Coexpression of Alginate Lyase with Hyperthermophilic Archaea Chaperonin in E. coli

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When the alginate lyase gene (aly) from Pseudoalteromonas elyakovii IAM 14594 was expressed in E. coli, most of the gene product expressed was produced as aggregated insoluble particles known as inclusion bodies. In order to produce with an elevated level of a soluble and active form of alginate lyase in E. coli, the hyperthermophilic chaperonins (ApCpnA and ApCpnB) from archaeon Aeropyrum pernix K1 were employed as the coexpression partners. At 25℃ culture temperature, the level of alginate lyase activity was increased from 10.1 unit/g-soluble protein in aly single expression to 83.1 unit/g-soluble protein by coexpressing with ApCpnA and to 100.3 unit/g-soluble protein by coexpressing with ApCpnB. This results indicate that the coexpression of aly with ApCpnA and ApCpnB revealed a marked enhancement, about 8~10 fold, in the production of alginate lyase as a soluble and active form. Based on the results of various examinations on the expression variables, the optimal conditions for the maximal production of alginate lyase were determined as 1.0 mM IPTG for the inducer concentration, 25℃ for the culture temperature after IPTG induction, and ApCpnB for the coexpression partner. The coexpression set in the present report may be useful in the industrial production of functionally or medically important recombinant proteins in E. coli.

Key words: Aeropyrum pernix, alginate lyase, chaperonin, coexpression, E. coli

Introduction

Brown seaweed contains up to 67% (w/w) carbohydrates including materials such as alginate, laminaran, mannitol, etc. The most abundant carbohydrate in the brown seaweed is alginate that constitutes 17-45% of total biomass [25]. Alginates are linear uronic acid polymers in which β-D-mannurionate and its C-5 epimer, α-L-gulurionate residues are linked to form blocks of polyguluronate and random sequences. These residues are arranged in block structures which can be homopolymeric [poly (β-D-mannosyluronic acid) and poly (α-L-gulosyluronic acid)] or heteropolymeric, i.e. containing random blocks [6].

The proportion and arrangement of the block structures vary greatly in alginates from different sources and determine the physical properties of the polymer, particularly the ability to form gels in the presence of divalent cations [6]. Marine alga alginate is used widely in the food, pharmaceutical, textile, and oil industries due to its ability to chelate metal ions and form a highly viscous solution, recently, polymers and oligosaccharides with novel physicochemical and physiological functions are sought by biopolymer-based industries in order to expand the application areas of polysaccharides [27]. Alginate degrading enzyme, alginate lyases, also known as alginas or alginate depolymerases, catalyze the degradation of alginate by a β-elimination mechanism that has yet to be fully elucidated. Alginate lyases are isolated from various sources, such as marine algae, marine mollusks, fungi, bacteria, and viruses. Many bacteria, such as Pseudoalteromonas elyakovii, Sphingomonas sp., Klebsiella pneumoniae, and Pseudomonas sp. have been found to produce a large amount of alginate lyases [2, 26]. Of these enzymes, unique substrate specificity has been discovered in an alginate lyase produced by P. elyakovii, which is capable of degrading all block structures derived from sodium alginate and produces a series of tri- to octa-oligouronates [19, 20].

When the alginate lyase gene (aly) from P. elyakovii was expressed in E. coli, it yielded inactive aggregates known as inclusion bodies [12, 21].
Various plasmid systems have been constructed for the coexpression of molecular chaperones and foldases from E. coli, and it has been successfully demonstrated that their coexpression increases the formation of active proteins [8, 9, 13, 16-18, 21, 22]. However, there is relatively little information on the comparative effects of coexpressing a series of folding accessory proteins on the productivity of active target proteins, even though such knowledge is important to elucidate the criteria for selecting appropriate folding accessory proteins according to the properties of target proteins.

