

Cat. # HB131/HB132

For Research Use

TAKARA

***Brevibacillus* Expression System
Intracellular Expression Vectors**

pNI DNA (Cat. #HB131)

pNI-His DNA (Cat. #HB132)

Product Manual

v201910

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I. Introduction

Brevibacillus (Bacillus brevis) Expression System is a high-efficiency protein production system. This gram-positive bacterium is characterized by its ability to secrete/produce large amount of proteins¹⁾. This characteristic has been successfully used for secretory production of a large number of heterologous proteins.

Recent studies have shown that this bacterium is also capable of intracellular protein expression. Proteins that are insoluble and precipitate when produced in *E. coli* can be recovered in a soluble state when produced by this bacterium. This system is recommended for the production of functional proteins that are endogenously produced intracellularly but are insoluble and cannot undergo *in vitro* refolding when produced in *E. coli*. This system has the following features:

- Efficient intracellular production of soluble heterologous proteins
- Simple procedures for genetic manipulation and culturing
- Almost no detectable protease activity
- Production of active protein
- Safe host bacteria

Using an *E. coli-Brevibacillus* shuttle vector, expression vectors can be constructed in *E. coli*. A vector with a his-tag inserted at the N-terminus that allows target protein purification using a nickel purification column is also available. The his-tag can be removed by treating the purified protein with enterokinase.

II. Component

pNI DNA (Cat. #HB131)	10 μ g (0.2 μ g/ μ l)
pNI-His DNA (Cat. #HB132)	10 μ g (0.2 μ g/ μ l)

【 Form 】 10 mM Tris-HCl, pH 8.0
1 mM EDTA

III. Storage

-20°C

*2 years from date of receipt under proper storage conditions.

IV. Overview of Intracellular Expression Vectors

The workflow to produce target proteins using this system is described below.

IV-1. Selection of expression vectors

(1) pNI DNA

pNI DNA is an intracellular expression vector constructed by removing the secretory signal segment from the secretory expression vector pNCMO2 DNA (Cat. #HB112) to allow intracellular accumulation of expressed proteins. The rest of its sequence is the same as pNCMO2 DNA. It is a shuttle vector between *Brevibacillus* and *E. coli*; the target gene is cloned into the vector in *E. coli*, and then the plasmid is transferred to *Brevibacillus* for protein expression.

pNI DNA uses the P2 promoter from the *Brevibacillus* host cell wall protein as promoter for expression of the target gene. The P2 promoter exhibits weak activity in *E. coli* and is useful for cloning target genes. It is a very strong promoter in *Brevibacillus* and is suitable for efficient protein production in *Brevibacillus*.

(2) pNI-His DNA

pNI-His DNA is an intracellular expression vector containing a his-tag sequence (6 x His) and an enterokinase recognition sequence for tag removal. This vector allows for easy purification of expressed proteins using nickel chelate resin. The rest of its sequence is the same as pNI DNA.

