보 문

Exploration and functional expression of homologous lipases of *Candida antarctica* lipase B

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Candida antarctica lipase B의 상동체 효소 탐색과 발현

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ABSTRACT: *Candida* (also known as *Pseudozyma*) *antarctica* lipase B (CAL-B) has been intensely studied in academic and industrial fields. However, the research related to its homologous enzymes has been rarely reported. In the current investigation, protein sequence similarity search of CAL-B has been conducted and six homologous protein sequences were identified. After the syntheses of their codon-optimized genes, the synthetic genes have been cloned into a periplasmic expression vector to express in *Escherichia coli*. Among six homologous sequences, four sequences were successfully expressed in *E. coli*. The hydrolytic activities of the expressed proteins towards 4-nitrophenyl acetate and 4-nitrophenyl butyrate were measured and compared with those of CAL-B to identify whether the expressed proteins work as a hydrolase. It has been revealed that the expressed proteins can hydrolyze the substrates and the specific activities were determined as $(1.3-30) \times 10^{-2} \, \mu mol/min/mg$, which are lower than those of CAL-B. Among these homologous enzymes, *Pseudozyma hubeiensis* SY62 exhibits the comparable enantioselectivity to that of CAL-B towards the hydrolysis of (\pm) -1-phenylethyl acetate.

Key words: heterologous expression, homology, lipase

Lipases (E.C. 3.1.1.3), which catalyze the hydrolysis of triglycerides in nature, are one of the most widely used enzymes in academic and industrial fields (Bornscheuer and Kazlauskas, 2006; Wu *et al.*, 2013). Lipases structurally belong to an α/β -hydrolase fold family, and possess the catalytic triad composed of Ser-His-Asp/Glu. The amino acids of the catalytic triad are connected by hydrogen bonding to increase the nucleophilicity of the catalytic serine. The catalytic serine works as a nucleophile and attacks the carbonyl carbon atom of an ester substrate during catalysis.

Many lipases exhibit high stereoselectivity as well as chemoand thermo-stability. Especially, *Candida antarctica* lipase B (CAL-B, also called as *Pseudozyma antarctica* lipase B; Boekhout, 1995), one of the fungal lipases, exhibits extraordinary broad enantioselectivity towards a wide range of chiral secondary alcohols and high stability in various organic solvents including alkanes, ethers, and acetonitrile. Such properties make CAL-B suitable in the organic synthetic applications. Hence, researchers have employed CAL-B in the preparation of a variety of chiral building blocks (Rotticci, 2003; Bornscheuer and Kazlauskas, 2006). For instance, it has been demonstrated that CAL-B catalyzes kinetic resolution of racemic alcohols and amines, desymmetrization of diols and diacetates, and polymerization of lactones (Schmid and Verger, 1998; Gross *et al.*, 2001; van der Mee *et al.*, 2006; Takwa *et al.*, 2011; Andualema and Gessesse, 2012).

Although CAL-B is classified as one of the lipases (Ollis *et al.*, 1992) and also structurally belongs to α/β -hydrolase fold family, CAL-B exhibits distinct characteristics compared to

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other lipases. For instance, the kinetics of typical lipases exhibit an interfacial activation behavior (Verger, 1997), while CAL-B does not follow the interfacial activation mechanism (Martinelle et al., 1995). The interfacial behavior of lipases is structurally understood by the presence of a helical segment, which is referred to as a lid, covering the active site (Grochulski et al., 1993). Before a hydrophobic substrate approaching, the lid is covering the active site and the lipase remains as a closed structure. As a hydrophobic substrate approaches to a lipase, the lid opens up and the lipase is converted to the open structure. Both crystal structures for the closed and open forms of many lipases have been reported (Brady et al., 1990; Brzozowski et al., 1991; Grochulski et al., 1994). Typical lipases exhibit low activity at a low concentration of a hydrophobic substrate because of their closed structures. According to the increase of the concentration of the hydrophobic substrate, the activity of lipase rapidly increases at a particular point. This kinetic behavior of lipase is known as an interfacial activation. Interestingly, CAL-B does not exhibit such interfacial activity and a crystal structure of the closed form of CAL-B has not been reported (Martinelle et al., 1995; Uppenberg et al., 1995). Such distinctive characteristics of CAL-B probably suggest that CAL-B can accept a broad range of compounds including hydrophilic compounds as a substrate.

