

Genomic DNA Isolation Kit (Plant)

Cat. No.: SN025-0100 Size: 100 Reactions
Cat. No.: SN025-0004 Size: 4 Reactions
Sample: Up to 100 mg of fresh plant tissue
 Up to 50 mg of dry plant tissue
Format: spin column
Column capacity: up to 50 µg
Operation time: within 60 minutes



Description

The Genomic DNA Isolation Kit (Plant) is designed specifically for genomic DNA isolation from plant samples. This unique buffer system ensures total DNA with high yield and good quality from samples. The spin column system is designed to purify or concentrate DNA samples which have been previously isolated using buffers. The entire procedure can be completed in one hour without phenol / chloroform extraction. The isolated DNA is suitable for PCR or other enzymatic reactions.

Features

- Delivering high-quality genomic DNA with the fast procedure.
- Ready-to-use genomic DNA for high performance in any downstream application.
- Highly purified and high yield genomic DNA can be extracted from various samples.
- Optimized lysis buffer for the efficient lysis.
- Designed to rapidly purify high-quality DNA using spin column format.

Applications

- Gene cloning.
- Southern blotting.
- PCR.
- SNP genotyping.

Kit Contents

Contents	SN025-0100	SN025-0004
Buffer PL	55 ml X 1 bottle	2 ml X 1 vial
Buffer W1	45 ml X 1 bottle	2 ml X 1 vial
Buffer W2 (Add ethanol)	15 ml X 1 bottle (60 ml X 1 bottle)	300 µl X2 vials (1.2 ml X 2 vials)
Buffer BE	10 ml X 1 bottle	1 ml X 1 vial
PC Column	50 pieces X 2 bags	4 pieces X 1 bag
Collection tube	50 pieces X 2 bags	

Quality Control

The quality of the Genomic DNA Isolation Kit (Plant) is tested on a lot-to-lot basis to ensure consistent product quality.

Required Materials

- Mortar and pestle
- Isopropanol
- Microcentrifuge tubes
- RNase A (50 mg/ml)
- Ethanol (96-100%)
- Water bath / Dry bath

Buffer Preparation

- TE buffer, pH8.0 (Selective): 10 mM Tris-HCl, pH 8.0 with 1 mM EDTA

Genomic DNA Isolation Kit (Plant) Protocol

Step 1 Sample Preparation

1. Cut off the fresh plant tissue (up to 50 mg) or the dry plant tissue (up to 25 mg).
2. Grind the sample under liquid nitrogen to a fine powder using a mortar and pestle.

Step 2 Lysis

1. Add 500 µl of the Buffer PL and 0.5 µl of the RNase A (50 mg/ml) to the sample in the mortar and grind the sample until it is completely dissolved.
2. Transfer the dissolved sample to a 1.5 ml microcentrifuge tube.
3. Incubate at 75°C for 30 minutes. (Invert the tube every 10 minutes)
4. Centrifuge at 14-16,000 x g for 5 minutes.
5. Transfer the supernatant to a new 1.5 ml microcentrifuge tube.
#Pre-heat the Buffer BE to 75°C for Step 5 DNA Elution.

Step 3 DNA Binding

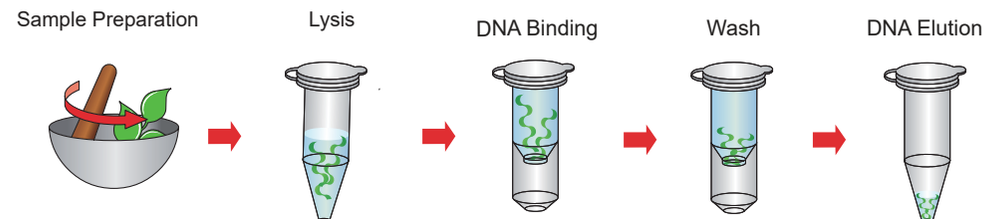
1. Add the same volume of the isopropanol to the clear supernatant from the previous step and vortex immediately for 5 seconds (e.g. add 350 µl Isopropanol to the 350 µl supernatant)
2. Place a Column PC in a 2 ml Collection Tube.
3. Transfer the mixture to the Column PC.
4. Centrifuge at 14,000 x g for 30 seconds.
5. Discard the flow-through and place the Column PC back in the 2 ml Collection Tube.

Step 4 Wash

1. Add 400 µl of the Buffer W1 into the Column PC.
2. Centrifuge at 14,000 x g for 30 seconds.
3. Discard the flow-through and place the Column PC back into the same Collection tube.
4. Add 600 µl of the Buffer W2 (Ethanol added) into the Column PC.
5. Centrifuge at 14,000 x g for 30 seconds.
6. Discard the flow-through and place the Column PC back into the same Collection tube.
7. Centrifuge at 14,000 x g again for 2 minutes to remove the residual Buffer W2.

Step 5 DNA Elution

1. Transfer the dried Column PC to a new 1.5 ml microcentrifuge tube.
2. Add 50-200 µl of the Pre-Heated Buffer BE or TE buffer (not provided) into the center of the column matrix.
3. Let stand at 75°C for 3 minutes.
4. Centrifuge for 2 minutes at 14,000 x g to elute the purified DNA.



Troubleshooting

Refer to the table below to troubleshoot problems that you may encounter when you did RNA isolation with the kit.

Problem	Cause	Solution
Low yield of DNA	Incomplete lysed sample	Use the appropriate method for the lysate preparation based on the amount of the starting materials.
		Increase the digestion time
		Make sure that the tissue is completely immersed in the Buffer PL.
	Ethanol not added to Buffer W2	Check the ethanol (96-100%) has been added to Buffer W2, and shake before use.
Incorrect elution conditions	Perform incubation at 75°C with Buffer BE before centrifugation.	
	Poor quality of starting material	Be sure to use fresh sample and process immediately after collection or freeze the sample at -80°C or in liquid nitrogen. The yield and quality of DNA isolated depends on the type and age of the starting material.
DNA degrade	Sample not fresh	Avoid repeated freeze / thaw cycles of the sample
		Use a new sample for the DNA isolation. Perform the extraction of the fresh material when possible.
	DNase contaminant	Maintain a sterile work environment to avoid contamination from DNases.
Presence of RNA	RNA contamination	Perform RNase A digestion step during Step Lysis
Inhibition of downstream enzymatic reactions	Presence of ethanol in purified DNA	Discard the ethanol of Buffer W2 flow through from the collection tube. Place the spin cartridge into the collection tube and centrifuge the spin cartridge at maximum speed for 2-3 minutes to completely dry the cartridge.

Related Ordering Information

Cat. No.	Description	Size
SM101-0500	<i>Taq</i> DNA polymerase	500 U
SM201-0100	Hot Star SUPERMIX	100 Reactions
SM255-0100	Ultrapure Proteinase K	100 mg
SA001-0500	AGAROSE Tablet, 0.5g	100 Tablets
SL001-1000	Novel Juice Supplied in 6X Loading Buffer	1 ml
SD010-R600	1 Kb DNA Ladder RTU	600 µl
SD013-R600	XLarge DNA Ladder RTU	600 µl
ST040-4000	100 mM dNTP Set	4 x 1 ml

Caution

- During operation, always wear a lab coat, disposable gloves, and protective equipment.
- Check buffers before use for salt precipitation. Redissolve any precipitate by warming to 37°C.
- When using 100 reaction assays, add 60 ml of the ethanol (96-100%) to the Buffer W2, and shake before use.
- When using 4 reaction assays, add 1.2 ml of the ethanol (96-100%) to the each vial of the Buffer W2, and shake before use.
- All products are for research use only.