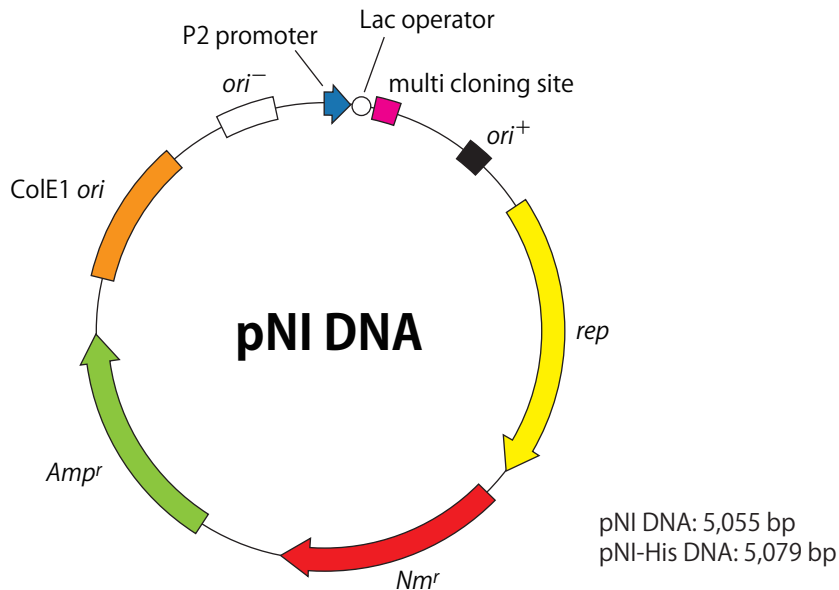
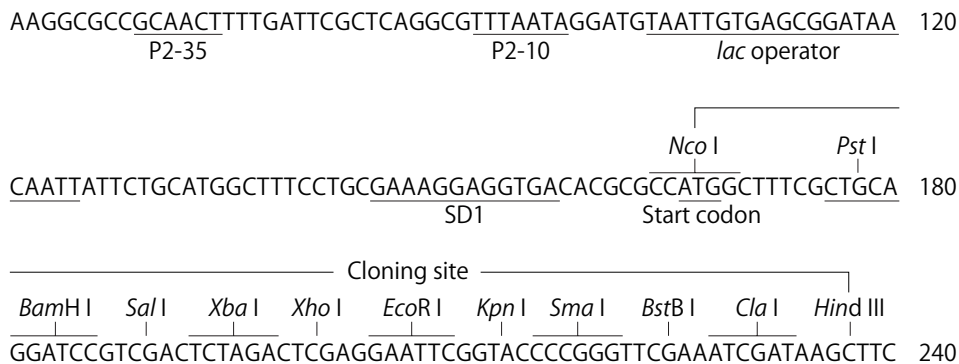


Figure 1. Vector Map of pNI DNA

<Features of pNI and pNI-His DNA>

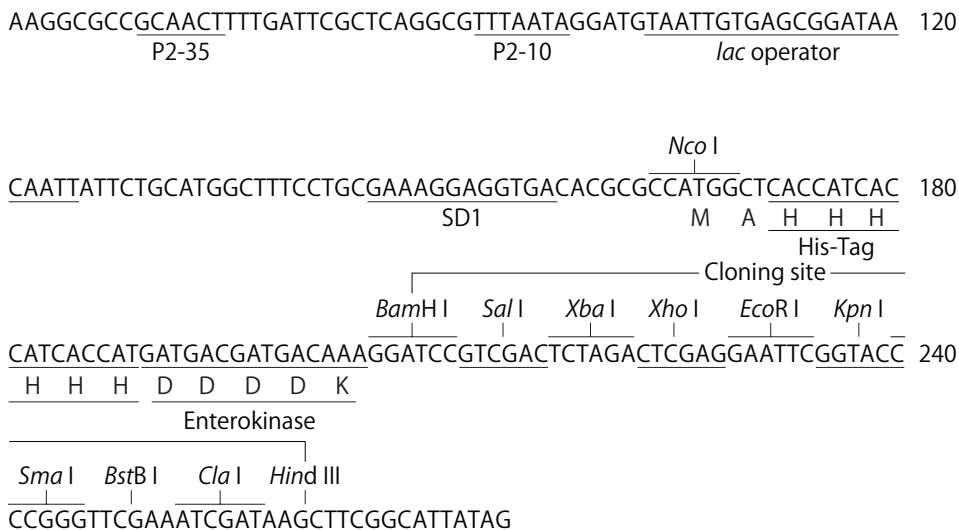
P2 promoter	A portion of 5' sequence upstream of the cell wall protein (HWP) gene, which is expressed strongly in <i>Brevibacillus</i>
His-tag + enterokinase recognition sequence	pNI-His DNA only
Multiple cloning site	12 restriction enzyme cleavage sites (pNI DNA) 10 restriction enzyme cleavage sites (pNI-His DNA)
Terminator	A 46-bp termination signal inserted downstream from the multiple cloning site.
<i>rep</i>	Gene involved in plasmid replication (derived from pUB110)
<i>Ori</i>	Replication origin allowing replication and maintenance of the plasmid in <i>Brevibacillus</i> (derived from pUB110)
<i>Nm^r</i>	Neomycin resistance gene, selection marker in <i>Brevibacillus</i>
<i>ColE1 ori</i>	Replication origin allowing replication and maintenance of the plasmid in <i>E. coli</i> (derived from pUC).
<i>Amp^r</i>	Ampicillin resistance gene, selection marker in <i>E. coli</i>

