

## ExiProgen™ EC-Bulk Protein Synthesis Kit (Cat. No. K-7340)

### Step I Protein Expression

#### A. Preparation of Experiment

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

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

Use 5-20 µg of plasmid DNA or 30-50 µg of PCR product.

(Purity:  $A_{260/280} > 1.7$ ,  $A_{260/230} > 1.5$ ).

- 2) Take out the 'Expression Cartridge ①, ②' from Box ② and thaw them at room temperature.
- 3) Take out 'E. coli extract', 'Master Mix', and 'DEPC DW' from a Box ② and thaw them on ice.
- (Note) Please make sure that all solutions are completely thawed. It takes about 3-4 hours.
- 4) Punch all holes in the sealing films of 'Expression Cartridge ①, ②'.
- 5) Install 'Bulk SECF device' from a Box ① on a 'Bulk Reactor (Cat. No. KA-7340-1)' and fill the 'Bulk SECF device' with the 'Reaction Mixture' as belows.

(Note) 'Bulk SECF device' is filled with solution containing 0.05% NaNa<sub>3</sub>.

Remove the solution in the device before use.

Reaction Mixture	
Template DNA	X ml
E. coli extract	5.34 ml
Master Mix	9.34 ml
DEPC DW	(5.32·X) ml
Total volume	20 ml



#### B. Protein Expression with ExiProgen™

- 1) Turn on the ExiProgen™ and tap 'Press to Start' button.

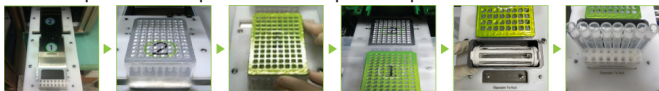
Wait until the [MENU] screen appears.

- 2) Open the door of ExiProgen™ and pull out the baseplate.

- 3) Load 'Cartridge ①, ②', 'Waste Tray', and 'Bulk Reactor'.

Follow in order: 'Cartridge ②' → 'Cartridge ①' → 'Waste Tray' → 'Bulk Reactor'

, Place 8 'Disposable Filter Tips' on row B of 'Disposable Tip Rack'.



(Note I) Make sure that the first and second lines of 'Cartridge ②' 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) Push the baseplate in until you hear the click sound, then close the door.

- 5) Press [MENU] 'Start' → [PREP SETUP], '906 (Bulk Expression)', 'Enter' → [PREP SETUP, Elution Volume] 'OK' → [PREP SETUP, Reaction Temperature] '30°C', 'OK' → [CHECK LIST] 'OK' → [Running Mode] 'RUN'.



- 6) The [Work Completion] screen will appear once the protocol is completed.

It takes approximately 25 hours.

- 7) Expressed protein samples will be collected in the 'Bulk SECF device'.

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### Step II Protein Purification

#### A. Preparation of Experiment

- 1) Take out 'Purification Cartridge ①, ②' from a **Box ①** and punch all holes in the sealing films.
- 2) Load the **1 ml/well (Total 16 ml)** expressed protein samples (Final sample of Step I) on the '**Sample Loading Well (16 wells)**' of 'Purification Cartridge ①'.
- 3) Install the new 'Bulk SECF device' from a **Box ①** on a 'Bulk Reactor (Cat. No. KA-7340-1)' and **fill 20 ml of sterile distilled water in the device.**  
(Note) 'Bulk SECF device' is filled with solution containing 0.05% NaN<sub>3</sub>.  
Remove the solution in the device before use.

#### B. Protein Purification with ExiProgen™

- 1-4) 1-4 steps are the same as 'Protein Expression' protocol. Refer to front page.
- 5) Press [MENU]'Start' → [PREP SETUP], **907 (Bulk Purification)**, 'Enter' → [PREP SETUP, Elution Volume]'OK' → [PREP SETUP, Reaction Temperature]'OK' → [CHECK LIST]'OK' → [Running Mode]'RUN'.
- 6) The [Work Completion] screen will appear once the protocol is completed.  
It takes approximately **15 hours.**  
(Note) The '907' protocol includes [Ni-NTA affinity purification] and [Dialysis to storage buffer] processes. If you want to separate two processes, use [908 (Bulk\_Ni-NTA purify)] and [909 (Bulk\_Storage)] protocol.
- 7) **Purified protein samples in storage buffer will be collected in the 'Bulk SECF device'.**  
(Note I) Purified protein samples may contain trace amounts of magnetic beads. The beads do not influence to proteins and can be easily removed through spin-down.  
(Note II) Composition of Storage Buffer: 50 mM Tris-HCl, 100 mM NaCl, 1 mM DTT, 0.1 mM EDTA, 0.05% NaN<sub>3</sub>, 50% Glycerol, pH 7.6.

### Step III Analysis of Sample

#### A. Position of each sample (in 'Purification Cartridge ①, ②)

The diagram shows two purification cartridges, labeled 1 and 2. Cartridge 1 is a 4x4 grid with wells labeled F2, F1, E2, E1, D2, D1, C2, C1, B2, B1, A2, A1. A large circle with the number '1' is in the center. Cartridge 2 is a 4x4 grid with wells labeled L2, L1, K2, K1, J2, J1, I2, I1, H2, H1, G2, G1. A large circle with the number '2' is in the center. A red box highlights wells L2 and L1 in Cartridge 2, with a label 'Bead' pointing to it. A green box highlights wells A2 and A1 in Cartridge 1, with a label '1<sup>st</sup> washing' pointing to it. A blue box highlights wells B2 and B1 in Cartridge 1, with a label 'Unbound' pointing to it.

- Unbound sample**: Samples not bound to Ni-NTA magnetic bead
- 1<sup>st</sup> washing sample**: Samples after 1<sup>st</sup> 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.

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

(Note) Add 360-400 µl/well of sterile distilled water to the well containing the beads for suspension. Proceed with the sampling afterwards.

- 2) Incubate the samples at 95°C for 5-10 min.
- 3) Load 5 µl of each sample to a well of SDS-PAGE gel [10 x 8 (cm), 0.75 mm thick, 10-well].

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