Although CAL-B has been commercialized and utilized in many researches, the homologous lipases of CAL-B have not been studied. Finding and studying homologous enzymes of interest will likely provide not only an evolutionary knowledge but also an opportunity of creating a new enzyme by gene shuffling. Gene shuffling is one of the strong tools to produce diversity of an enzyme and then to create a new enzyme through directed evolution (Crameri *et al.*, 1998; Carbone and Arnold, 2007). This paper describes the discovery of the homologous enzymes of CAL-B by comparison of protein sequences in a pool of protein sequences and the functional expression of the homologous enzymes in *E. coli*.

Materials and Methods

Chemicals, buffers, and lysozyme were purchased from Sigma-Aldrich Korea. Restriction enzymes and pfu DNA polymerase were purchased from Elpisbiotech. The Ni-NTA agarose resin and antibodies were purchased from Qiagen Korea Ltd. and Abcam, respectively. A centrifugal device (Amicon Ultra-15) was purchased from Millipore Korea. DNA oligomers were obtained from Sigma-Proligo. The synthetic genes were prepared by GenScript USA. DNA sequencing was performed by Solgent Co.

Exploration of homologous sequences of CAL-B by Protein BLAST

The protein sequence of CAL-B was put into Protein BLAST (http://blast.ncbi.nlm.nih.gov) and homologous sequences were explored with default parameters. Six sequences with more than 50% sequence identity were selected. The corresponding DNA sequences were obtained from GenBank (http://www.ncbi.nlm. nih.gov/nuccore). After the codons of the nucleotide sequences of the homologous proteins were optimized for *E. coli*, the codon-optimized genes were synthesized by GenScript and used as the template DNAs for PCR amplifications.

Construction of the expression vectors of the homologous lipase genes of CAL-B

A pair of gene specific primers for each gene containing restriction sites for NcoI and SalI were used to introduce the restriction sites in the genes and the amplified genes by PCR were cloned into pBAD/gIIIa vector. The following primers were used for PCR: 5'-AGCACCATGGCTCTGCCGCA-3' and 5'-GATGGTCGACGAAGCGCGTAACGATACCG-3' for CCF54401; 5'-AGCACCATGGCTCTGCCGTCA-3' and 5'-GATGGTCGACCAGGATAACGCCAGAACAGGT-3' for CBQ 70828; 5'-AGCACCATGGCTCTGCCGATTG-3' and 5'-GAT GGTCGACGAAACCCGTATTAACGCCGC-3' for GAC93661; 5'-AGCACCATGGCTCTGCCGAGT-3' and 5'-GATGGTCG ACGAACGTGGTAAAACCAGAACAGG-3' for EST06015; 5'-AGCACCATGGCTGCCAGCTCT-3' and 5'-GATGGTCG ACATTGATCGTACCGCTACAGGTTT-3' for XP 011387292; 5'-AGCACCATGGCTCTGCCGCG-3' and 5'-GATGGTCGA CCATGATAACGCCGCTACACG-3' for GAC96817. The reaction mixture for PCR contains the synthesized gene (10 ng), dNTP (0.2 mM), a pair of primers (each 0.4 µM), and pfu polymerase (1 μ l of 5 unit/ μ l) in a standard pfu reaction buffer $(50 \,\mu l)$. The PCR was carried out using the following conditions: initial denaturation at 95°C for 30 sec, followed by twenty five cycles (95°C for 30 sec, 57°C for 30 sec, and 72°C for 2 min), and a final extension at 72°C for 7 min. After digestion of the PCR products with *NcoI* and *SalI*, the digested products were ligated with the pBAD/gIIIa restricted by the same enzymes. The constructed expression vectors were transformed into Top 10 *E. coli* competent cells.

Expression and purification of homologous lipases of CAL-B

Overnight culture of Top10 (2.5 ml) was inoculated to LB medium (250 ml; ampicillin, 100 µg/ml) and incubated at 37°C for 3 h. A sterile L-arabinose solution (1.25 ml; 4% w/v) was added to induce protein expression and then incubated at 20°C for 20 h. The cells were harvested by centrifugation (10 min, 10,000 rpm, 4°C). The cell pellet (~0.4 g) was resuspended in a lysis buffer (2.5 ml; NaH₂PO₄, 50 mM; NaCl, 300 mM; imidazole, 10 mM; pH 8.0) containing lysozyme (1 mg/ml). After incubation on ice for 45 min, the suspension was frozen and thawed. The viscous lysate was drawn through a sterile 20-gauge syringe needle several times and centrifuged (20 min, 10,000 rpm, 4°C). Ni-NTA agarose resin (1 ml of 50% slurry) was added to the cleared lysate (2.5 ml) and the mixture was shaken at 4°C for 1 h. The lysate-Ni-NTA mixture was loaded on a Poly-Prep column (Bio-Rad), drained, and then washed three times with a wash buffer (2 ml; NaH₂PO₄, 50 mM; NaCl, 300 mM; imidazole, 20 mM; pH 8.0). The proteins were eluted from the column with an elution buffer (2 ml; NaH₂PO₄, 50 mM; NaCl, 300 mM; imidazole, 250 mM; pH 8.0). Eluate (2 ml) was exchanged from the elution buffer to a BES buffer (5 mM, pH 7.2) using a centrifugal device.