< Cloning Sites of pNI DNA >



GGCATTATAG

< Cloning Sites of pNI-His DNA >



IV-2. Cloning into expression vectors

For pNI DNA, the gene of interest is inserted into the multiple cloning site (MCS). If the *Nco* I site is usable, extra amino acids derived from the MCS will not be included in the resulting protein. Using a restriction site other than *Nco* I will result in the addition of extra amino acids to the N-terminus, depending on the site.

For pNI-His DNA, the gene of interest is inserted into a site downstream of *Bam*HI.

IV-3. Transformation of *Brevibacillus*

Transformation of *B. choshinensis* cells can be performed using the New Tris-PEG (NTP) method (Section V-4). The selection marker is neomycin resistance. When the shuttle vector is used in combination with *E. coli* host, ampicillin resistance can be used as the selection marker in *E. coli*.

IV-4. Detection of protein production and scale-up

Culture a negative control concurrently to confirm the expression of the target protein. Culture transformants harboring the target protein expression plasmid in the specified liquid medium with shaking for 48 to 64 hours to express the target protein. The bacterial cultures will be used for SDS-PAGE analysis, or a similar test, to confirm the presence/absence of protein expression. Scale up the culture volume when a large amount of product is required.

V. Protocol

V-1. *Brevibacillus* strain

Standard genetic engineering techniques are applicable.

V-1-1. Genotype

Because genes relating to sporulation have been disrupted, it is easy to perform sterilization of the strain. Additionally, disruption of the intracellular protease gene (*imp*) and extracellular protease gene (*emp*) minimizes degradation of expressed recombinant proteins.

V-1-2. Storage of recombinant *Brevibacillus*

Short-term storage (about 1 week)

1. Pick a single colony and spread on a MTNm plate.
2. Place the plate in an incubator at 30°C overnight.
3. Seal the plate and store it at room temperature (about 20°C).

Note: Do not store the plates in a refrigerator.

Long-term storage (1 month or longer)

1. Pick a single colony and inoculate into 2SYNm medium (see V-8. Medium components). Culture overnight on a shaker.
2. Transfer the broth to a freezing vial and add an equal amount of LB medium containing 40% glycerol.
3. Store at -80°C.
4. Thaw each vial immediately before use and inoculate at 0.1 - 1.0% volume of liquid medium.

Note: Do not subject cell stocks to repeated freeze-thaw cycles.

V-2. *E. coli* host

A lac operator is inserted in pNI DNA and pNI-His DNA to weaken the promoter activity in *E. coli*. For that reason, JM109 or another F factor integrated strain inserted with an F factor (*lacI^q*) must be used. The genotype of JM109 is shown below for reference.

JM109: *recA1, endA1, gyrA96, hsdR17, thi-1, hsdR17 (r_K⁻ m_K⁺), e14⁻ (mcrA⁻), supE44, relA1, Δ (lac-proAB) /F' [traD36, proAB⁺, lac I^q, lacZ Δ M15]*

V-3. Construction of an expression plasmid using pNI DNA and pNI-His DNA

Precautions for the construction of expression plasmids using pNI DNA and pNI-His DNA

E. coli strains containing *lac I^q* and *recA⁻*, such as JM109, are recommended as the host for plasmid construction.

Insert DNA should be cloned in-frame, downstream from the tag sequence.

