User Manual

FAVORGEN

FavorPrep[™] 96-Well Plasmid Kit

- For 96-well high-throughput extraction of plasmid from 1~5 ml overnight culture

Kit contents:

Cat. No.: (Qanity)	FAPWE 96001 (1 plate)	FAPWE 96002 (2 plates)	FAPWE 96004 (4 plates)
FAPD1 Buffer	30 ml	65 ml	130 ml
FAPD2 Buffer	30 ml	65 ml	130 ml
FAPD3 Buffer	40 ml	85 ml	175 ml
Wash Buffer (Concentrate)	15 ml 🗖	35 ml 🔺	35 ml≜x 2
Elution Buffer	15 ml	30 ml	65 ml
RNase A Solution	75 µl	170 µl	170 µl x 2
Filter Plate (96-Well Plasmid Binding pla	1 plate ate)	2 plates	4 plates
Collection Plate (96-Well 2 ml Plate)	3 plates	6 plates	12 plates
Elution Plate (96-Well PCR plate)	1 plate	2 plates	4 plates
Adhesive Film	4 pcs	8 pcs	16 pcs

■, ▲: Add ethanol (96~100%) to Wash Buffer at the first use.

Storage

- 1. Kit components except RNase A Solution should be stored at room temperature (15~25°C).
- 2. Store RNase A Solution at -20°C upon receipt of kit.
- 3. After adding RNase A, FAPD1 Buffer should be stored at 4~8°C.

Quality Control

The quality of 96-Well Plasmid Kit is tested on a lot-to-lot basis. The plasmid is checked by restriction enzyme digestion and optical density ratio 260/280.

Specification

Principle: Filter Plate (96-well plate, glass fiber membrane) Sample size: 1~5 ml culture/preparation Processing: vacuum or centrifugation Operation time: within 60 min/96 preparations Plasmid Binding capacity: up to 60 µg/well Elution volume: 50~75 µl

Product Description

FavorPrep[™] 96-well Plasmid Kit is designed for 96 wells high-throughput isolation of plasmid. The technology is based on alkaline lysis followed by adsorption of DNA onto silica membrane in the presence of high salt. Plasmid DNA purified with this product is immediately ready for use. High-quality plasmid DNA is eluted in a small volume of Elution Buffer or deionized water. Plasmid prepared by FavorPrep[™] 96-well Plasmid Kit is suitable for a variety of routine applications including restriction enzyme digestion, sequencing, library screening, ligation and transformation, *in vitro* translation, and transfection of robust cells.

Important Note

1. Buffers provided in this kit contain irritants. Wear gloves and lab coat when handling these buffers.

Cat. No.: FAPWE 96001, 1 plate

FAPWE 96002, 2 plates

FAPWE 96004, 4 plates

For Research Use Only

- Check FAPD2 Buffer before use, warm the FAPD2 Buffer at 60°C for 5 mins if any precipitate formd.
- 3. Add RNase A to FAPD1 Buffer when first use. (see Preparation of working buffers)
- 4. Add ethanol (96~100%) to Wash Buffer when first use. (see Preparation of working buffers)

Additional Materials Required

For All Protocol:

- Pipets and pipet tips, sterile.
 96~100% ethanol (for preparation of Wash Buffer).

For vacuum processing:

- A centrifugator is required for the clarification of lysate and for the alternative of elution step, capable of 5,600~6,000 xg,
- with a swing -bucket rotor and the adaptor for 96-well plates.
 A vacuun manifold for 96-well plate and a vaccum source reached to -12 inches Hg are required.

For centrifuge processing:

• A centrifugator is required, capable of 5,600~6,000 xg, with a swing-bucket rotor and the adaptor for 96-well plates.

Preparation of Working Buffers

1. Working FAPD1 Buffer

Add indicated volume of RNase A Solution into FAPD1 buffer, mix well and store the FAPD1 buffer at 4°C.

