DsbC Set

#PF005-10ML-EX For 2 mL x 5 Reaction

PUREfrex® is NOT included.

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in vitro research use only Store at -80°⊂ before opening

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Kit components

| GSSG (Green) ^{*1} | 100 µL x 5 |
|---|------------------------------|
| Oxidized glutathione (60 mM) | Store at -20°C |
| DsbC (Green) ^{*2} E.coli DsbC (320 μM) | 100 µL x 5 |
| | Store at -80°C ^{*3} |
| Dilution Buffer (Clear) 30% glycerol buffer | 500 μL x 1 |
| | Store at -20°C |

Introduction

1. About PUREfrex®

PUREfrex[®] is a reconstituted cell-free protein synthesis kit which GeneFrontier has developed based on the PURE system technology. The target protein can be synthesized by adding the template DNA (or mRNA) to the reaction mixture. The PURE system is a unique cell-free protein synthesis system, which has originally developed by Professor Takuya Ueda at the University of Tokyo, and consists of only purified factors necessary for transcription, translation and energy regeneration (Ref. 1). Therefore it enables to adjust the composition of the reaction mixture.

PURE *frex*[®] has been raised in purity by improving the methods for preparing ribosomes, tRNAs and all proteins in the reaction mixture compared with the original PURE system (Ref. 2). As the result, the contaminating lipopolysaccharide from E. coli is reduced to around 0.1 EU per 1 µL of reaction and other contaminants, such as RNase and β-galactosidase, are also reduced.

Because all of proteins in PURE frex® have no tags, the synthesized protein can be purified and detected by any tags.

References) 1. Shimizu et al. (2001) Nat. Biotecnol., vol. 19, p. 751 2. Shimizu et al. (2005) Methods, vol. 36, p. 299

Introduction

2. About DsbC Set / PDI Set

Formation of a disulfide bond is one of an important process for folding and stability of most of secretory proteins such as enzymes or antibodies.

Oxidative environment is necessary to form a disulfide bond because a disulfide bond is formed by the oxidation of sulfhydryl groups (-SH) of adjacent cysteine residues. Disulfide bond isomerase, which can catalyze the exchange of disulfide bridges, may be also required for a correct pairing of cysteines.

DsbC Set (#PF005) includes oxidized glutathione (GSSG) and E. coli DsbC protein (disulfide bond isomerase in the periplasm of E. coli).

PDI Set (#PF006) includes oxidized glutathione (GSSG), human PDI (protein disulfide isomerase) and human Ero1 a (ER oxidoreductin-1 to reoxidize PDI).

Addition of DsbC Set or PDI Set to PURE frex® enables a protein containing disulfide bonds to be synthesized in an active form.

Efficiency of the formation of disulfide bonds is dependent on reducing agent in the reaction mixture. We recommend the use of PURE frex® 2.1 (#PF213) in which the suitable reducing agent can be selected.

Protocol

This is a standard protocol for synthesizing proteins containing disulfide bonds. Each solution of DsbC Set and PURE frex @ 2.1 (#PF213) are mixed together in a same tube. For example, 20 µL of reaction is assembled as below, which final concentration of each reagent will be 0.5 mM Cysteine, 4 mM GSH and 3 mM GSSG and 4 µM DsbC.

(Materials)

PUREfrex® 2.1 (#PF213) contains Solution I, Solution II, Solution III, Cysteins and GSH. DsbC Set (#PF005) contains DsbC and GSSG.

- 1. Thaw Solution I, Cysteine, GSH and GSSG by incubation at room temperature or 37 °C for 1 minute completely, and then cool on ice.
- 2. Thaw Solution II. III and DsbC on ice.
- 3. Mix each solution by vortex and centrifuge briefly to collect each solution at the bottom.
- 4. Dilute DsbC 4-fold with Dilution Buffer.
- 5. Assemble the reaction mixture in a tube as follows. (Add the template DNA to 1-3 ng/µL per 1 kbp)

Note

DsbC Set is developed for in vitro research use only. DsbC Set should not be used for the therapy, diagnostic or administration to animals including human and should not be used as food or cosmetics etc.

To avoid the contamination of nuclease, nuclease-freetreated water, reagents and materials should be used. We also recommend wearing gloves and mask.

For information concerning commercial use of DsbC Set, please contact GeneFrontier.

e-mail : purefrex@genefrontier.com



Distributor

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Protocol

| Water | 5-Χ μL |
|----------------|--------|
| Solution I *4 | 8 μL |
| 10 mM Cysteine | 1 μL |
| 80 mM GSH | 1 μL |
| 60 mM GSSG | 1 μL |
| Solution II | 1 μL |
| Solution III | 2 µL |
| 80 µM DsbC *⁵ | 1 μL |
| Template DNA | XμL |
| Total | 20 µL |

6. Incubate the tube at 37°C for 2-6 hours.

Protein synthesis reaction is almost done until 6 hours, but some proteins require longer incubation (e.g. 24 hours) to form disulfide bonds between the correct pair of cysteine residues.

7. Analyze the synthesized product.

*4)

Please note that the volume of Solution I of PUREfrex® 2.1 (#PF213) is different from PURE frex® 2.0 (#PF201).

*5) Please use 4-fold diluted DsbC

We recommend the optimizing the concentration of any reducing agent to achieve higher activity because it is depends on the synthesized protein.

The concentration unit has been changed from "mg / mL" to "µM" . 7.5mg / mL is equal to 320µM.

Standard final concentration of DsbC is 1 - 16 $\mu M.$ We recommend the optimizing the concentration of any reducing agent to achieve higher activity because it is depends on the synthesized protein Please use attached Dilution Buffer for dilution of DsbC.

*3)

For storage at -80°C, the rest of solution should be frozen rapidly in liquid nitrogen or dry ice/ethanol. Divide into aliquots, if necessary, and avoid refreeze and thaw as much as possible.

Kit components

Store at -80 °C before opening

- *1) Standard final concentration of GSSG is 3 mM.
- *2)