Insert a stop codon at the end of the target gene to be cloned.

V-3-1. Cloning insert DNA into pNI DNA or pNI-His DNA

< Gene amplification using PCR >

Design PCR primers so that the target gene is inserted downstream of the tag sequence. Introduce two types of restriction enzyme sites at the ends of the PCR product to establish directionality. Amplify the target gene by PCR. Select PCR conditions based on the primers and PCR enzyme used in the reaction. The use of a high-fidelity PCR enzyme, such as PrimeSTAR[®] Max DNA Polymerase (Cat. #R045A), is recommended.

< Construction of the expression plasmids through ligation >

Select two restriction enzyme sites in the expression vector multiple cloning site that match both ends of the target DNA insert to establish directionality. Treat 0.5 - 1.0 μg of the insert and vector DNA with the restriction enzymes. Perform agarose gel electrophoresis on both the vector and insert DNA restriction enzyme reactions, then collect and purify the target fragments.

Ligate - 100 ng of each purified DNA fragment using a ligation reagent such as DNA Ligation Kit <Mighty Mix> (Cat. #6023). Use a portion of the reaction solution for transformation of *E. coli*.

< Cloning with In-Fusion[®] Cloning System >

The In-Fusion HD Cloning Kit can be used to simply and quickly carry out directional cloning even when there are no appropriate restriction enzyme sites. Refer to the In-Fusion Cloning System protocol.

< Transformation of *E. coli* >

Use an *E. coli* host with high transformation efficiency. *E. coli* JM109 Competent Cells (Cat. #9052) or *E. coli* JM109 Electro-Cells (Cat. #9022) are recommended.

V-3-2. Analysis of recombinants

After spreading 100 - 200 μ l of the transformation mixture onto LB plates containing 50 - 100 μ g/ml ampicillin, incubate the plates at 37°C for 15 - 18 hours. Select 10 - 20 ampicillin-resistant colonies and inoculate into 2 ml of LB liquid medium containing 50 - 100 μ g/ml ampicillin. Grow for 15 - 18 hours at 37°C with shaking and isolate plasmid DNA. Perform restriction enzyme analysis using an appropriate amount of DNA. Usually the same enzymes that were used for the construction of the expression plasmid are used. After confirming the presence of the insert, perform sequencing to confirm that the gene contains no PCR-introduced errors and is in-frame with tag-sequence in pNI-His.

V-3-3. Sequencing

The following forward and reverse primers can be used to confirm the insert DNA sequence. (Primer sequences are shared by pNI DNA and pNI-His).

Forward Sequencing Primer: 5'-TCGAAGGCGCCGCAAC-3'

Reverse Sequencing Primer: 5'-CAATGTAATTGTTCCCTACCTGC-3'

V-4. Transformation of *Brevibacillus*

V-4-1. Preparation

Brevibacillus Competent Cells (Cat. #HB116)
(Contents) *Brevibacillus* Competent Cells
MT medium
Solution A
Solution B
Plasmid for target gene expression
Plasmid for negative control (blank vector)
MTNm plates*
Culture tubes
Sterilized microtubes

* Refer to V-8. Medium Components

V-4-2. NTP Transformation Method

- (1) Thaw Solution A, Solution B, and MT medium.
- (2) Remove only the number of tubes of *Brevibacillus* Competent Cells needed for transformation from storage, and keep on dry ice/ethanol.
- (3) Thaw the *Brevibacillus* Competent Cells quickly (approximately 30 sec) in a 37°C water bath.
- (4) Centrifuge the cells (12,000 rpm for 30 sec to 1 min) to form a cell pellet and remove the supernatant with a micropipette.

Perform the following procedures at room temperature.