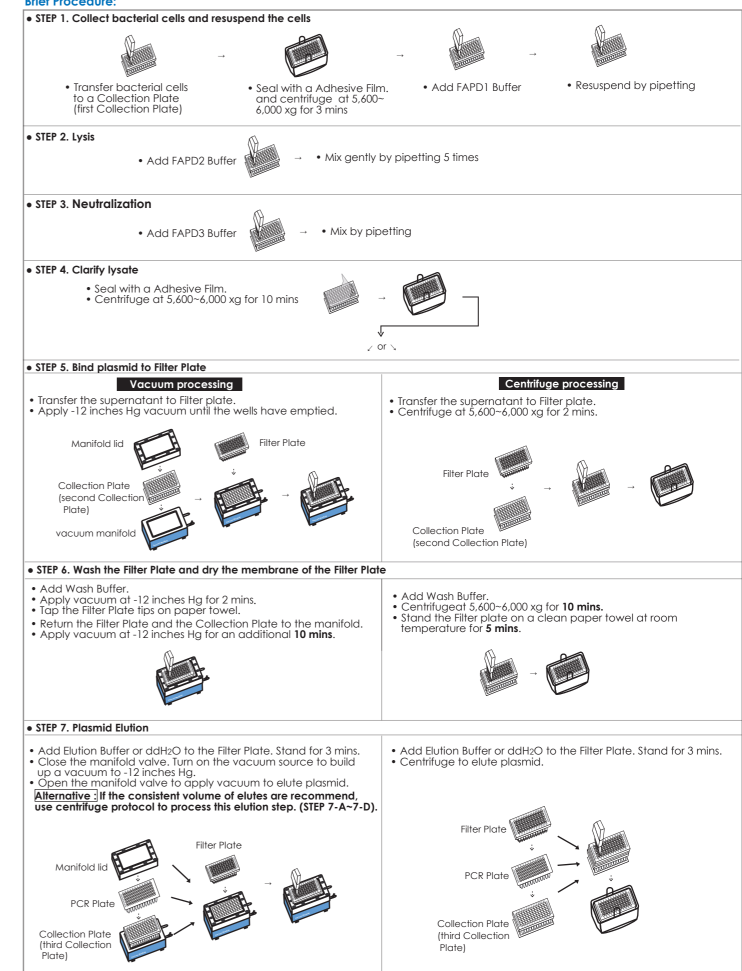
Cat. No.	FAPWE 96001	FAPWE 96002	FAPWE 96004
Volume of RNase A Solution for FAPD1 Buffer	60 µl	130 µl	260 µl

2. Working Wash Buffer

Add 96~100% ethanol to Wash Buffer when first use. Store the buffers at room temperature (15~25°C).

Cat. No.	FAPWE 96001	FAPWE 96002, FAPWE 96004
Ethanol for Wash Buffer	■ 60 ml	▲140 ml





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Safety Information:

- 1. FAPD2 Buffer and FAPD3 Buffer provided in this system contain irritants. Wear gloves and lab coat when handling these buffers.
- 2. CAUTION: FAPDE3 Buffers contain guanidinium salts which can form highly reactive compounds when combined with bleach. DO NOT add bleach or acidic solutions directly to the preparation waste.

Kit Component: FAPD2 Buffer		
Hazard contents Sodium hydroxide CAS-No. 1310-73-2 EC-No. 215-185-5	2	
Hazard statement(s) H290 H314	May be corrosive to metals. Causes severe skin burns and eye damage.	
Precautionary stateme P260	0	
P280	Wear protective gloves/ protective clothing/ eye protection/ face protection.	
P303 + P361 + P353	IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water/shower.	
P304 + P340 + P310	IF INHALED: Remove person to fresh air and keep comfortable for breathing. Immediately call a POISON CENTER/ doctor.	
P305 + P351 + P338	IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.	

Kit Component: FAPD3 Buffer	
Hazard contents Guanidine hydrochk CAS-No. 50-01-1 EC-No. 200-002-3	oride
Hazard statement(s) H302 + H332 H315 H319	Harmful if swallowed or if inhaled. Causes skin irritation. Causes serious eye irritation.
Precautionary stateme P261 P301 + P312 + P330 P305 + P351 + P338	nt(s) Avoid breathing dust/ fume/ gas/ mist/ vapours/ spray. IF SWALLOWED: Call a POISON CENTER/ doctor if you feel unwell. Rinse mouth. IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.

Kit Component: RNase A Solution		
Hazard contents RNase A CAS-No. 65742-22-5 EC-No. 232-646-6		
Hazard statement(s) H334	May cause allergy or asthma symptoms or breathing difficulties if inhaled.	
Precautionary statemen P261 P285	t(s) Avoid breathing {dust/fume/gas/mist/ vapors/spray}. In case of inadequate ventilation wear respiratory protection	
Response Statement(s)		
P304+340 P342+311	IF INHALED: Remove victim to fresh air and keep at rest in a position comfortable for breathing. If experiencing respiratory symptoms call a POISON CENTER or doctor/ physician	
	Hazard contents RNase A CAS-No. 65742-22-5 EC-No. 232-646-6 Hazard statement(s) H334 Precautionary statement P261 P285 Response Statement(s) P304+340	

Protocol: (vacuum processing)

Please read Important Notes and Additional Materials Required before starting the following steps.

