User Manual



Kit Constants

FavorPrep™ 96-well Genomic DNA Kit

- For 96-well high-throughput extraction of DNA from whole blood, buffy coat, serum, plasma, body fluids, cultured cells and animal tissues

Cat. No.: (Q'ty)	FADWE 96001 1 plate	FADWE 96002 2 plates	FADWE 96004 4 plates
FATG1 Buffer	40 ml	80 ml	80 ml X 2
FATG2 Buffer	40 ml	80 ml	80 ml X 2
W1 Buffer * (concentrate)	44 ml	88 ml	88 ml X 2
Wash Buffer 🗖 (concentrate)	17.5 ml	35 ml	35 ml X 2
Elution Buffer	30 ml	60 ml	60 ml X 2
Proteinase K ^a	24 mg	48 mg	48 mg X 2
Filter Plate (96-Well DNA binding plate)	1 plate	2 plates	4 plates
Collection Plate (96-Well 2 ml Plate)	3 plate	6 plates	12 plates
Elution Plate (96-Well PCR plate)	1 plate	2 plates	4 plates
Adhesive Film	2 pcs	4 pcs	8 pcs

Preparation of working buffers

Add RNase-free ethanol (96~100%) to Wash Buffer 1 and Wash Buffer 2 when first use.

	FADWE 96001	FADWE 96002	FADWE 96004
* Ethanol volume for Wash Buffer 1	16 ml	32	ml
Ethanol volume for Wash Buffer 2	70 ml	140	ml

Add required ddH2O to Proteinase K bottle and disslove well. Store the prepared proteinase K at 4°C.

	FADWE 96001	FADWE 96002	FADWE 96004
^a ddH2O volume for Proteinase K	2.4 ml	4.8 ml	

Quality Control

The quality of 96-Well Genomic DNA Kit is tested on a lot-to-lot basis. The purified DNA is checked by real-time PCR and capillary electrophoresis,

Specification:

Principle: 96- well DNA Binding Plate (silica membrane) Sample size/ preparation : up to 200 µl of fresh/ frozen whole blood, buffy coat, serum, plasma, body fluids up to 25 mg of animal tissue up to 5 X 10⁶ animal cultured cells Processing: centrifugation protocol or vacuum & centrifugattion protocol Operation time: < within 90 min/ 96 preparation DNA Binding capacity: up to 30 µg/ well Elution volume: 75 ~200 µl

Reagent to be provided by user

1.96~100 % ethanol 2. (Optional) RNAse A (50 mg/ml)

Important notes:

- 1. Buffers provided in this system contain irritants. Wear gloves and lab coat when handling these buffers.
- 2. The maximum sample size is described on specification, do not use the sample more than the limitation.
- 3. Add ethanol (96~100 %) to W1 Buffer and Wash Buffer when first open.
- 4. Add ddH2O to proteinase K to prepare the 10 mg/ml proteinase K solution and store the solution at 4 °C.
- 5. Set incubator or ovens to 60°C and 70°C before the operation.
- 6. Preheat the Elution Buffer to 70°C for DNA elution.
- 7. Equipments required:
- 60°C and 70°C shaker incubators or ovens
- For centrifugation protocol: A centrifuge is required, capable of 5,600 ~ 6,000 X g, with a swing -bucket rotor and the adaptor
- For vacuum protocol: A vacuum manifold for 96-well plate and a vacuum source reached to 15 inches Hg are required.
 - (Alternative): If using centrifugation for Elution Step (STEP 6), a centrifuge equipment is required, capable of 5,600 ~ 6,000 X g, with a swing -bucket rotor and the adaptor for 96-well plate.

Sample amount and yield

Sample (maximum sample size)	Recommended amount of sample used	Average yield (µg)
Whole Blood (up to 200 µl)	200 µl	4~12
Low yield Tissue (Mouse)(up to 25 mg)	Heart, 25 mg Brain, 25 mg Kidney, 25 mg Lung, 25 mg Intestine, 10 mg	5~15 5~25 20~30 5~10 5~10
High yield Tissue (Mouse)(up to 10 mg)	Spleen, 10 mg	5~30
Animal cells (up to 5×10^6 cells)	5 x 10 ⁶ cells	15~20

Brief procedure:

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Protocol: Centrifugation processing

Please Read Important Notes Before Starting The Following Steps.

Required hardware

- 60°C and 70°C shaker incubators or ovens
- Centrifuge equipment capable of 5,600 ~ 6,000 X g with a swing -bucket rotor and the adaptor for 96-well plate

Hint

Preheat required Elution Buffer or ddH₂O (75 \sim 100 µl per well) to 70°C for DNA elution step (STEP 6)

STEP 1: Sample lysis

•Whole blood, buffy coat, serum, plasma, body fluids

- · Add 20 µl Proteinase K (10 mg/ml) to each well of a Collection Plate (provided, 96-well 2 ml plate; first collection plate).
- Add 200 µl of sample to each well and mix by pipetting.
- Add 200 µl FATG2 Buffer to each well and mix by pipetting.
- Seal with Adhesive Film. Incubate the plate with shaking at 100 rpm, 60°C for 20 min.
- (Optional) If RNA-free genomic DNA is required, add 5 µl of RNase A (50 mg/ml, not provided) to each well and incubate at room temperature for 4 min.
- Proceed to STEP 2.

