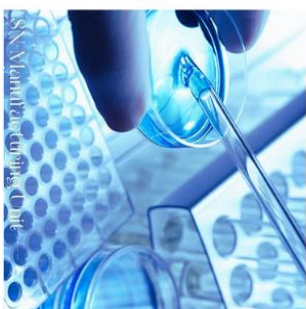




Contact Us -

☎ +91- 80959 61994 / 99005 05076

✉ sales@snlifesciences.com / drmondal@snlifesciences.com

📍 No.57/90, 3rd Road, 4th Phase Bommasandra Industrial Area,
Bengaluru - 560099



2025 - 2026

#startupindia



SN Plant/Feed RNA extraction kit

Column Based



SN PLANT/FEED RNA EXTRACTION KIT (Column Based)

DESCRIPTION:

SN PLANT/FEED RNA Extraction Kit is a specialized tool used to isolate high purity RNA from various PLANT/FEED samples. It is a column based method which is reliable and efficient way to isolate high quality of RNA from PLANT/FEED samples. Kit is essential for various molecular biology and genetic research applications.

CONTENTS OF KIT:

Sl. No	Components	Volume		
		5 Rxn	50 Rxn	100 Rxn
1	RS Buffer	2.5ml	25ml	50ml
2	LS Buffer	3ml	30ml	60ml
3	STB Buffer	2.5ml	25ml	50ml
4	W1 Buffer	1.5ml	15ml	30ml
5	Elution Buffer	250µl	2.5ml	5ml
6	SN Spin Columns	5 No.	50 No.	100 No.

NOTE: Preparation for first use after receiving the kit (Add 100% Ethanol to Wash Buffer)
(5Rxn: 1ml, 50Rxn:10 ml, and 100Rxn: 20ml) mix well and store the buffer at room temperature.

- ❖ If Proteinase K shipped in lyophilized form, upon receiving resuspend with Nuclease free water (5Rxn:100µl,50Rxn: 1ml, 100Rxn: 2ml)
- ❖ If DNase shipped in lyophilized form, upon receiving resuspend with Nuclease free water (5Rxn:50µl, 50Rxn:500µl, 100Rxn: 1ml)
- ❖ **Proteinase -K and DNase store at -20°C either liquid form or lyophilized form.**

REQUIRED MATERIALS NOT PROVIDED

- ✓ 100% Ethanol & 70 % Ethanol
- ✓ Dry bath
- ✓ 1.5ml Centrifuge tubes
- ✓ micro centrifuge
- ✓ Vortex

STORAGE / SHIPPING:

- ❖ Shipped at: Ambient Temperature.
- ❖ Storage: All Buffers can be stored at Room temperature.

SPECIFICATIONS:

Principle	Spin column
Recommended Input Amount: 200mg PLANT/FEED Sample	
Elution Volume : 50µl	
Purity: A260/280 - 1.8±0.1	
Compatible Downstream Application: PCR, qPCR, Sequencing	
Expected Yield : depends upon the sample	

PROCEDURE:

- 1) Weigh 200mg of clean PLANT/FEED sample and transfer to mortar and pestle grind like a fine paste by adding 500µl of RS Buffer.
- 2) Transfer the above mixture to 1.5 ml centrifuge tube using a clean spatula.
- 3) Add 600 µl of LS buffer into the sample along with 20µl of Proteinase K and 10µl of DNase. Vortex for 1min.
- 4) Incubate at 60°C for 15mins with intermittent vortexing for 3-4 times to lysed the cells are completely. Centrifuge the tube for 2mins at 10,000rpm, then transfer the supernatant to a fresh centrifuge tube.
- 5) Add 500µl of 100 % ethanol to the supernatant, keep the tube for 2mins at room temperature, and observe precipitation in the supernatant.
- 6) Transfer the precipitate to the Spin Column, then centrifuge at 15,000 rpm for 2mins and discard the flow-through in the collection tube.
- 7) Add 500 µl of STB buffer to the SN spin column and centrifuge at 15,000 rpm for 1min, discard the flow through into the collection tube.
- 8) Add 500µL of W1 buffer solution to the SN spin column and centrifuge at 15,000 rpm for 1min, discard the flow through into the collection tube.