The role of the subunit heterogeneity or α- and β-subunit cooperativity in group II chaperonins has not been systematically addressed [5, 7, 23]. One intriguing possibility is that the heterogeneity is directly linked to substrate specificity. A number of biochemical studies using endogenous model substrates suggest that each subunit contributes to the recognition of specific motifs within the substrates [3, 11]. Especially, the substrate specificity of each subunit on the heterologous proteins in E. coli has not been elucidated in details.

To study the substrate specificity and to improve the production level of active alginate lyase enzyme in E. coli, α-subunit and/or β-subunit of hyperthermophilic group II chaperonin from A. pernix K1 were employed to coexpress with alginate lyase (aly) from P. elyakovii. In this work, we have investigated and reported on the effects of hyperthermophilic chaperonin α- and/or β-subunits from A. pernix K1 on the soluble and active production of alginate lyase in E. coli.

**Materials and Methods**

**Bacterial strain and plasmids**

E. coli Rosetta (DE 3) strain was used in all experiments. Recombinant plasmids, pALP4, pET3d-ApCpnA, pET21a-ApCpnB, and pG-KJE6 were used in this work [11, 21]. The alginate lyase gene (aly, 1.19 kb) from P. elyakovii [19] was subcloned into the plasmid pALP4 (6.7 kb) [21]. The hyperthermophilic chaperonin A and B genes (ApCpnA and ApCpnB) from A. pernix K1 was subcloned into the plasmid pRSFDeut-1 vector (Invitrogen, USA), resulting in the plasmid ApCpnA (5.4 kb) and ApCpnB (5.4 kb), respectively. The plasmid pG-KJE6 encoding dnaK-dnal-grpE and GroES-GroEL of E. coli was also used as a coexpression partner for the active production of alginate lyase [21]. The plasmids, pALP4, ApCpnA, ApCpnB, and pG-KJE6 were transformed into E. coli Rosetta (DE 3) by CaCl2 method, respectively or simultaneously [14]. The transformed E. coli cells were selected on LB agar plates containing 50 μg/ml ampicillin (selection for pALP4), 50 μg/ml kanamycin (selection for ApCpnA and ApCpnB), and 50 μg/ml chloramphenicol (selection for pG-KJE6).

**Culture media and culture condition**

The transformed E. coli cells were grown and selected on LB agar plates containing 50 μg/ml ampicillin, 50 μg/ml kanamycin, and 50 μg/ml chloramphenicol. Isopropyl-1-thio-β-D-galactopyranoside (IPTG) was used for the expression of aly, ApCpnA, and ApCpnB. To induce the expression of groEL/ES and dnaK/dnak/grpE, tetracycline and L-arabinose were used, respectively. After all cells were preincubated in LB media supplemented with appropriate antibiotics at 37°C, when the cell density (OD600) reached at 0.6~0.8, IPTG was added to LB medium at a concentration of 0~2 mM, followed by continuous incubation for 6 hr.

**Measurement of cell growth and protein concentration**

Cell growth was estimated by optical density at 600 nm (OD600) with a spectrophotometer (Shimadzu, Japan). Cells were disrupted by sonication (1 min, 70 Watt, and 7 sec cycle on ice) with a sonicator (Sonoplus HD2070, Bandelin, Germany), and then centrifuged at 9,800×g for 10 min for the separation into the soluble and insoluble fractions. The protein concentrations of total cell lysates and soluble and insoluble fractions were determined by the modified micro method of Lowry by using bovine serum albumin as a standard.

**SDS-PAGE analysis**

The soluble and insoluble fractions obtained from 10 μg cell lysate protein was analyzed by SDS-PAGE (14% gel). The ApCpnA and ApCpnB proteins were detected by staining gels with Coomassie brilliant blue RT250.

**Assay of alginate lyase activity**

The assay for alginate lyase activity was conducted as follows. In each assay, reaction products were confirmed to increase proportionally with time and enzyme concentration in the reaction mixture. Alginate lyase assayed in a mixture containing 0.2% sodium alginate (Sigma, USA) in 1 M Tris-HCl (pH 7.5) and 0.3 M NaCl. The reaction was monitored at 37°C for 5 min, depending on the increase in absorbance
at 235 nm in comparison with that without enzyme. One unit of alginate lyase was defined as the amount of enzyme required to increase the absorbance by 0.1 at 235 nm per min [21].

