



## FavorPrep™ Stool DNA Isolation Mini Kit

Cat.No. : FASTI 000, 4 preps  
 FASTI 001, 50 Preps  
 FASTI 001-1, 100 Preps  
**(For Research Use Only)**

### Kit Contents:

|                              | FASTI 000<br>(4 preps_sample) | FASTI 001<br>(50 preps) | FASTI 001-1<br>(100 preps) |
|------------------------------|-------------------------------|-------------------------|----------------------------|
| SDE1 Buffer                  | 1.8 ml                        | 20 ml                   | 40 ml                      |
| SDE2 Buffer                  | 1.2 ml                        | 7 ml                    | 14 ml                      |
| SDE3 Buffer                  | 1.2 ml                        | 15 ml                   | 30 ml                      |
| SDE4 Buffer                  | 3 ml                          | 20 ml                   | 40 ml                      |
| Wash Buffer (concentrate) *  | 1.5 ml                        | 20 ml                   | 35 ml                      |
| Elution Buffer               | 1.5 ml                        | 15 ml                   | 30 ml                      |
| Proteinase K (lyophilized) * | 1.1 mg                        | 11 mg                   | 11 mg x 2                  |
| SDE Mini Columns             | 4 pcs                         | 50 pcs                  | 100 pcs                    |
| Collection Tubes             | 8 pcs                         | 100 pcs                 | 200 pcs                    |
| Elution Tubes                | 4 pcs                         | 50 pcs                  | 100 pcs                    |
| Beads tube                   | 4 pcs                         | 50 pcs                  | 100 pcs                    |
| User Manual                  | 1                             | 1                       | 1                          |

\* Preparation of Proteinase K solution and Wash Buffer for first use:

| Cat. No:  | FASTI 000 | FASTI 001 | FASTI 001-1 |
|---|-----------|-----------|-------------|
| ddH <sub>2</sub> O volume for Proteinase K Solution | 0.11 ml   | 1.1 ml    | 1.1 ml      |
| ethanol volume for Wash Buffer                      | 6 ml      | 80 ml     | 140 ml      |

### Specification:

Principle: spin column (silica membrane)

Sample: 50 ~100 mg

Operation time: < 60 min

Elution volume: 50~200 µl

### Important Notes:

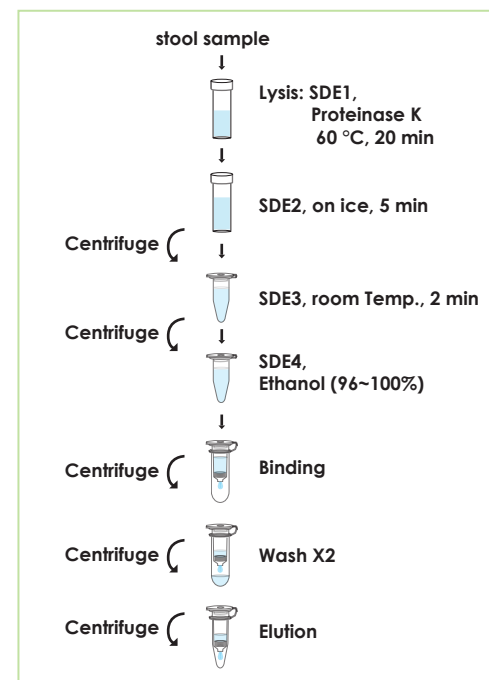
1. Buffers provided in this system contain irritants. Wear gloves and lab coat when handling these buffers.
2. Check SDE1 Buffer before use, Warm SDE1 Buffer at 60°C for 10 minutes if any precipitate formd.
3. Add required sterile ddH<sub>2</sub>O to Proteinase K tube to make a 10 mg/ml stock solution. Vortex and make sure that Proteinase K has been completely dissolved. **Store the stock solution at 4 °C.**
4. Add indicated volume of ethanol (96-100%) to Wash Buffer before use.
5. Prepare a heating block or a water bath to 60 °C. If DNA is isolated from gram positive bacteria, prepare a heating block or a water bath to 95 °C for another incubation.
6. All centrifuge steps are done at full speed (~18,000 x g) in a microcentrifuge.
7. Preheat Elution Buffer or ddH<sub>2</sub>O to 60°C for elution step.

### General Protocol:

Please Read Important Notes Before Starting Following Steps.

1. **Add up to 200 mg of stool sample to a bead tube and place the tube on ice.**  
 --If the sample is liquid, add 200 µl of sample into a bead tube.
2. **Add 300 µl of SDE1 Buffer and 20 µl of proteinase K (10 mg/ml) to the sample.**  
**Vortex at maximum speed for 5 minutes. Incubate the sample mixture at 60 °C for 20 minutes and vortex the sample for every 5 minutes during the incubation.**  
 -- Make sure stool sample is homogenized completely.  
 -- For isolation of DNA from gram positive bacteria, do a further incubation at 95 °C for 5 minutes after proteinase K lysis.
3. **Briefly spin the tube to remove drops from the inside of the lid.**
4. **Cool down the sample mixture and add 100 µl of SDE2 Buffer. Mix well by vortexing and incubate the sample mixture on ice for 5 minutes.**
5. **Centrifuge at full speed ( ~18,000 x g) for 5 minutes.**
6. **Carefully transfer the supernatant to a 1.5 ml microcentrifuge tube (not provided) and discard the stool pellet.**  
 --Avoid pipetting any debris and pellet.
7. **Add 200 µl of SDE3 Buffer. Mix well by vortexing and incubate the sample mixture at room temperature for 2 minutes.**  
 --**Note:** SDE3 Buffer must be suspended completely by vigorously vortexing before every using.  
 -- Cut off the end of 1 ml tip to make it easier for pipetting the SDE3 Buffer.
8. **Centrifuge at full speed for 2 minutes.**
9. **Carefully transfer 250 µl of supernatant to a 1.5 ml microcentrifuge tube (not provided).**  
 --Avoid pipetting any debris and pellet.