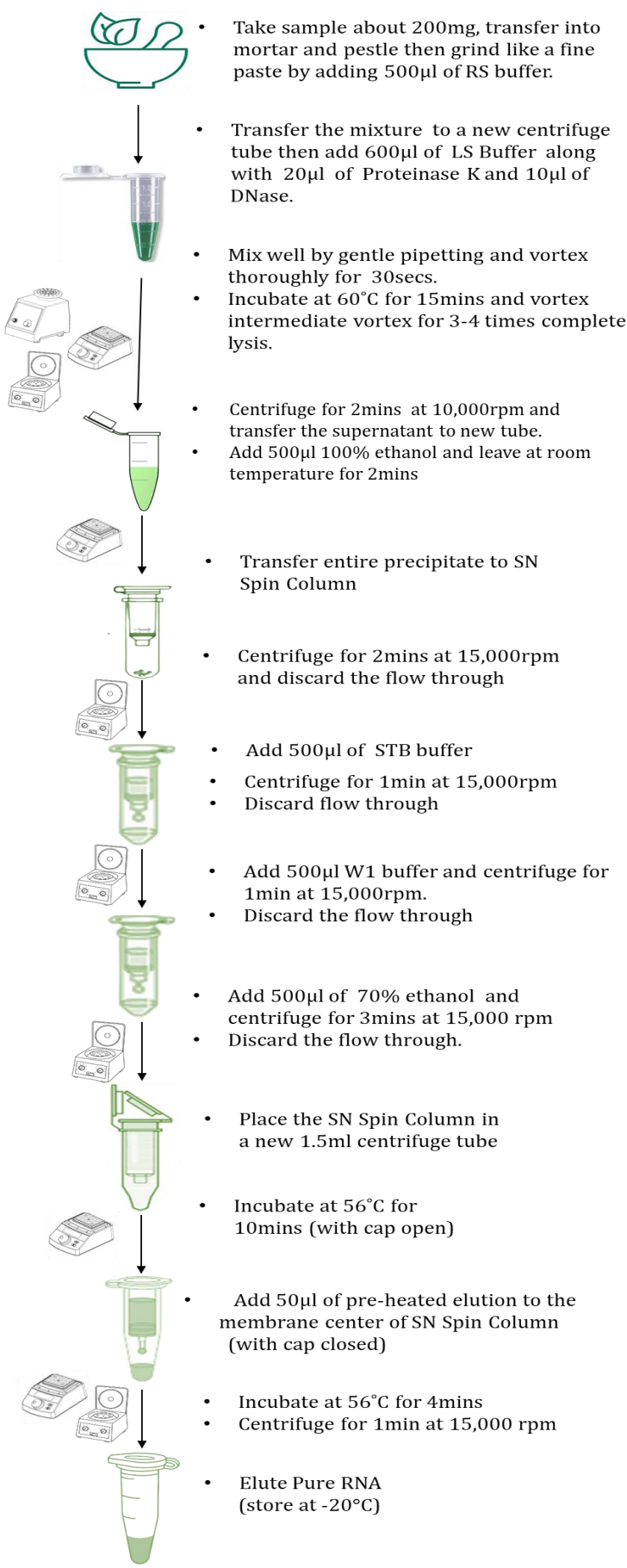
NOTE: Note : (W1 buffer concentration per reaction: W1 buffer - 300µl: 100% ETOH - 200µl)

- 9) Add 500µL of 70 % Ethanol Centrifuge the column at 15,000 rpm for 1min and centrifuge additional 1min.
- 10) Place the SN Spin Column in a new 1.5ml centrifuge tube and incubate for 10mins at 56°C(with cap open). This step is very important to remove the residual of ethanol.

NOTE: Elution should be pre-heated for 10mins before adding into the SN Spin Column.

- 11) Add 50µl of pre-heated Elution to the membrane center of SN Spin Column and incubate for 4mins at 56°C (with cap close).
- 12) Centrifuge the tube at 15,000rpm for 1min to elute pure RNA(store at -20°C)

FLOW CHART



TROUBLE SHOOT

Problems	Possible reasons	Solutions
Low or none recovery of RNA fragment	Weigh 200mg of PLANT/FEED sample	If product is more, then separate it into multiple tubes.
	Elution of RNA fragment is not efficient	Make sure the pH of Elution Buffer or ddH ₂ O is between 7.5-8.0.
Poor Performance in the downstream application	Salt residue remains in eluted RNA	make sure that the elution has been completely absorbed by the column Pre-heat the elution solution to 56°C before use.
	Ethanol residue remains in eluted RNA	Do discard the flow-through after washing with W1 Buffer and centrifuge for an additional 1min.

IMPORTANT NOTES:

- a) Buffer provided in this kit contains irritants. Wear gloves and lab coat when handling these buffers.
- b) Add required volume of ethanol (96-100%) to wash Buffer before use.
- c) Centrifugation steps are done by a micro centrifuge capable of the speed at 15,000rpm.
- d) Check STB buffer for salt precipitation before use. Re-dissolve the precipitated salt by warming it in 37°C
- e) Fresh TBE electrophoresis buffer is recommended on electrophoresis for experiments requiring high purity.