**Results and Discussion**

**Expression of alginate lyase (aly) in E. coli**

To optimize the expression of alginate lyase (aly), the E. coli transformant harboring pALP4 plasmid was cultivated on LB medium with various IPTG concentrations. As shown in Fig. 1, the 51 kDa protein band corresponding to the molecular weight of alginate lyase was clearly detected in the insoluble fraction of cell lysate and had no differences in the protein intensity between IPTG concentrations. However, the activity of alginate lyase of the cell lysate was peaked at 1 mM IPTG (Fig. 2). Since the most of alginate lyase was expressed as an insoluble body in the insoluble fraction [21], the enzyme activity in the cell lysate was very low at the level of 3.6 unit/g-soluble protein. The majority of alginate lyase activity was detected in the soluble fraction and is probably due to a high expression rate leading the protein to accumulate with an abnormal conformation [21].

**Expression of hyperthermophilic archaea chaperonin in E. coli**

For the maximal expression of ApCpnA and ApCpnB, IPTG with different concentrations was added in the culture of E. coli cell harboring ApCpnA or ApCpnB plasmid. After IPTG induction for 6 hr, the proteins in the soluble and insoluble fractions were analysed by SDS-PAGE. As shown in Fig. 3, the protein bands corresponding to ApCpnA (60.7 kDa) and ApCpnB (61.2 kDa) were found in the both soluble and insoluble fractions at 1 mM IPTG. After heat treated at 85°C for 20 min of the soluble fractions, most of intracellular proteins of E. coli itself were removed as the precipitates by centrifugation and the supernatant revealed the homogeneous protein bands of thermophilic ApCpnA and ApCpnB (Fig. 4).

**Effect of culture temperature and coexpression of APCpnA and ApCpnB on the soluble production of alginate lyase**

In order to investigate the effects of culture temperature and coexpression system of archaea chaperonin on the synthesis and solubilization of alginate lyase, two transformants harboring plasmids of pALP4 and ApCpnA, and pALP4 and ApCpnB were cultivated on LB media for 37°C, respectively. When the cell concentration of OD<sub>600</sub> reached at 0.8, 1 mM IPTG was added into the culture media and then the cells were incubated for more 6 hr at different temperatures (15~37°C). The highest activity of alginate lyase in both cases of aly single expression and coexpression with archaea chaperonins was detected at 25°C (Fig. 5). At 25°C, the level of alginate lyase activity was increased from 10.1 unit/g-soluble protein in aly single expression to 83.1 unit/g-soluble protein by coexpressing with ApCpnA and to 100.3 unit/g-soluble protein by coexpressing with ApCpnB. Therefore, it can suggested that the optimal culture temperature for the soluble and active production of alginate lyase with or without chaperonin is 25°C and ApCpnB is the preferable coexpression partner than ApCpnA. Due to
the decreased or retarded rate of protein synthesis, low culture temperature below 37°C has been reported as one of prerequisite parameters for the overexpression of various heterologous genes in *E. coli* [24, 28]. The reason of ApCpnB as the optimal coexpression partner than ApCpnA for the overproduction of alginate lyase is not clear at present, but the specificity or affinity toward unfold misfolded substrate, i.e., alginate lyase, of ApCpnB seems to be greater than that of ApCpnA [1, 7].

**Comparison with archaea chaperonin and *E. coli* molecular chaperones**

To compare the coexpression effect on the production of active alginate lyase, molecular chaperones such as GroEL/GroES and DnaK/DnaJ/GrpE of *E. coli* itself, and archaea chaperonins (ApCpnA and ApCpnB) were employed as co-expression partners of *aly*, in which IPTG concentration were 20 μM and 1 mM. As shown in Fig. 6, 1 mM IPTG for all expression systems resulted in higher expression of *aly* over 20 μM. In our previous report [20], the activity of alginate lyase in *E. coli* by coexpressing GroEL/GroES and DnaK/DnaJ/GrpE was about 20.0 unit/g-soluble protein at 20 μM IPTG. In addition, the coexpression system of archaea chaperonin gave greater benefit on the expression of *aly* than *E. coli* molecular chaperones.

