel



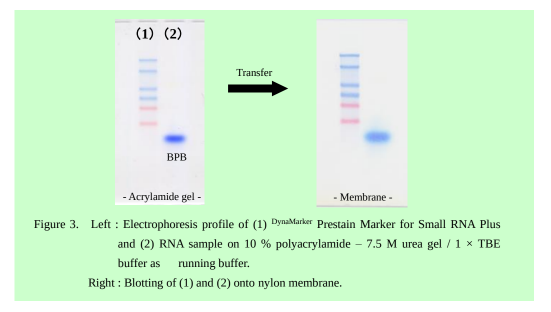
40 % acrylamide : bis solution	5.0 ml
Urea	9.0 g
$10 \times TBE$	2.0 ml
H2O	to 20 ml

After urea is dissolved completely, add 20 μ l of TEMED and 160 μ l of 10 % ammonium persulfate. Mix quickly then pour the gel into the mold of a vertical gel apparatus.

2) Loading and electrophoresis.

Thaw the ^{DynaMarker} Prestain Marker for Small RNA Plus completely before use. Load the denatured RNA sample and 5 μ l of ^{DynaMarker®} Prestain Marker for Small RNA Plus into a well and run the gel using 1 × TBE electrophoresis buffer at 20 – 40 V / cm.

- 3) Transfer the ^{DynaMarker} Prestain Marker for Small RNA Plus and RNA from gel to membrane (figure 3).
 - 3-1) Cut a piece of positive charged nylon membrane slightly larger than the gel. Soak the membrane and four sheets of blotting paper of appropriate size in $0.5 \times \text{TBE}$ buffer.
 - 3-2) Place two sheets of blotting paper on the anode platform of the transfer cell.
 - 3-3) Place the membrane on top of the blotting paper.
 - 3-4) Transfer the gel from the glass plate to the top of the membrane and press out any air bubbles. (*Make sure that there are no air bubbles between the membrane and the gel.)
 - 3-5) Place another two sheets of blotting paper onto the gel and set the cathode assembly.
 - 3-6) Transfer for 30 60 min at 300 mA.
 - 3-7) After ensuring the marker has transferred successfully onto the membrane, remove both paper and gel. Rinse the membrane in $2 \times SSC$.
 - 3-8) Fix the RNA to the membrane with a UV crosslinker.
 - 3-9) Cut off the marker lane.
 - 3-10) Carry out northern hybridization.





<u>References</u>:

- 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.