### Brief Procedure:



10. (Optional) If RNA-free DNA is required, add 1  $\mu$ l of 100 mg/ml RNase A (not provided). Mix well and incubate the sample mixture at room temperature for 2 min.
11. Briefly spin the tube to remove drops from the inside of the lid.
12. Add 250  $\mu$ l of SDE4 Buffer and 250  $\mu$ l of ethanol (96~100%). Mix thoroughly by pulse-vortexing.
13. Place a SED Column into a Collection. Transfer all of the sample mixture to the SDE Column. Centrifuge at full speed for 1 min and discard the flow-through then place the SDE Column into a new Collection Tube.
14. Add 750  $\mu$ l of Wash Buffer (ethanol added) to the SDE Column. Centrifuge at full speed for 1 min then discard the flow-through. Return the SDE Column back to the Collection Tube.  
--Make sure that ethanol (96~100%) has been added into Wash Buffer when first use.
15. Repeat step 15.
16. Centrifuge at full speed for an additional 3 min to dry the SDE column.  
--Important step! This step will avoid the residual liquid to inhibit subsequent enzymatic reactions.
17. Place the SDE Column into a 1.5 ml microcentrifuge tube (not provided). Add 50 ~ 200  $\mu$ l of preheated Elution Buffer or ddH<sub>2</sub>O to the membrane center of the SDE Column. Stand the SDE Column for 2 min at room temperature.  
--Important step! For effective elution, make sure that the Elution Buffer or ddH<sub>2</sub>O is dispensed onto the membrane center and is absorbed completely.
18. Centrifuge at full speed for 1 min to elute DNA.

## Troubleshooting

| Problem   | Possible reasons  | Solutions  |
|---|---|--|
| <b>Low or no yield of genomic DNA</b>                   |   |  |
|   | Sample stored incorrectly   | Store the stool sample at -20 °C.  |
|   | Low amount of cells in the sample   | Increase the sample size   |
| <b>Poor cell lysis</b>                                  |   |  |
|   | Poor cell lysis because of insufficient beads beating time                                  | Extend the beads beating time.   |
|   | Poor cell lysis because of insufficient mixing with SD1, SD2 Buffer and Proteinase K        | Mix the sample mixture immediately thoroughly by pulse-vortexing after adding SD1, SD2 Buffer and proteinase K solution.   |
|   | Poor cell lysis because of insufficient Proteinase K activity                               | Use a fresh or well-stored Proteinase K stock solution.  |
| <b>Insufficient binding of DNA to column's membrane</b> |   |  |
|   | Ethanol is not added into sample lysate before DNA binding                                  | Make sure that the correct volumes of ethanol (96- 100 %) is added into the sample lysate before DNA binding.  |
|   | Ethanol and sample lysate did not mix well before DNA binding                               | Make sure that Ethanol and sample lysate have been mixed completely before DNA binding   |
| <b>Incorrect preparation of Wash Buffer W1/W2</b>       |   |  |
|   | Ethanol is not added into Wash Buffer when first use  | Make sure that the correct volumes of ethanol (96- 100 %) is added into Wash Buffer when first use.  |
|   | The volume or the percentage of ethanol is not correct for adding into Wash Buffer          | Make sure that the correct volumes of ethanol (96- 100 %) is added into Wash Buffer when first use.  |
| <b>Elution of DNA is not efficient</b>                  |   |  |
|   | pH of water (ddH <sub>2</sub> O) for elution is acidic                                      | Make sure the pH of ddH <sub>2</sub> O is between 7.0-8.5.<br>Use Elution Buffer (provided) for elution .  |
|   | Elution Buffer or ddH <sub>2</sub> O is not completely absorbed by column membrane          | After Elution Buffer or ddH <sub>2</sub> O is added, stand the SD Column for 5 min before centrifugation.  |
| <b>Poor quality of genomic DNA</b>                      |   |  |
| A260/A280 ratio of eluted DNA is low                    | <b>Poor cell lysis</b>  |  |
|   | Poor cell lysis because of insufficient beads beating time                                  | Extend the beads beating time.   |
|   | Poor cell lysis because of insufficient mixing with SD1 buffer, SD2 Buffer and Proteinase K | Mix the sample mixture immediately thoroughly by pulse-vortexing after adding SD1, SD2 Buffer and proteinase K solution.   |
|   | Poor cell lysis because of insufficient Proteinase K activity                               | Use a fresh or well-stored Proteinase K stock solution.  |
| A260/A280 ratio of eluted DNA is high                   | A lot of residual RNA in eluted DNA   | Add 8 $\mu$ l of RNase A (50 mg/ml) to the eluate and incubate at 37 °C for 10 minutes. After incubation, add 200 $\mu$ l of SD2 Buffer and 200 $\mu$ l of ethanol (96~100%), mix well by plus-vortexing. Then follow the general Protocol starting from step 7. |