- (5) Mix the plasmid DNA solution (in a volume of 5 μ l or less)*¹ with 50 μ l of Solution A.
- (6) Add all of the DNA solution to the bacterial cell pellet (from step 4) and vortex to completely suspend the bacterial pellet.*²
- (7) Allow to stand for 5 minutes at room temperature.
- (8) Add 150 μ l of Solution B (PEG solution)*³ and vortex until the solution is uniform (5 - 10 sec).
- (9) Centrifuge the cells (5,000 rpm for 5 min) and remove the supernatant.
- (10) Centrifuge briefly (5,000 rpm for 30 sec) and remove the supernatant completely.
- (11) Add 1 ml of MT medium and suspend completely with a micropipette.
- (12) Incubate at 37°C in an orbital shaker (120 rpm for 2 hours).
- (13) Use a sterile inoculating loop to remove a small sample from the culture. Streak on the MTNm plates and culture overnight at 37°C.
- (14) Select isolated colonies for plasmid analysis or protein expression.

*1 When DNA ligation solution is used, mix 5 μ l of the reaction solution with Solution A. When using purified plasmids, use 10 - 100 ng.

*2 Be sure to suspend well, as incomplete dispersion of cells will decrease the efficiency of transformation.

*3 Solution B (PEG solution) is highly viscous-use a 1,000 μ l micropipette and pipette slowly.

V-5. Target proteins expression with recombinant *Brevibacillus*

After the construction of expression strain is complete, perform a small-scale protein expression analysis. A standard protocol for verifying expression is given here.

V-5-1. Overview

After a positive clone has been identified, check expression together with a negative control.

The production of the target protein may vary among transformants depending on the nature of the protein. The size of colonies may also vary. Thus, six to ten colonies should be selected at random (including both large and small colonies) for culturing in test tubes. If the plates are left for several days after transformation, protein production may be decreased. If this occurs, repeat the transformation.

V-5-2. Culture media

Use 2SYF medium as a basic medium for expression analysis.

V-5-3. Culture for protein production

By growing cultures of a transformant expressing the target protein and a negative control, expression of a target protein can be confirmed.

The protocol for expression analysis is shown below.

- (1) Select individual colonies and inoculate into 3 ml 2SYFNm liquid medium in culture tubes (\varnothing 16 mm). Incubate at 30 - 33°C with shaking at 120 rpm for 48 - 64 hours. Perform sampling every 24 hours to check target protein production.
- (2) At the end of incubation, isolate the cells by centrifugation at 5,000g for 5 minutes.
- (3) Suspend the cells in PBS*¹ and sonicate to disrupt the cells*². Make sure to perform this step in an ice bath to avoid increases in temperature.
- (4) Separate the supernatant and precipitate in the cell homogenate by centrifugation at 20,000g for 10 minutes. Suspend the precipitate in a volume of PBS equal to the volume of the supernatant.
- (5) Evaluate protein expression in the pellet fraction using SDS-PAGE (CBB staining or Western blotting) or measure protein activity.

* 1 Can be prepared using PBS Tablets (Cat. #T900).

* 2 The standard sonication conditions when using a BRANSON SONIFIER 350 are 50% duty, output 4 to 5, and 30 sec x 2. Select the optimal conditions when using a different device.

V-6. SDS-PAGE analysis

Use an SDS-PAGE gel suitable for isolating the target protein.

V-6-1. Sample preparation

Add 10 μ l of 5X SDS-PAGE loading buffer to 40 μ l of cell supernatant and precipitate suspension.

Mix and heat at 100°C for 10 minutes to prepare samples for electrophoresis.

V-6-2. Control

Use following samples as controls.

- a. Molecular weight marker
- b. Target protein standard
- c. Sample from the culture of *B. choshinensis* SP3 harboring a vector without the insert (negative control)

V-6-3. Analysis of protein expression

Target protein expression can be confirmed by comparing the position and the density of the specific band with the standard protein by SDS-PAGE. If protein production is low, it is insoluble, or there are other protein contaminants, it may be difficult to detect the target protein by SDS-PAGE. In those cases, Western blot analysis using a tag-specific antibody, functional evaluation (e.g., specific activity), or protein purification may be useful for detection of protein production.

