

# Total RNA Isolation Kit -- Plant





# **Plant Total RNA Isolation Kit**

Storage: Stored at room temperature (15-25°C).

For mat: spin column

For research use only
Sample: up to 100 mg of fresh plant tissue or 25 mg of dry plant tissue.

### Introduction:

The EBL Total RNA Isolation Kit (Plant) provides a fast, simple, and cost-effective method for isolation of total RNA from tissue sample. Detergents and chaotropic salt are used to lyse cells and inactivate RNase. The specialized high-salt buffering system allows RNA species bases to bind to the glass fiber matrix of the spin column while contaminants pass through the column. Impurities are efficiently washed away, and the pure RNA is eluted with EB Buffer without phenol extraction or alcohol precipitation. RNA purified with The Total RNA Isolation Kit is suitable for a variety of routine applications including RT-PCR, cDNA Synthesis, Northern Blotting, Differential display, Primer Extension and mRNA Selection. The entire procedure can be completed within 25-40 minutes.

### About the kits:

Cat. No.	FBRP-0100	
SR Columns	100 pcs	
Collection Tubes	100 pcs	
PRL Buffer	110 ml	
W1 Buffer	45 ml	
W2 Buffer	15 ml	
EB Buffer	10 ml	

# Additional requirements:

- \* Ethanol (96~100%)
- \* Isopropanol
- \* B-mercaptoethanol

# Optional Step < DNA Residue degradation >

### NOTE:

- 1. Add ethanol (96–100%) to Buffer W2 before use (see bottle label for volume).
- 2. If necessary, re-dissolve any precipitate by warming to 37°C.
- 3. Buffers W1 contain irritants. Wear gloves when handling these buffers.

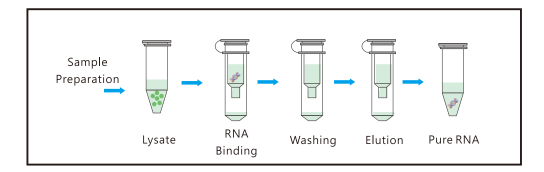
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# Related product and Ordering information:

CAT NO:	Product	Quantity
MGK-T0100	EBL Genomic DNA Isolation Kit (Tissue)	100 rxn.
MGK-T0300	EBL Genomic DNA Isolation Kit (Tissue)	300 rxn.
MGK-C0100	EBL Genomic DNA Isolation Kit((ell/ Blood)	100 rxn.
MGK-C0300	EBL Genomic DNAIsolation Kit (Cell/ Blood)	300 rxn.
MGK-P0100	EBL Genomic DNA Isolation KitP(lant)	100 rxn.
MGK-P0300	EBL Genomic DNA Isolation KitP(lant)	300 rxn.
MPD-01300	EBL Plasmid Miniprep DNA Kit	300 rxn.
MPD-02025	EBL 200PLUS Plasmid Midiprep DNA Kit	25 rxn
MPD-03020	EBL 800PLUS Plasmid Maxiprep DNA Kit	20 rxn
FBRE100	Blood/ cell Total RNA Isolation kit	100 rxn.
FBRT100	Tissue Total RNA Isolation kit	100 rxn.
FBRP	Plant Total RNA Isolation kit	100 rxn.

# Procedure:



### Protocol:

# Step 1. Sample Preparation

- > Cut off 100 mg of fresh plant tissue or 25 mg of dry plant tissue.
- > Grind the sample under liquid nitrogen to a fine power using a mortar and pestle.

# Step 2. Lysis

- > Add 1 ml of PRL Buffer and 10  $\mu$ l of  $\beta$ -mercaptoethanol to the sample in the mortar and grind the sample until it is completely dissolved.
- > Transfer the dissolved sample to a RNase-free 1.5 ml microcentrifuge tube. Incubate at 70°C for 30 minutes. (invert the tube every 10 minutes)
- > Centrifuge at 16,000 x g for 10 minutes. Transfer the supernatant to a new 1.5 ml microcentrifuge tube.

# Step 3. RNA Binding

- > Add 1/2 volume of Isopropanol the sample from Step 2 and shake vigorously.
- > Place a SR Column in a Collection Tube. Apply  $600\mu l$  of the mixture to the SR Column.
- > Centrifuge at 14,000 x g for 30 seconds. Discard the flow-through and place the SR Column in the same Collection tube. Transfer the remaining mixture to the same SR Column.
- > Centrifuge at 14,000 x g for 30 seconds. Discard the flow-through and place the SR Column in the same Collection tube.

# optional DNase treatments can be followed on-column Dnase I digestion to remove unwanted DNA residue.

### Procedure:

- > Add the DNase I incubation mix (80  $\mu$ I, 4U DNase I in the reaction buffer) directly to the SR column membrane center, and place on the benchtop (20–30°C) for 15 min.
- > Following W1 Buffer wash step.

# Step 4. Wash

- > Add 400 µl of W1 Buffer into the SR Column. Centrifuge at 14,000 x g for 30 seconds.
- > Discard the flow-through and place the SR Column back into the same Collection tube.
- > Add 600  $\mu$ l of W2 Buffer (**Ethanol added**) into the SR Column. Centrifuge at 14,000 x g for 30 seconds.
- > Discard the flow-through and place the SR Column back into the same Collection tube.
- > Centrifuge at 14,000 x g again for 2 minutes to remove residual W2 Buffer.

# Step 5. RNA Elution

- > Transfer the dried SR Column to a new 1.5 ml microcentrifuge tube.
- > Add 50-100  $\mu$ I EB Buffer to the center of the SR column matrix and let stand for 2-3 minutes.

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> Centrifuge for 2 minutes at 14,000 x g to elute the purified RNA.

# Troubleshooting

Problem	Comments and suggestions
Degraded RNA / low integrity > RNases contaminantion	Clean everything, use barrier tips, wear gloves and a lab coat, and use RNase-free enzymes, EX: RNase inhibitor.
Low yields of RNA > Incomplete lysis and homogenization	Don`t use more samples than the suggested limit.
>Incorrect elution conditions	Add 100 $\mu$ l of the EB Buffer to the center of each SR Column, let it stand for 2 minutes, and centrifuge at 14,000 x g for 2 minutes.
Inhibition of downstream enzymatic reactions > Presence of ethanol in the purified RNA	Repeat the wash step: Centrifuge at 14,000 x g again for 2 minutes to remove the residual W2 Buffer.

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