Recently, it was reported that the promotion of proteolysis of target proteins was observed in DnaK and GroEL sets as folding assistant partner, resulting in the reduced yield or decreased expression level of foreign polypeptides [4, 15]. In our archaea chaperonin system, this proteolysis on alginate lyase produced is likely less profound than GroEL/GroES and DnaK/DnaJ/GrpE sets.

In conclusion, the coexpression system employing archaea chaperonins rather than *E. coli* itself molecular chaperones (GroEL/GroES and DnaK/DnaJ/GrpE sets) could effectively

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**Fig. 3.** Effect of IPTG concentration on the expression of ApCpnA (left) and ApCpnB (right) in the recombinant *E. coli* Rosetta (DE 3) harboring ApCpnA or ApCpnB plasmids.

**Fig. 4.** Homogeneity of ApCpnA and ApCpnB after 85°C for 20 min heat-shock treatment.

**Fig. 5.** Effect of coexpression of ApCpnA and ApCpnB on the alginate lyase production in the recombinant *E. coli* Rosetta/pALP4+ApCpnA and *E. coli* Rosetta/pALP4+ApCpnB at different culture temperatures.
improve the production of active and soluble alginate lyase. The optimal combination for the maximal production of alginate lyase was determined as 1.0 mM IPTG, 25°C for the culture temperature after IPTG induction, and ApCpnB co-expression partner. The coexpression system in the present report may be useful in the industrial production of functionally or medically important recombinant proteins in E. coli.