If pNI-His DNA is used as the expression vector, Western blotting with an anti-his-tag antibody can be used to detect the target protein.

V-7. Purification of expressed protein

Purification methods vary according to the type of target protein. To purify a non-his-tagged target protein, use an ordinary purification method (e.g., ion exchange, hydrophobic, or affinity chromatography).

If pNI-His DNA is used as the expression vector, the target protein can be easily purified using TALON® Metal Affinity Resin (Cat. #635501) or another histidine-tagged protein purification resin. After purification, if needed, the his-tag may be removed by treatment of the purified product with enterokinase. For purification using a histidine-tagged protein purification resin and enterokinase treatment, refer to the manual for each product.

V-8. Medium Components

- 2SYF liquid medium

Components

Fructose*	20.0 g/L
Bacto Soytone	40.0 g/L
Bacto Yeast Extract	5.0 g/L
CaCl ₂ · 2H ₂ O	0.15 g/L
Adjust to pH 7.2 with NaOH	

* Mix fructose and CaCl₂ and filter-sterilize separately from the other components. Mix after sterilization.

- 2SYF Nm liquid medium

Add neomycin solution (stock solution 50 mg/ml) to 2SYF liquid medium to the concentration of 50 μg/ml.

- 2SY liquid medium

Components

Glucose*	20.0 g/L
Bacto Soytone	40.0 g/L
Bacto Yeast Extract	5.0 g/L
CaCl ₂ · 2H ₂ O	0.15 g/L
Adjust to pH 7.2 with NaOH	

* Mix Glucose and CaCl₂ and filter-sterilize separately from the other components. Mix after sterilization.

- 2SY Nm liquid medium

Add neomycin solution (stock solution 50 mg/ml) to 2SYF liquid medium to the concentration of 50 μg/ml.

- TM liquid medium

Components

Glucose	10.0 g/L
Phytone Peptone	10.0 g/L
35%Ehrlich Bonito Extract	5.75 g/L
Yeast extract Blue label	2.0 g/L
FeSO ₄ · 7H ₂ O	10 mg/L
MnSO ₄ · 4H ₂ O	10 gm/L
ZnSO ₄ · 7H ₂ O	1 mg/L
Adjust to pH 7.0 with NaOH	

* Autoclave glucose separately from the other components. Mix after sterilization.

- TMNm Broth
Add neomycin solution (stock solution: 50 mg/ml) to TM liquid medium to the concentration of 50 μ g/ml.

- MT liquid medium

Components

Glucose	10.0 g/L
Phytone Peptone	10.0 g/L
35%Ehrlich Bonito Extract	5.75 g/L
Yeast extract Blue label	2.0 g/L
FeSO ₄ · 7H ₂ O	10 mg/L
MnSO ₄ · 4H ₂ O	10 gm/L
ZnSO ₄ · 7H ₂ O	1 mg/L
MgCl ₂	4.1 g/L
Adjust to pH 7.0 with NaOH	

* Autoclave glucose separately from the other components. Mix after sterilization.

- MTNm plate
Add 7.5 g of agar to 500 ml of MT liquid medium and sterilize by autoclaving. Allow the medium to cool to about 50°C before adding neomycin solution (stock solution 50 mg/ml) to a final concentration of 50 μ g/ml, mix gently, and dispense to plates.

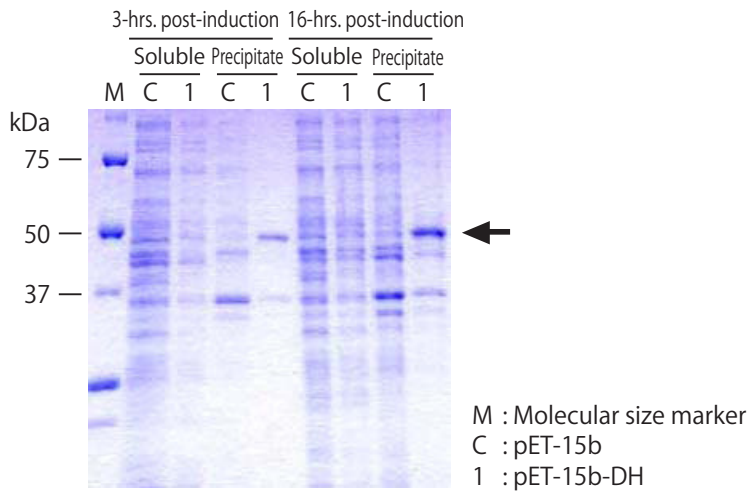
For the components of the 2SYF medium, the 2SY medium, the TM medium and the MT medium, the following manufacturers are recommended.

