#### Manual



# FAVORGEN® FavorPrep<sup>TM</sup> Fungi/ Yeast Genomic DNA Extraction Mini Kit

Cat. No.: FAFYG 000 FAFYG 001 FAFYG 001-1

(For Research Use Only)

# **Kit Contents:**

Cat. No:	FAFYG 000 (4 preps_sample)	FAFYG 001 (50 preps)	FAFYG 001-1 (100 preps)
Beads Tube FA Buffer FB Buffer TG1 Buffer TG2 Buffer W1 Buffer a (concentrate) Wash Buffer b (concentrate) Elution Buffer Lyticase solution Proteinase K TG Mini Column Collection Tube Elution Tube	4 pcs 5 ml 2.7 ml 2 ml 2 ml 1.3 ml 1 ml 0.5 ml 250 µl 1 mg 4 pcs 8 pcs 4 pcs	50 pcs 60 ml 32 ml 22 ml 15 ml 22 ml 10 ml 7 ml 550 µl x 5 11 mg 10 pcs x 5 100 pcs 50 pcs	100 pcs 120 ml 65 ml 45 ml 30 ml 44 ml 20 ml 15 ml 550 µl x 10 11 mg x 2 10 pcs x10 100 pcs x 2

Preparation of W1 Buffer and Wash Buffer by adding ethanol (96 ~ 100%) and <b>Store at RT.</b>					
Ethanol volume for W1 Buffer <sup>a</sup>	0.5 ml	8 ml	16 ml		
Ethanol volume for Wash Buffer <sup>b</sup>	4 ml	40 ml	80 ml		

Preparation of Proteinase K solution by adding ddH2O and <b>Store the solution at 4 °C.</b>				
ddH2O volume for Proteinase K <sup>c</sup>	0.1 ml	1.1 ml	1.1 ml x 2	

### **Specification:**

Principle: mini spin column (silica matrix)

Sample size:  $1 \sim 5 \times 10^6$ Operation time:  $\sim 60$  minutes Binding capacity:  $\sim 60$  µg/ column

Column applicability: centrifugation and vaccum

# Additional requirement to be provided by user

- 1. Microcentrifuge capable of speed at ~18,000 x g
- 2. 1.5 ml microcentrifuge tube
- 3.96~100 % ethanol
- 4. Vortex
- 5. Heating block or water bath

# **Important Notes:**

- 1. Buffers provided in this system contain irritants. Wear gloves and lab coat when handling these buffers.
- 2. Store the Lyticase solution at -20  $^{\circ}\text{C}$  on arrival.
- Caution: Lyticase solution and FB Buffer containing 14 mM of 
  ß-mercaptoethanol is hazardous to human health. perform the
  procedures involving Lyticase solution and FB Buffer in a chemical fume hood.
- 4. Add sterilized ddH2O to a Proteinase K tube to make a 10 mg/ml stock solution. Vortex and make sure that Proteinase K has been completely dissolved. Store the stock solution at 4 °C.
- 5. Add required volume of ethanol (96-100%) to W1 Buffer and Wash Buffer when first open. Store the solution at room temperature.
- 6. Prepare a heating block or a water baths to 37 °C for step 4 and to 55 °C for step 9 before operation.



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#### **General Protocol:**

Please Read Important Notes Before Starting Following Steps.

- 1. Transfer 1~5 x10<sup>6</sup> of cultures (fungal/yeast cells) to a 1.5 ml microcentrifuge tube. (not provided)
- 2. Add 1 ml of FA Buffer to the cells and resuspend the cells by pipetting.
- 3. Descend the cells by centrifuging at 5,000 x g for 2 min and discard the supernatant completely.
- Resuspend the cells in 550 μl of FB buffer and add 50 μl of lyticase solution, mix well by vortexing. Incubat the sample at 37 °C for 30 min.
  - Caution: Lyticase solution and FB Buffer containing 14 mM of  $\beta$ -mercaptoethanol is hazardous to human health. perform the procedures involving Lyticase solution and FB Buffer in a chemical fume hood.
- 5. **(Optional)** If RNA-free genomic DNA is required, add 8 µl of 50 mg/ml **RNase A** (not provided) and incubate for 2 min at room temperature.
- 6. Descend the cells by centrifuging at 5,000 x g for 10 min. Remove the supernatant completely.
- 7. Add 350 µl TG1 Buffer and mix well by pipetting. Transfer the sample mixture to a bead tube. (provided)
- 8. Mix well by Plus-vortexing for 5 minutes.
- 9. Add 20 µl of **Proteinase K** (10 mg/ml) **and mix well by vortexing.** Incubate at 55 °C for 15 min, vortex 30 seconds for every 5 minutes incubation.
- 10. Descend the cells by centrifuging at 5,000 x g for 1 min and transfer 200 µl of supernatant to a new 1.5 ml microcentrifuge tube. (not provided)
- 11. Add 200 µl of **TG2 Buffer** and mix well by pipetting.
- 12. Add 200 µl of **ethanol** (96-100%) and mix well by pulse-vortexing for 10 seconds.
- 13. Place a **IG Mini Column** in **Collection Tube**. Transfer the sample mixture (including any precipitate) carefully to **IG Mini Column**. Centrifuge at 11,000 x g for 30 second **then place the IG Mini Column to a new Collection Tube**.
- 14. Add 400 µl of **W1 Buffe**r to the **TG Mini Column.** Centrifuge at 11,000 x g for 30 seconds and discard the flow-through. Place the TG Mini Column back to the Collection Tube.
  - --Make sure ethanol has been added into W1 Buffer when first use.
- 15. Add 750 µl of **Wash Buffe**r to the **TG Mini Column.** Centrifuge at 11,000 x g for 30 seconds and discard the flow-through. Place the TG Mini Column back to the Collection Tube.
  - --Make sure ethanol has been added into Wash Buffer when first use.
- 16. Centrifuge at full speed (~ 18,000 x g) for an additional 3 min to dry the column. Important Step! This step will avoid the residual liquid to inhibit subsequent enzymatic reactions.
- 17. Place the **TG Mini Column** to a **Elution Tube**.
- 18. Add 50 ~100 µl of Elution Buffer or ddH2O to the membrane center of the TG Mini Column. Stand TG Mini Column for 3 min. Important Step! For effective elution, make sure that the elution solution is dispensed onto the membrane center and is absorbed completely.
- 19. Centrifuge at full speed (~ 18,000 x g) for 1 min to elute total DNA.
- 20. Store total DNA at 4°C or -20°C.