STEP 1. Collect bacteria cells and resuspend the cells

- 1-1. Transfer up to 2 ml bacterial culture to each well of a Collection Plate. (provided, 96-well 2 ml plate; first Collection Plate).
- 1-2. Seal with a Adhesive Film on the Collection Plate. Place the plate in a rotor bucket and centrifuge at 5,600~6,000 xg for 3 mins. Repeat step 1-1 & 1-2 if more than 2 ml culture should be collected.

-Note: Do not exceed culture 5 ml culture.

1-3. Add 250 µl of FAPD1 Buffer (RNase A added) and resuspend the cells by pipetting.

-Note: Make sure that the cells be thoroughly resuspensed.

STEP 2. Lysis

- 2-1. Add 250 µl of FAPD2 Buffer. Mix immediately by gently pipetting the sample mixture 5 times.
- 2-2. Stand for 3~5 mins at room temperature until lysate clear.

STEP 3 Neutralization

3-1. Add 350 µl of FAPD3 Buffer. Mix immediately by pipetting. -Note: make sure that buffers have been mixed completely.

STEP 4. Clarify lysate

4-1. Seal with a new Adhesive Film. Place the plate in a rotor bucket and centrifuge at 5,600~6,000 xg for 10 mins.

STEP 5. Bind plasmid to Filter Plate

- 5-1. Fix a clean Collection Plate (provided, second Collection Plate) on the rack of vacuum manifold and cover the manifold lid. Place a Filter Plate (provided, 96-Well Plasmid binding plate) on top of the second Collection Plate.
- 5-2. Transfer the sample mixture to the Filter Plate and discard the first Collection Plate.
- 5-3. Apply vacuum at -12 inches Hg until the wells have emptied.
- 5-4. Release vacuum from the manifold.
- 5-5. Discard the flow-through. Return the Filter Plate and the second Collection Plate back to the manifold.

STEP 6. Wash the Filter Plate and dry the membrane of the Filter Plate

- 6-1. Add 650 µl of Wash Buffer (ethanol added) to each well of the Filter Plate
- 6-2. Apply vacuum at -12 inches Hg for 2 mins.
- 6-3. Release vacuum from the manifold and discard the flowthrough. Return the Filter Plate and the second Collection Plate back to the manifold.
- 6-4. Gently tap the tips of the Filter Plate on a clean paper towel to remove residual liquid. Return the Filter Plate to the second Collection Plate fixed in the manifold.
- 6-5. Apply vacuum at -12 inches Hg for an addition 10 mins.
- 6-6. Release vacuum from the manifold and discard the flow-

through and the second Collection Plate.

STEP 7. Plasmid Elution

Alternative: If the consistent volume of eluates are recommended. use "centrifuge processing step 7-A~7-D", to proceed this elution.

- 7-1. Place a Elution Plate (provided, 96-Well PCR plate) on top of a clean Collection Plate (provided, third collection plate) and fix plates on the rack of manifold. Cover the manifold lid and place the Filter Plate on the Elution Plate. (top: Filter Plate, middle: 96-well PCR Plate, bottom: third Collection Plate)
- 7-2. Add 50~75 µl of Elution Buffer or ddH2O to the membrane center of the Filter Plate. Stand for 3 mins.
 - -Note! The eluates averaged about 25 µl less than the adding volume of elution buffers. For example, adding 50 µl of Elution Buffer will recover ~25 µl of eluate. -Note! Do not use Elution Buffer or ddH2O less than the
 - suggested volume (<50 µl). It will lower the plasmid vield.
 - -Note! For effective elution, make sure that Elution Buffer or ddH2O is dispensed on the membrane center and is absorbed completely.

- 7-3. Close the manifold valve. Turn on the vacuum source to build up a vacuum to -12 inches Ha.
- 7-4. Open the manifold valve to apply vacuum to elute plasmid.
- 7-5. Release vacuum from the manifold.
- 7-6. Take out the Elution Plate (96-well PCR plate) and seal with a Adhesive Film (provided). Store the plasmid at -20°C before use

Protocol: (centrifuge processing)

Please read Important Notes and Additional Materials Required before starting the following steps.

STEP 1. Collect bacteria cells and resuspend the cells

- 1-1. Transfer up to 2 ml bacterial culture to each well of a Collection Plate. (provided, 96-well 2 ml plate; first Collection Plate).
- 1-2. Seal with a Adhesive Film on the Collection Plate. Place the plate in a rotor bucket and centrifuge at 5,600~6,000 xg for 3 mins. Repeat step 1-1 & 1-2 if more than 2 ml culture should be collected

-Note: Do not exceed culture 5 ml culture.