Determination of the amount of functional enzymes

The amount of purified enzymes was determined by Bio-Rad Protein Assay kit (Bio-Rad) according to the manufacturer's instructions.

Measurement of the hydrolytic activity towards the hydrolysis of 4-nitrophenyl acetate and 4-nitrophenyl butyrate

The assay solution was prepared by mixing 4-nitrophenyl

acetate or 4-nitrophenyl butyrate (20 μ l of 200 mM in acetonitrile), acetonitrile (870 μ l), and a BES buffer (5 mM, pH 7.2, 11,110 μ l). After mixing the assay solution (100 μ l) and the enzyme solution (5 μ l), the absorbance change was measured at 404 nm for 10 min by a microplate reader. The specific activity was calculated according to the method of Janes *et al.* (1999).

Results

Exploration of homologous sequences of CAL-B and heterologous expression in *E. coli*

The amino-acid sequence of a crystal structure (pdb code: 1TCB) of CAL-B was used for the blast search. Searching the sequence revealed that six protein sequences possess above 50% identity: CCF54401, 78%; CBO70828, 75%; GAC93661, 74%; EST06015, 72%; XP_011387292, 71%; GAC96817, 67% (Fig. 1). These six sequences were selected for the current investigation because a structure alignment from the FSSP (families of structurally similar proteins) database revealed that enzymes with above 50% sequence identity well conserve their functions (Bateman et al., 2002). As CAL-B is one of the fungal proteins, all identified sequences were originated from fungi such as Ustilago hordei, Sporisorium reilianum SRZ2, Pseudozyma hubeiensis SY62, Pseudozyma brasiliensis GHG 001, and Ustilago maydis 521 (Table 1). The homologous sequences possess the identical amino acids corresponding to the amino acids of the catalytic triad and oxyanion hole of CAL-B (i.e. Ser-His-Asp for the catalytic triad, and Thr and Gln for the oxyanion hole). In addition, the identified sequences contain a sequence for a potential signal peptide and the sequences were removed during gene syntheses.

The sequences were cloned into an expression vector (pBAD/ gIIIa) using a pair of sequence-specific primers containing the *NcoI* and *SalI* restriction sites for each gene. The encoded proteins contain additional alanine at the C-terminal, and valine, aspartate and 6×His at the N-terminal. After addition of L-arabinose for the induction of the protein expression, the culture was incubated at 20°C for 20 h. The expression of the homologous proteins were confirmed with an anti-6×His antibody by Western blotting analyses (Fig. 2). Among six homologous genes, only four homologous proteins were