Bacto Soytone	(Becton Dickinson, Code. 243620)
Bacto Yeast Extract	(Becton Dickinson, Code. 212750)
Phytone Peptone	(Becton Dickinson, Code. 211906)
35%Ehrlich Bonito Extract	(Kyokuto Pharmaceutical, Code. 551-01212-5)
Yeast extract Blue label	(Oriental Yeast Co., Ltd.)

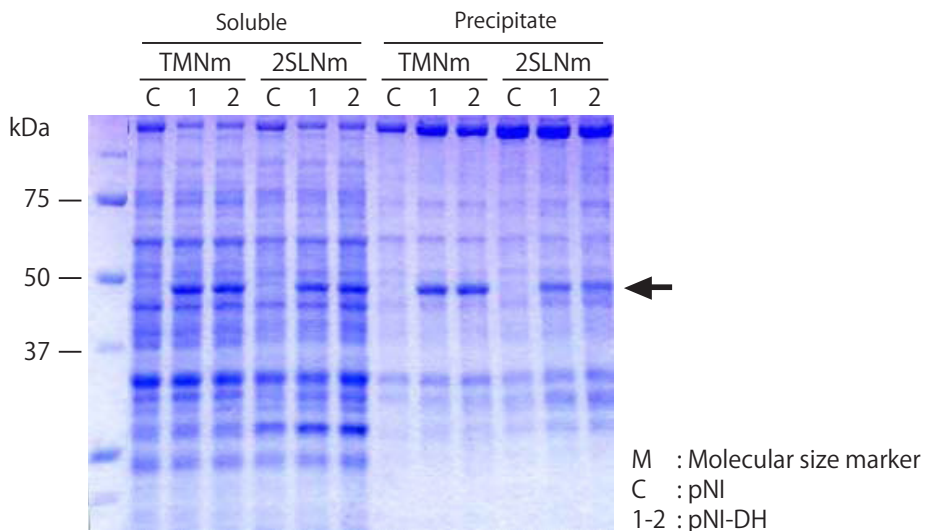
VI. Experimental Example: Production of a Eukaryotic Protein Using pNI

Eukaryotic dehydrogenase (DH, approximately 50 kDa) was produced using the pNI vector. When this protein was expressed using the pET vector system in *E. coli*, most of the protein was present in inclusion bodies (Figure 2. A). The secretory expression system using *Brevibacillus* did not yield any transformants. Intracellular expression (30°C, 48 hours) using the pNI vector yielded target protein in the soluble fraction from the cell homogenate (Figure 2. B) and 10 to 20-fold greater enzyme activity compared with protein expressed in *E. coli* (Figure 2. C). Optimization of the culture conditions improved the yield further by approximately 5-fold.