Animal Tissue

Cut up to 25 mg of animal tissues (or 0.5 cm of mouse tail) and transfer into each well of a Collection Plate (provided, 96-well 2 ml plate; first collection plate). Reduce the sample size to up to 10 mg for high DNA content tissue such as liver and spleen.

- Add 200 µl FATG1 Buffer and 20 µl Proteinase K (10 mg/ml) to each well of a Collection Plate (provided, 96-well 2 ml plate; first collection plate).
- Seal with Adhesive Film. Incubate the plate with shaking at 100 rpm, 60°C for 1~2 hours or more time until the tissue sample is lysed completely. To reduce the incubation time, grind the sample in liquid nitrogen or homogenize the sample mechanically in advance.
- (Optional) If RNA-free genomic DNA is required, add 5 µl of RNase A (50 mg/ml, not provided) to each well and incubate at room temperature for 4 min.
- Add 200 µl FATG2 Buffer to each well and mix by pipetting.
- Seal with Adhesive Film. Incubate the plate with shaking at 100 rpm, 70°C for 20 min until the sample lysate is clear.

• Proceed to STEP 2.

- •Animal Cultured Cell
- Transfer cultured cells to each well of a Collection Plate (provided, 96-well 2 ml plate; first collection plate).
- \cdot Centrifuge at 1,000 x a for 10 min to pellet the cells, discard the supernatant.
- Add 200 µl FATG1 Buffer and 20 µl Proteinase K (10 mg/ml) to each well and resuspend the pellet by pipetting.
- Seal with adhesive film and incubate the plate with shaking at 100 rpm, 60°C for 10 ~ 20 min to lyse the sample.
- · (Optional) If RNA-free genomic DNA is required, add 5 µl of RNase A (50 mg/ml, not provided) to each well and incubate at room temperature for 4 min.
- Add 200 µl FATG2 Buffer to each well and mix by pipetting.
- Seal with Adhesive Film. Incubate the plate with shaking at 100 rpm, 70°C for 20 min until the sample lysate is clear.
- Proceed to STEP 2.

STEP 2. Adjust binding condition:

• Add 200 μ l ethanol (96~100%) to each well. Mix immediately by pipetting 5 ~ 10 times.

STEP 3. DNA Binding

- Place a Filter Plate (provided, 96-Well DNA binding plate) on a clean Collection Plate (provided, second collection plate).
- Transfer the sample mixture to each well of the Filter Plate and discard the Collection Plate (first collection plate).
- Place the plates in a rotor bucket and centrifuge at $5,600 \approx 6,000 \times g$ for 2 min.
- Discard the flow-through and return the Filter Plate to the Collection Plate.

STEP 4. Wash the Filter Plate with W1 Buffer and Wash Buffer

- · Add 400 µl of W1 Buffer (ethanol added) to each well of the Filter Plate.
- Place the combined plate in a rotor bucket and centrifuge at $5,600 \sim 6,000 \times q$ for 2 min.
- Discard the flow-through and return the Filter Plate to the Collection Plate.
- Add 650 µl of Wash Buffer (ethanol added) to each well of the Filter Plate.
- Place the combined plate in a rotor bucket and centrifuge at $5,600 \sim 6,000 \times g$ for 15 min.
- Discard the flow-through and return the Filter Plate to the Collection Plate.

STEP 5. Dry the membranes of Filter Plate

Place the Filter Plate on top of a clean paper towel (not provided) and stand at room temperature for 10 min.

STEP 6. DNA Elution

- Place an 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: Collection Plate)
- Add 75 ~ 200 µl of preheated Elution Buffer or ddH2O to the membrane center of the Filter Plate. Stand for 3 min.
- -- Important Step! For effective elution, make sure that Elution Buffer or ddH2O is dispensed on the membrane center and is absorbed completely.
- -- Important : Do not elute the DNA using Elution Buffer or ddH2O less than suggested volume (< 75 µl). It will lower the DNA yield.
- Place the plates in a rotor bucket and centrifuge at 5,600 ~ 6,000 x g for 5 min to elute DNA.
- Seal the Adhesive Film and store the DNA at -20°C.

Protocol: Vacuum processing

Please Read Important Notes Before Starting The Following Steps.