1TCB_A CCF54401 CBQ70828 GAC93661 EST06015 XP_757569 GAC96817	1 1 1 1 1	LPSGSDPAFSQPKSVLDAGLTCQGASPSSVSKPILLVPGTGTTGPQSFD MKLSMIVSLLTLAAS-AVTASPLVRRLPQGFHPPFSTPKSLLDAGLTCQNGSPSSQSKPILLVPGTGATGKQSFD MKFLTALTVLASCSA-LASATPLVKRLPSGSDPAYTLSKAULDSVLAC0NGSPSSQSKPILLVPGTGTGPQSFD [102] MKFTSTVTALAALVC-AASATPLVKRLPIGPDPAFSVPQSQLASVLECQNGSPSSQSNPILLVPGTGVGGQSFD MKFASIITALVALASSAVVASPLVKRLPSGSDPALSTPQAVLDAGLFCQNGSPSSQNPILLVPGTGVDGRQNFD MKTSSVISALVTAS-IIRAAPLASSDPAFSTPKATLDAGLECQTGSPSSQTKPILLVPGTGVDGQQSFA MKT1PFVSALIVVLS-VVTATPLLSLPSDPAFSTPKATLDAGLECQTGSPSRQTRPILLVPGTGVDGQQSFA	49 74 176 75 69 74
1TCB_A	50	SNW I PLSTQLGYTPCW I SPPPFMLNDTQVNTEYWVNA I TALYAGSGNNKLPVLTWSQGGLVAQWGLTFFPS I RSKVDRLM	129
CCF54401	75	SNW I PLSARLGYNPCW I SPPPFMLNDTQVNAEY I VNAVN I LYAGSRSKKVPVLTWSQGGLAAQWALTFFPS I RSKVDRLM	154
CBQ70828	75	SNW I PLSTQLGYSPCW SPPPFMLNDTQVNAEY I VNAVKLVSASGA-KVPVLTWSQGGLAAQWALTFFPS I RTQVDRLM	153
GAC93661	177	SNW I PLSSQLGYSPCW SPPPFMLNDSQVNAEY I VNAVKLVAGSGSKKVPVLTWSQGGLATQWALTFFPS I RSKVDRFM	256
EST06015	76	SNW I PLSTMLGYSPCWVSPPPFMLNDSQ I NAEYVVNAVRRLYAGSGSTKLPVLTWSQGGLATQWALTFFPS I RNAVDRFM	155
XP_757569	70	SNW I PLSTMLGYSPCWI SPPPFMLNDSQ I NAEYVVNAVRRLYAGSGSKKVPVLTWSQGGLATQWALTFFPS I RNAVDRLM	149
GAC96817	75	SNW I PLSTLLGYSPCWI SPPFFMLNDSQ I NSEY I VNAVHTLYSGSGG-KVPVVTWSQGGLAVQWALTFFPS I RGKVDRLM	153
1TCB_A	130	AFAPDYKGTVLAGPLDALAVSAPSVWQQTTGSALTTALRNAGGLTQIVPTTNLYSATDEIVQPQVSNSPLDSSYLFNg-K	208
CCF54401	155	AFAPDYKGTIEAGLLNAVGLSSQSVWQQTAGSAFVTALQNAGGLNQIVPTTNLYSATDEIVQPQITNSPLDSSHLFN-AK	233
CBQ70828	154	AFAPDYKGTVLAAFLTTPGLASGSVWQQAQSALTTALANAGGLTKIVPTTNLYSATDEIVQPQTFNGPLDSGYLNGGAK	233
GAC93661	257	AFAPDYKGTVEAVFLTVPGLASQSVWQQAQSAYLTALQNAGGLTKIVPTTNLYSVFDDIVQPQVSNSPLDSSYLFN-AK	335
EST06015	156	AFVPDYKGTVNAYPLTATDTASPSIWQQPAQSAYLTALQNAGGLTKIVPTTNLYSVFDDIVQPQVSNSPLDSSYLFN-AK	234
XP_757569	150	AFVPDYKGTVNAYPLTATDTASPSIWQQPAQSAFTTALPNAGGLNKIVPTTNLYSVFDEIVQPQITNSPLDSSYLFN-AK	228
GAC96817	154	AFAPDYKGTVEAVFLAYGLASQSVWQQQAQSAFVTALKNAGGLTSFVPTTNLYSVFDEIVQPQVSNSPLDSSYLFN-AK	232
1TCB_A CCF54401 CBQ70828 GAC93661 EST06015 XP_757569 GAC96817	209 234 234 336 235 229 233	NVQAQAVCGPLFVIDHAQSLTSQFSYVVGRSALRSTTGQARSADYGITDCNPLPANDLTPEQKVAAAALLAPAAAAIVAG NIQAQSVCGPLFVIDHAQSLTSQFSYVVGRSALASPTGQAQSKDYRSSDCNPLPADPLSPQNKKEASALLLVAGANLVAG NIQAQSVCGPLFVVDHAQTLTSQFSFVVGRSALRSTTGQAQSSDYGVTDCNPLPADSLTPDQKLRAEGLLLVAGANVAAG NIQAQTVCGPAFLIDHAQSLTSQFSYIVGRSALASGTGEAQSSDYSITDCNPLPADPLTPQQKAESSTLLLVAGANVAAG NIQQQTVCQGFFVADHAQSLVNQFAYVVGRSALRSSTGQAQSSDYSITDCNPLPADPLTPQQKAESSTLLLVAGANVAAG NIQQQTVCQGFFVIDHAQSLTSQFSYVVGRSALRSSTGQAQSSDYSIADCNPLPADPLTPQQKAESSTLLLVAGANVAAG NIQQQTVCGGFFVIDHAQSLTSQFSYVVGRSALSSTGQAQSSDYSIADCNPLPADPLTPQQKAESSTLLLVAGANVAAG NIQQQTVCGGFFVIDHAQSLTSQFSWVGRSALGSKTGEAVSGEFGQDCNATAAEPLSAQQKSDAGGLLVAGANLAG	288 313 313 415 314 308 312
1TCB_A	289	PKQNCEPDLMPYARPFAVGKRTCSGIVTP- 317	
CCF54401	314	PKQNCEPDLKPYARRFAIGKRTCSGIVTRf 343	
CBQ70828	314	PKQNCEPDLMPYARQVAVGKRTCSGVIL 341	
GAC93661	416	PKQNCEPDLKPYARRFAIGKKTCSGVITGf 445	
EST06015	315	PKTNCEPDLKPYARPFAPGSQTCSGFTTF- 343	
XP_757569	309	PKQNCEPDLKPYARQFAVGKKTCSGTIN 336	
GAC96817	313	PKVRCEPDLKPYARQFAVGKKTCSGVIN 340	

