

ExiProgen™EC-Disulfide Protein Synthesis Kit (Cat. No. K-7330)

Step | Preparation of Template DNA

1) Prepare the 'Template DNA' for protein expression.

(Note I) The Template DNA' must contain the following: T7 Promoter, Ribosome binding site (RBS) Target gene (with His-taq), and T7 terminator.

(Note II) Use 1 µg/kb (insert size of DNA) of plasmid DNA (Purity: A260/280 > 1.7, A260/230 > 1.5).

Step | Preparation of Experiment

- 1) Take out the 'Cartridge 2)' from **Box** 2. Thaw them at room temperature.
- 2) Take out 'E coli extract,' Master Mix', 'DEPC DW' and 'Storage Buffer' from Box ② and thaw them on ice. Take out 'Cartridge ①,' 'Disposable Filter Tip', and 'Protection Cover' from Box ①. (Note) Make sure that all the solutions are completely thawed. It takes about 2 hours.
- 3) Prepare the 'Reaction Block (ExiProgen™ accessory)'.
- 4) Take out twice as many 'Dialysis Tube' as the number of samples from **Box** (1).
- 5) Remove the solution in the 'Dialysis Tube' and rinse inside and out using squeeze bottle with sterile distilled water.

(Note) The solution in the 'Dialysis Tube' is 20%(v/v) ethanol.

6) Install the 'Dialysis Tube' on the 'Reaction Block,' as shown in the diagram.



7) Remove water in the 'Dialysis Tubes' and prepare expression solution as shown in the table.

The expression solution should be loaded at the 'Dialysis Tube' in row A (Protein expression section). Fill 500 µl of sterile distilled water in the 'Dialysis Tube' on row B (Dialysis section).

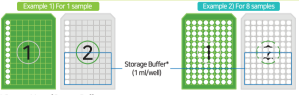
*Composition of Expression Solution

		Positive control DNA (Optional)	
Template DNA	Xμl	2 μΙ	
E.coli extract	120 µl	120 µl	
Master Mix	210 µl	210 µl	
DEPC DW	(120-X) µl	118 µl	
Total volume	450 µl	450 µl	



- 8) Place the 'Protection Cover' on the 'Reaction Block'.
- 9) Punch holes in the sealing films of 'Cartridge (1), (2).
- 10) Add 1ml of 'Storage Buffer' G1-I2 rows of 'Cartridge 2)'.

(Note) The number of columns punched should match with the sample number.



- * Composition of Storage Buffer
- $-50\,mM\,Tris-HCl, 100\,mM\,NaCl, 1\,mM\,DTT, 0.05\% (v/v)\,NaN_3, 50\% (v/v)\,glycerol, pH\,7.6.$

Step III Protein Synthesis with ExiProgen™

Turn on the ExiProgen™ and tap 'Press to Start' button.
 Wait until the [MENU] screen appears.

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- 2) Open the door of ExiProgen™ and pull out the baseplate.
- 3) Load 'Cartridge 1), 2), 'Waste Tray', and 'Reaction Block'.

Follow in order: 'Cartridge $\textcircled{2}' \rightarrow$ 'Cartridge $\textcircled{1}' \rightarrow$ 'Waste Tray' \rightarrow 'Reaction Block'.

(Note I) Make sure that the first and second lines of 'Cartridge @' are firmly fixed on the heating block.

(Note II) There are 'silicon rings' embedded on both sides of the 'Cartridge 1' installation position. Place the left side first, and then the right side.

4) Place the 'Disposable Filter Tips' on row B of 'Disposable Tip Rack'.

(Note) Tips should be placed in the same columns with the punched holes of the Cartridges.













- 5) Push the baseplate in until you hear the click sound, then close the door.
- 6) Tap the following: [MENU] 'Start' → [PREP SETUP] '905 (Synthesis_DS), 'Enter' → [PREP SETUP, Elution Volume] 'OK' → [PREP SETUP, Reaction Temperature] '26°C, 'OK' → [CHECK LIST] 'OK' → [Running Mode] 'RUN'.













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- 7) The [Work Completion] screen will appear once the protocol is completed. It takes approximately 36 hours.
- 8) Purified protein samples in the 'Storage Buffer' can be collected from the 'Dialysis Tube', located at the row B of the 'Reaction Block'.

Step IV Analysis of Sample

A. Position of each sample (in 'Cartridge 2)')



Bead sample
Expression sample
Unbound sample
and sample

IExpression sample

 $: Samples \ after \ expression, but \ without \ purification$

IUnbound sample

: Samples not bound to Ni-NTA magnetic bead

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: Samples after 1st washing step in purification process

| Bead sample |

: Used bead samples for purification

B. Sampling for SDS-PAGE analysis

1) Prepare loading mixture as shown in the table. Incubate the samples at 95°C for 5-10 min.

	Expression/Unbound/ 1st washing sample	Purified protein sample	Bead sample
Sample	5 µl	10 µl	15 µl
4X Loading dye	5 µl	5 µl	5 µl
Sterile distilled water	10 µl	5 µl	-
Total volume	20 µl	20 µl	20 µl

(Note) Add 100 µl of sterile distilled water to the well containing the beads for suspension. Proceed with the sampling afterwards.

Load each sample to wells of SDS-PAGE gel [10 x 8 (cm), 0.75 mm thick, 10-well].
 (Note) Expression, Unbound and 1st washing samples: 5 µl/well,

Purified proteins and Bead samples: 10 µl/well

※ For more information, visit our website (www.bioneer.com).