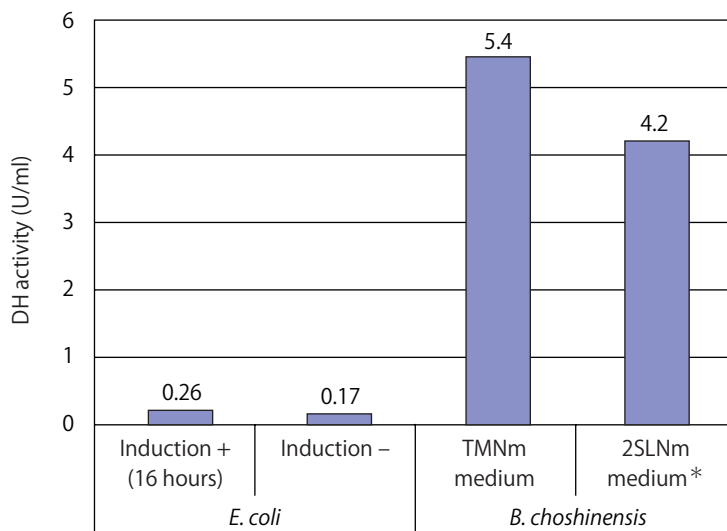
A. Host: *E. coli* BL21



B. Host: *Brevibacillus* SP3



C. Comparison of Enzyme Activity



* 2SLNm medium:

2% glucose, 4% soy peptone, 0.5% yeast extract, 0.001% iron sulfate, 0.001% manganese sulfate, 0.0001% zinc sulfate, 50 μ g/ml neomycin (pH 7.2)

Figure 2. A Comparison of Eukaryotic Dehydrogenase (DH) Expression Using pNI and pET vectors.

VII. Related Products

[*Brevibacillus* Secretory Expression System]

- BIC System (Cat. #HB300)
- pBIC DNA Set (Cat. #HB310)
- Brevibacillus* Expression System II (Cat. #HB200)
- Brevibacillus* Competent Cells (Cat. #HB116)
- pNY326 DNA (Cat. #HB111)
- pNCMO2 DNA (Cat. #HB112)
- pNY326-BLA DNA (Cat. #HB114)
- pNC-HisT DNA (Cat. #HB121)
- pNC-HisF DNA (Cat. #HB122)
- pNC-HisE DNA (Cat. #HB123)

[Histidine Tagged Protein Purification]

- HisTALON™ Superflow Cartridge (Cat. #635650)
- HisTALON™ Buffer Set (Cat. #635651)
- TALON® Metal Affinity Resin (Cat. #635501/635502/635503/635504/635652/635653)

[Other]

- E. coli* JM109 Competent Cells (Cat. #9052)
- E. coli* JM109 Electro-Cells (Cat. #9022)
- DNA Ligation Kit <Mighty Mix> (Cat. #6023)
- PrimeSTAR® Max DNA Polymerase (Cat. #R045A)
- PBS (Phosphate Buffered Salts) Tablets (Cat. #T900)

VIII. References

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- 2) T. Takano, A. Miyauchi, H. Takagi, K. Kadowaki, K. Yamane, and S. Kobayashi. Expression of the Cyclodextrin Glucanotransferase Gene of *Bacillus macerans* in *Bacillus brevis*. *Biosci Biotech Biochem.* (1992) **56**(5): 808-809.
- 3) H. Tojo, T. Asano, K. Kato, S. Udaka, R. Horinouchi, and A. Kakinuma. Production of Human Protein Disulfide Isomerase by *Bacillus brevis*. *J Biotechnol.* (1994) **33**(1): 55-62.
- 4) H. Yamagata, K. Nakahama, Y. Suzuki, A. Kakinuma, N. Tsukakoshi, and S. Udaka. Use of *Bacillus brevis* for efficient synthesis and secretion of human epidermal growth factor. *Proc Natl Acad Sci USA.* (1989) **86**: 3589-3593.
- 5) Y. Takimura, M. Kato, T. Ohta, H. Yamagata, and S. Udaka. Secretion of Human Interleukin-2 in Biologically Active Form by *Bacillus brevis* Directly into Culture Medium. *Biosci Biotechnol Biochem.* (1997) **61**(11): 1858-1861.
- 6) K. Yashiro, J. W. Lowenthal, T. E. O'Neil, S. Ebisu, and H. Takagi. High-Level Protein Production of Recombinant Chicken Interferon- γ by *Brevibacillus choshinensis*. *Protein Expression and Purification.* (2001) **23**: 113-120.

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