Fig. 1. Alignment of CAL-B with its homologous proteins. The yellow highlighted sequences are assumed as a signal sequence and the sequences were removed in syntheses of the genes. The red colored letters are considered as the catalytic triad and the blue ones as the oxyanion holes.

Origin (organism)	Locus number (NCBI protein)	Sequence identity	Number of amino acids ^a	Molecular weight (Da)
Ustilago hordei	CCF54401	78%	318	33,782
Sporisorium reilianum SRZ2	CBQ70828	75%	316	33,016
Pseudozyma hubeiensis SY62	GAC93661	74%	318	33,666
Pseudozyma brasiliensis GHG001	EST06015	72%	317	33,588
Ustilago maydis 521	XP_011387292	71%	314	32,910
Pseudozyma hubeiensis SY62	GAC96817	67%	316	33,379

Table 1. Identified homologous sequences of CAL-B

^a The signal peptides were removed.



- 4. Pseudozyma brasiliensis GHG001
- 5. Ustilago maydis 521
- 6. Pseudozyma hubeiensis SY62

Fig. 2. Western blotting analyses of the expressed proteins. The corresponding bands (lanes 1 and 6) to the proteins from U. hordei and P. hubeiensis SY62 were not or poorly detected, respectively.

overexpressed. The proteins from Ustilago hordei and Pseudozyma hubeiensis SY62 were not or poorly expressed, respectively. Thus, the rest four proteins were used for further study. The four expressed proteins were purified using Ni-NTA agarose resins (Fig. 3) and the protein amount was determined by the Bradford method after the buffer was exchanged from the elution buffer to a 5 mM BES buffer (pH 7.2).

Measurement of hydrolytic activities of the homologous proteins and CAL-B

Hydrolytic activities of the expressed proteins were determined using 4-nitrophenyl acetate and 4-nitrophenyl



Fig. 3. SDS-PAGE and Native PAGE analyses of the recombinant homologous proteins and CAL-B expressed in *E. coli*. (A) protein from *Sporisorium reilianum* SRZ2, (B) protein from *Pseudozyma hubeiensis* SY62, (C) protein from *Pseudozyma brasiliensis* GHG00, (D) protein from *Ustilago maydis* 521, (E) lipase B from *Candida antarctica*, (F) Native PAGE analyses (Lanes: 1, protein from *Sporisorium reilianum* SRZ2; 2, protein from *Pseudozyma hubeiensis* SY62; 3, protein from *Pseudozyma brasiliensis* GHG00; 4, protein from *Ustilago maydis* 521; 5, lipase B from *Candida antarctica*). SDS-PAGE and Native PAGE were performed on a 12% polyacrylamide gel and stained using the Coomassie brilliant blue. M, molecular weight marker; L, lysis buffer fraction; F, flow-through fraction; W, wash buffer fraction; E, elution buffer fraction.