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References

1. Bigotti, M. G. and Clarke, A. R. 2005. Cooperativity in the thermosome. J. Mol. Biol. 348, 13-26.
2. Chavagnat, F., Duez, C., Guinand, M., Potin, P., Barbeyron, T., Henrisat, B., Wallach, J. and Ghuysen, J. M. 1996. Cloning, sequencing and overexpression in Escherichia coli of the alginate lyase-encoding aly gene of Pseudomonas algivora: identification of three classes of alginate lyases. Biochem. J. 319, 575-583.
3. Frydman, J. 2001. Folding of newly translated proteins in vivo: the role of molecular chaperones. Annu. Rev. Biochem. 70, 603-647.
4. García-Fruitós, E., Martínez-Alonso, M., González-Montaltan, N., Valls, M., Mattanovich, D. and Villaverde, A. 2007. Divergent genetic control of protein solubility and conformational quality in Escherichia coli. J. Mol. Biol. 374, 195-205.
5. Gutsche, I., Essen, L. O. and Baumeister, W. 1999. Group II chaperonins: new TRiC (k)s and turns of a protein folding machine. J. Mol. Biol. 293, 295-312.
6. Haug, A., Larsen, B. and Baardseth, E. 1969. Comparison of the constitution of alginates from different sources. Proc. Int. Seaweed Symp. 6, 443-451.
7. Kim, J. H., Lee, J. W., Shin, E. J. and Nam, S. W. 2011. Cooperativity of α- and β-subunits of group II chaperonin from the hyperthermophilic archaeum Aeropyrum pernix K1. J. Microbiol. Biotechnol. 21, 212-217.
8. Kohda, J., Endo, Y., Okumura, N., Kurokawa, Y., Nishihara, K., Yanagi, H., Yura, T., Fukuda, H. and Kondo, A. 2002. Improvement of productivity of active form of glutamate racemase in Escherichia coli by coexpression of folding accessory proteins. J. Biochem. Eng. 10, 39-45.
9. Kurokawa, Y., Yanagi, H. and Yura, T. 2000. Overexpression of protein disulfide isomerase DsbC stabilizes multiple-disulfide-bonded recombinant protein produced and transported to the periplasm in Escherichia coli. Appl. Environ. Microbiol. 66, 3960-3965.
10. Lee, J. W., Kim, S. W., Kim, J. H., Jeon, S. J., Kwon, H. J., Kim, B. W. and Nam, S. W. 2013. Functional characterization of the α- and β-subunits of a group II chaperonin from Aeropyrum pernix K1. J. Microbiol. Biotechnol. 23, 818-825.
11. Ma, L. Y., Chi, Z. M., Li, J. and Wu, L. F. 2008. Overexpression of alginate lyase of Pseudoalteromonas elyakovii in Escherichia coli, purification, and characterization of the recombinant alginate lyase. World J. Microbiol. Biotechnol. 24, 89-96.
12. Makino, T., Skretas, G. and Georgiou, G. 2011. Strain engineering for improved expression of recombinant proteins in bacteria. Microb. Cell Fact. 10, 32-41.
13. Mandel, M. and Higa, A. 1970. Calcium-dependent bacteriophage DNA infection. J. Mol. Biol. 53, 159-162.
14. Martínez-Alonso, M., García-Fruitós, E., Ferrer-Miralles, N., Rinas, U. and Villaverde, A. 2010. Side effects of chaperone gene co-expression in recombinant protein production. Microb. Cell Fact. 9, 64-69.
15. Nishihara, K., Kanemori, M., Yanagi, H. and Yura, T. 2000. Overexpression of trigger factor prevents aggregation of recombinant proteins in Escherichia coli. Appl. Environ. Microbiol. 66, 884-889.
16. Priya, S., Sharma, S. K. and Goloubinoff, P. 2013. Molecular chaperones as enzymes that catalytically unfold misfolded polypeptides. FEMS Lett. 587, 1981-1987.
17. Qiu, J., Swartz, J. R. and Georgiou, G. 1998. Expression of active human tissue-type plasminogen activator in Escherichia coli. Appl. Environ. Microbiol. 64, 4891-4896.
18. Sawabe, T., Takahashi, H., Ezura, Y. and Gacesa, P. 2001. Cloning, sequence analysis and expression of Pseudoalteromonas elyakovii IAM 14594 gene (aly/PEEC) encoding the extracellular alginate lyase. Carbohydr. Res. 335, 11-21.
19. Sawabe, T., Takahashi, H., Saeki, H., Niwa, K. and Aono, H. 2007. Enhanced expression of active recombinant alginate lyase Aly/PEEC cloned from a marine bacterium Pseudoalteromonas elyakovii in Escherichia coli by calcium compounds. Enzyme Microb. Technol. 40, 285-291.
초록: 대장균에서 초고온성 샤페로닌과 alginate lyase의 공발현

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Pseudoalteromonas elyakovii 유래 alginate lyase 유전자 (aly)를 대장균에서 발현시켰을 때, 발현된 대부분의 유전자 산물은 내포체라는 불용성 응집체 형태로 생산되었다. Alginate lyase를 가용성 및 활성형으로 생산하기 위해 Aeropyrum pernix K1 유래의 초고온성 샤페로닌 ApCpnA와 ApCpnB를 공발현 파트너로 도입하였다. aly와 ApCpnA와 ApCpnB 각각과의 공발현 결과, aly 단독발현 때의 alginate lyase 활성 10.1 unit/g-soluble protein에서 ApCpnA와의 공발현 때는 83.1 unit/g-soluble protein, ApCpnB와의 공발현 때는 100.3 unit/g-soluble protein으로 가용성 및 활성형으로의 alginate lyase 생산이 8-10배 크게 향상되었다. 다양한 배양 조건들의 조사를 통해 alginate lyase 최대 생산을 위한 조건은 다음과 같았다: 1.0 mM IPTG, 25℃ 배양 온도(IPTG 유도 후), ApCpnB 공발현 파트너. 이러한 공발현 시스템은 대장균에서 기능적으로 또한 의학적으로 중요한 재조합 단백질의 산업적 생산에 크게 유용하게 사용될 것이다.