

# FavorPrep™ Plant Genomic DNA Extraction HE Mini Kit

## Kit Contents

Cat. No.	FAPG1030 (4 Preps)	FAPG1033 (50 Preps)	FAPG1034 (100 Preps)
PGB1 Buffer	5 ml	50 ml	100 ml
PGB2 Buffer	2 ml	25 ml	50 ml
Wash Buffer (Concentrate)▲	3 ml	17.5 ml	35 ml
Elution Buffer	0.5 ml	5 ml	7 ml
RNase A Solution	100 µl	900 µl	900 µl × 2
Micropestles	4 pcs	50 pcs	50 pcs x 2
Filter Columns	4 pcs	50 pcs	100 pcs
HE Columns	4 pcs	50 pcs	50 pcs x 2
HE Collection Tubes	8 pcs x 2	100 pcs x 2	100 pcs x 4
Elution Tubes	4 pcs	50 pcs	100 pcs
User Manual	1	1	1
Preparation of <b>Wash Buffer</b> by adding 96~100% ethanol.			
Volume of Ethanol for Wash Buffer 🔺	12 ml	70 ml	140 ml

1. All kit components are shipped at room temperature and should be stored at room temperature between 15~25°C upon receipt, **except RNase A Solution**.

2. Store RNase A Solution at -20°C upon receipt.

#### Specification

Format/Principle	Spin Column (silica matrix)
Binding Capacity	≤125 µg DNA/Column
Operation Time	<50 mins
Sample Size	≤125 mg
DNA yield	≤40 µg
Elution Volume	30 µl







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### Preparation Before Starting

- 1. Buffers provided in this system contain irritants. Wear gloves and lab coat when handling these buffers.
- 2. Additional materials: 96~100% ethanol,  $\beta$ -Mercaptoethanol ( $\beta$ -ME).
- 3. (Optional) For long-term DNA storage, immerse the plant tissue in FavorPrep<sup>™</sup> NApreserve Reagent (Cat. No. FNPR1084) as instructed in the user manual.
- 4. Set up a water bath or dry bath at 65°C and preheat the Elution Buffer to 65°C for elution step.
- 5. Check **PGB1 Buffer** before use. If precipitates are observed, vortex PGB1 Buffer until precipitates are completely dissolved.
- 6. Fresh preparation of **PGB1-RA mixture**, premix 800 μl of **PGB1 Buffer**, 16 μl of **RNase A Solution** and 2 μl β-ME per sample before executing DNA extraction.
- 7. Add indicated volume of ethanol (96~100%) into **Wash Buffer**, mix well and store at room temperature.
- 8. Caution:  $\beta$ -ME is hazardous to human health. Always perform procedures involving  $\beta$ -ME in a fume hood.

## General Protocol

- Note: All centrifugation steps should be performed at 18,000 xg at room temperature.
- Note: Avoid disturbing the pellet or debris while transferring the supernatant.
- 1. Grind 50 mg of wet weight (up to 125 mg) plant tissue or 20 mg (up to 50 mg) dry weight of plant tissue under liquid nitrogen to a fine powder and transfer to a new microcentrifuge tube (not provided).
  - Do not allow the sample to thaw and continue immediately to step 2.
- 2. Add 800 µl **PGB1-RA mixture** to the tube and vortex thoroughly.
- 3. Incubate mixture at 65°C for 20 mins and vortex occasionally during incubation.
  - If the mixture gradually turns brown, stop the incubation process and proceed to step 4.
- 4. Place a **Filter Column** to an HE Collection Tube and transfer the entire mixture to the Filter Column.
- 5. Centrifuge for 3 mins, then transfer all the supernatant of filtrate carefully into a new **HE Collection Tube**.
- 6. Add 400  $\mu l$  PGB2 Buffer and mix thoroughly by pipetting.
- 7. Add 700  $\mu l$  ethanol (96~100%) and mix thoroughly by pipetting.



- 8. Place an **HE Column** in a new HE Collection Tube.
- 9. Transfer 900  $\mu I$  mixture carefully into the HE Column and centrifuge for 1 min. Discard flow-through.
- 10. Repeat step 9 for the remaining mixture and place the HE Column in a new HE Collection Tube.
- 11. Add 900 µl **Wash Buffer** (ethanol contained) to the HE Column. Centrifuge for 1 min then discard flow-through.
- 12. Add 500 µl Wash Buffer (ethanol contained) to the HE Column. Centrifuge for2 mins to dry the membrane directly. Discard flow-through and HE CollectionTube.
- 13. Place the HE Column in an **Elution Tube**, then add 30  $\mu$ l prewarmed **Elution Buffer** or ddH<sub>2</sub>O (pH 7.5~9.0) directly onto the membrane. Stand the HE Column for 5 mins.
  - **Important step!** For effective elution, ensure that the elution solution is dispensed onto the membrane center and absorbed completely.
- 14. Centrifuge for 1 min to elute DNA.

For more product information, please visit https://www.favorgen.com/ For technical assistance, please email us at Technical@favorgen.com