butyrate as a substrate. 4-Nitrophenyl acetate is one of the typical esterase substrates and CAL-B also readily hydrolyzes

it. 4-Nitrophenyl butyrate contains a longer carbon chain than 4-nitrophenyl acetate and thus is better substrate for lipases (Jung and Park, 2008). The hydrolysis product of 4-nitrophenyl acetate and 4-nitrophenyl butyrate is 4-nitrophenol, which can be detected by a UV/Vis spectrophotometer. Table 2 shows the specific activities of the expressed proteins and CAL-B. All expressed proteins can hydrolyze both substrates and thus they can be considered as a hydrolase. Each enzyme exhibited different relative specific activities. Enzyme from S. reilianum SRZ2 possesses similar specific activities towards both substrates. Enzyme from P. hubeiensis SY62 exhibited three times higher activity towards 4-nitrophenyl butyrate, while enzyme from P. brasiliensis GHG001 and enzyme from U. maydis 521 displayed higher activity towards 4-nitrophenyl acetate by a factor of about 5. However, all homologous enzymes exhibited lower specific activities than those of CAL-B towards both substrates. The enantiomeric preferences of the homologous enzymes towards (±)-1-phenylethyl acetate were measured. CAL-B possesses high enantioselectivity towards (±)-1-phenylethyl acetate (E > 200). Among four homologous enzymes, enzyme from P. hubeiensis SY62 exhibits the comparable enantioselectivity (E > 200) towards (±)-1-phenylethyl acetate and the others showed lower enantioselectivity (E =21-80).

Table 2. Hydrolytic activity and selectivity of CAL-B and its homologous enzymes^a



^a Reaction condition: see the 'Materials and Methods' section.

^b Errors are standard deviations for at least three measurements

Discussion

Besides finding new enzymes from nature, it has become an important approach to create functionally improved mutant enzymes for industrial use. For the creation of mutant enzymes of interest, two approaches are mainly concerned: rational design and directed evolution (Bornscheuer and Pohl, 2001). Rational design of enzymes is based on the structural and mechanistic knowledges. On the other hand, directed evolution, which does not require the detailed mechanistic knowledges, uses generating random mutants and screening the mutants to find a functionally improved one. Researchers have used error-prone PCR, mutator strain, or gene shuffling to generate random mutants (Neylon, 2004). Among these approaches, gene shuffling is considered as one of the strongest tools to increase the genetic diversity. However, a homologous family is required to use gene shuffling to keep a conserved fold of an enzyme (Carbone and Arnold, 2007). Since the expression systems of CAL-B have been developed in low eukaryotes and in E. coli (Hoegh et al., 1995; Rotticci-Mulder et al., 2001; Zhang et al., 2003; Chodorge et al., 2005; Blank et al., 2006; Liu et al., 2006; Jung and Park, 2008), a directed evolution approach based on error prone PCR has been successfully employed to improve the functional properties of CAL-B (Zhang et al., 2003; Wu et al., 2013). However, the employment of gene shuffling on generating random mutants of CAL-B is still worth to be exploited. In the present study, six homologous hydrolases of CAL-B were identified from a protein sequence pool and four of them were successfully expressed in E. coli. Although the enzymes are originated from fungi, functional expression in E. coli benefits researchers from decrease of expression time and culture cost (Jung and Park, 2008). Identification of functionally active homologous enzymes of CAL-B could provide a useful source in direct evolution based on gene shuffling. In addition, the detailed sequence comparison of CAL-B with the homologous enzymes may provide a useful information for rational design to improve the property of CAL-B.

적 요

Candida (*Pseudozyma*로도 알려짐) *antarctica* lipase B (CAL-B)는 학문적으로 그리고 산업적으로 많이 활용되고 있

다. CAL-B 자체에 대한 연구는 많이 진행되어온 반면, CAL-B 상동체에 관한 연구는 그리 알려진 바가 없다. 본 연구에서는 단백질 유사성 검색을 통해서 CAL-B의 상동체 탐색을 수행 하였고, 6종의 단백질 서열을 찾았다. 해당하는 유전자들을 대 장균에 대한 코돈 최적화를 수행하였고, 이를 바탕으로 유전자 합성을 진행하였다. 이들 유전자를 대장균 발현용 벡터에 클로 닝한 후, 대장균 내에서 단백질 발현을 시도하여 이들 중4종의 단백질이 성공적으로 발현되었다. 이들 단백질들이 가수분해 효소로서의 활성이 있는지 확인하기 위해서, 4-nitrophenyl acetate와 4-nitrophenyl butyrate를 반응기질로 하여 가수분해 반응성을 확인하였다. 이들 단백질들의 비활성(specific activity) 값은 (1.3-30) × 10⁻² µmol/min/mg로 측정되었고, 이는 CAL-B 의 비활성 수치보다는 다소 낮은 값에 해당하였다. (±)-1phenylethyl acetate의 가수분해 반응에 대한 입체선택성은 이 들상동체 효소들 중에서 Pseudozyma hubeiensis SY62에서 유 래된 효소만이 CAL-B의 입체선택성과 유사함이 확인되었다.

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