- 1-3. Add 250 µl of FAPD1 Buffer (RNase A added) and resuspend the cells by pipetting.
 - -Note: Make sure that the cells be thoroughly resuspensed.

STEP 2. Lysis

- 2-1. Add 250 µl of FAPD2 Buffer. Mix immediately by gently pipetting the sample mixture 5 times.
- 2-2. Stand for 3~5 mins at room temperature until lysate clear.

STEP 3. Neutralization

3-1. Add 350 µl of FAPD3 Buffer. Mix immediately by pipetting. -Note: make sure that buffers have been mixed completely.

STEP 4. Clarify lysate

Residual ethanol in membrane because insufficient drying step. •Ensure that the step of dry the membrane of the Filter plate has 4-1. Seal with a new Adhesive Film. Place the plate in a rotor bucket been processed. and centrifuge at 5,600~6,000 xg for 10 mins.

STEP 5. Bind plasmid to Filter Plate

- 5-1. Place a Filter Plate (provided, 96-Well Plasmid binding plate) on a clean Collection Plate (provided, second Collection Plate).
- 5-2. Transfer the sample mixture to each well of the Filter Plate and discard the first Collection Plate.
- 5-3. Place the combined plates (Filter Plate + the second Collection Plate) in a rotor bucket and centrifuae at 5.600~6.000 xa for 2 mins.
- 5-4. Discard the flow-through and return the Filter Plate to the second Collection Plate.

STEP 6. Wash the Filter Plate and dry the membrane of the Filter Plate

- 6-1. Add 650 µl of Wash Buffer (ethanol added) to each well of the Filter Plate.
- 6-2. Place the combined plate in a rotor bucket and centrifuge at 5,600~6,000 xg for 10 mins.
- at room temperature for 5 mins.

- 7-A. Place a Elution Plate (provided, 96-Well PCR plate) on top of a clean Collection Plate (provided, third Collection Plate) then place the Filter Plate on the Elution plate. (top: Filter Plate, middle: 96-well PCR Plate, bottom: third Collection Plate)
- 7-B. Add 50~75 µl of Elution or ddH2O to the membrane center of the Filter Plate. Stand for 3 mins.
 - -Note! The eluates averaged about 25 µl less than the adding volume of elution buffers. For example, adding 50 µl of Elution Buffer will recover ~25 µl of eluate.
 - -Note! Do not use Elution Buffer or ddH2O less than the suggested volume (<50 µl). It will lower the plasmid vield.
 - -Note! For effective elution, make sure that Elution Buffer or ddH2O is dispensed on the membrane center and is absorbed completely.

- 6-3. Place the Filter Plate on top of a clean paper towel and stand
- **STEP 7. Plasmid Elution**

- 7-C. Place the combined plates in a rotor bucket and centrifuge at 5,600~6,000 x g for 5 mins to elute plasmid.
- 7-D. Take out the Elution Plate (96-well PCR plate) and seal with a Adhesive Film (provided). Store the plasmid at -20 °C before use

Problem Shooting:

low vield

- Bacterial cells were not lysed completely
 - Too many bacterial cells were used.
 - After FAPD3 Buffer addition, break up the precipitate by pipetting to ensure higher yield.
- Overgrown of bacterial cells
 - •Incubation time should not longer than 16 hrs.
- Bacterial cells were insufficient
 - •Ensure that bacterial cells have grown to an expected amount (OD600>1) after incubation under suitable shaking modes.
- Incorrect DNA elution step
 - Ensure that Elution Buffer or ddH2O was added and absorbed to the center of the Filter plate membrane.
 - If size of DNA fragments is larger than 10 kb, use preheated Elution Buffer or ddH2O (60~70°C) on slution step to improve the elution efficiency.
- Incorrect preparation of Wash Buffer
 - •Ensure that the correct volume of ethanol (96~100%) was added to Wash Buffer prior to using.

Genomic DNA Contaminates

Lysate prepared improperly

- •For the lysis step, gently pipett the sample mixture up an down to mix well after adding the FAPD2 Buffer and do not incubat longer than 5 mins.
- Do Not use overgrown bacterial culture.

RNA Contaminates

Insufficiency of RNase A activity in FAPD1 Buffer because of long -term storage

- Prior to using FAPD1 Buffer, ensure that RNase A was added. RNase A is not properly preserved.
- •Too many bacterial cells were used. reduce sample volume.