Required hardware

Hint

• 60°C and 70°C shaker incubators or ovens • Vacuun manifold for 96-well plate and vaccum source reached to -15 inches Hg Alternative: If using centrifugation for Elution Step (STEP 6), a centrifuge equipment is required, capable of 5,600 ~ 6,000 X g, with a swing -bucket rotor and the adaptor for 96-well plate. Preheat required Elution Buffer or ddH₂O (75 \sim 100 µl per well) to 70°C for DNA elution step (STEP 6) STEP 1: Sample lysis •Whole blood, buffy coat, serum, plasma, body fluids • Add 20 µl Proteinase K (10 mg/ml) to each well of a Collection Plate (provided, 96-well 2 ml plate; first collection plate). • Add 200 µl of sample to each well and mix by pipetting. · Add 200 µl FATG2 Buffer to each well and mix by pipetting. • Seal with Adhesive Film. Incubate the plate with shaking at 100 rpm, 60°C for 20 min. • (Optional) If RNA-free genomic DNA is required, add 5 µl of RNase A (50 mg/ml, not provided) to each well and incubate at room temperature for 4 min. • Proceed to STEP 2. • Animal Tissue Cut up to 25 mg of animal tissues (or 0.5 cm of mouse tail) and transfer into each well of a Collection Plate (provided, 96-well 2 ml plate; first collection plate). Reduce the sample size to up to 10 mg for high DNA content tissue such as liver and spleen. · Add 200 µl FATG1 Buffer and 20 µl Proteinase K (10 mg/ml) to each well of a Collection Plate (provided, 96-well 2 ml plate; first collection plate). Seal with Adhesive Film. Incubate the plate with shaking at 100 rpm, 60°C for 1~2 hours or more time until the tissue sample is lysed completely. To reduce the incubation time, grind the sample in liquid nitrogen or homogenize the sample mechanically in advance. (Optional) If RNA-free genomic DNA is required, add 5 µl of RNase A (50 mg/ml, not provided) to each well and incubate at room temperature for 4 min. · Add 200 µl FATG2 Buffer to each well and mix by pipetting. Seal with Adhesive Film. Incubate the plate with shaking at 100 rpm, 70°C for 20 min until the sample lysate is clear. • Proceed to STEP 2. Animal Cultured Cell Transfer cultured cells to each well of a Collection Plate (provided, 96-well 2 ml plate; first collection plate). \cdot Centrifuge at 1,000 x g for 10 min to pellet the cells, discard the supernatant. • Add 200 µl FATG1 Buffer and 20 µl Proteinase K (10 mg/ml) to each well and resuspend the pellet by pipetting. Seal with adhesive film and incubate the plate with shaking at 100 rpm, 60°C for 10 ~ 20 min to lyse the sample. · (Optional) If RNA-free genomic DNA is required, add 5 µl of RNase A (50 mg/ml, not provided) to each well and incubate at room temperature for 4 min. • Add 200 µl FATG2 Buffer to each well and mix by pipetting. Seal with Adhesive Film. Incubate the plate with shaking at 100 rpm, 70°C for 20 min until the sample lysate is clear. • Proceed to STEP 2. STEP 2: Adjust binding condition: • Add 200 µl ethanol (96~100%) to each well. Mix immediately by pipetting 5 ~ 10 times. STEP 3. DNA Binding · 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 DNA binding plate) on top of the Collection Plate. • Transfer the sample mixture to the Filter Plate and discard the Collection Plate (first collection plate). · Apply vacuum at 10 inches Hg until the wells have emptied. · Discard the flow-through and return the Filter Plate and the Collection Plate to the manifold. STEP 4. Wash the Filter Plate with W1 Buffer and Wash Buffer • Add 400 µl of Wash Buffer 1 (ethanol added) to each well of the Filter Plate. · Apply vacuum at 10 inches Hg until the wells have emptied. • Discard the flow-through and return the Filter Plate and the Collection Plate to the manifold. • Add 650 µl of Wash Buffer 2 (ethanol added) to each well of the Filter Plate. · Apply vacuum at 10 inches Hg until the wells have emptied. Discard the flow-through and return the Filter Plate and the Collection Plate to the manifold. STEP 5. Dry the membranes of Filter Plate · Gently tap the tips of the Filter Plate on a clean paper towel to remove residual liquid. · Return the Filter Plate to the Collection Plate fixed in the manifold. Apply vacuum for an addition 10 min. • Discard the flow-through and the Collection Plate (third plate). STEP 6. DNA Elution Place an 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: Collection Plate) Add 75 ~ 200 µl of preheated Elution Buffer or ddH2O to the membrane center of the Filter Plate. Stand for 3 min. -- Important Step! For effective elution, make sure that Elution Buffer or ddH2O is dispensed on the membrane center and is absorbed completely. -- Important : Do not elute the DNA using Elution Buffer or ddH2O less than suggested volume (< 75 µl). It will lower the DNA yield. Close the manifold valve. Turn on the vacuum source to build up a vacuum to 15 inches Hg. Open the manifold valve to apply vacuum to elute DNA. Seal the Adhesive Film and store the RNA at -20°C. Alternative: If the consistent volume of elutes are recommend use centrifuge protocol to proceed this elution step. (Page 3, STEP 6)

3