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## Cloning and Nucleotide Sequence of $\gamma$ -Polyglutamate Production Stimulating Factor on *Bacillus subtilis* (*natto*) Plasmid, pUH1

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The gene coding for  $\gamma$ -polyglutamate production stimulating factor (*psf*) from Bacillus *subtilis*(*natto*)plasmidpUH1 was cloned and sequenced in *Bacillus subtilis* host. The activity of y-glutamyltranspeptidase ( $\gamma$ -GTP) in *B. subtilis* host, which was introduced the *psf* gene was relatively high level, while its activity in *E. coli* host could not be detected. The nucleotide sequence of *psf* gene was determined and an open reading frame encoding a polypeptide composed of 420 amino acid residues (Mr, 49, 356) was identified. The putative -35 and -10 sequences, TTCAAA and TATTAT, were observed as the consensus sequence for the promoter recognized by the  $\sigma^{43}$  RNA polymerase of *B. subtilis*, and the ribosome binding site, whose sequence was AACGAG, was complementary to the binding sequence of *B. subtilis* 16SrRNA except for one base. The amino acid sequence of *psf* with the segment of putative protein C403 of staphylococcal plasmid pE194 indicates homology, whereas that with *E. coli* and mammalian  $\gamma$ -GTPs does not show any similarity at all.

#### INTRODUCTION

"Natto" is a Japanese traditional fermented food, manufactured by growing **Bacillus subtilis** (*natto*) on steamed soybeans. It is an adhesive, and consists of polysaccharide (levan-form fructan) and y-polyglutamate ( $\gamma$ -PGA). The adhesive material is mainly composed of  $\gamma$ -PGA containing D-and L-glutamate in varying proportions (Fujii, 1963). A plausible mechanism of the biosynthetic pathway of  $\gamma$ -PGA has been proposed by Thorne et al. (1955) for one of the  $\gamma$ -PGA-producing strains (*B. licheniformis* ATCC9945A). However, since it is known that various strains differ a great deal in the basic requirements for  $\gamma$ -PGA synthesis as a capsule, it is to be assumed that there is more than one biosynthetic pathway. We reported that a 5.8-kb plasmid designated pUH1, which is responsible for y-PGA synthesis, is distributed widely in *B. subtilis* (*natto*) strains isolated from a non-salty fermented soybean food, natto (Hara et al., 1983; 1993; unpublished results).

y-Glutamyltranspeptidase ( $\gamma$ -GTP, EC2.3.2.2) catalyzes the transfer of the  $\gamma$ -glutamyl residue from y-glutamyl compounds, such as glutathione, to amino acids and peptides, and the hydrolysis of y-glutamyl compounds (Tate and Meister, 1981), but its physiological role still remains controversial. The cDNAs of rat renal (Laperche *et al.*, 1986) and human hepatic (Sakamuro et *al.*, 1988) $\gamma$ -GTPs were cloned, and the nucleotide sequences were determined. Recently, Suzuki et *al.* (1989) reported the

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nucleotide sequence of *Escheyichia coli K-12* y-GTP. However, the homology percent of amino acid sequence between the *E. coli* and mammalian  $\gamma$ -GTPs was quite low.

We have succeeded in expressing  $\gamma$ -GTP activity in *B. subtilis* host strains, which were introduced a  $\gamma$ -PGA production stimulating factor (*psf*) gene. The *psf* gene might be responsible for  $\gamma$ -PGA synthesis in B. *subtilis* (*natto*). In the present work, we have sequenced the *psf* gene and compared with the *E. coli* and mammalian  $\gamma$ -GTPs.

### MATERIALS AND METHODS

Bacterial strains and **plasmids** 

Escheyichia coli JM101 (supE44, thi,  $\Delta$ (lac-proAB) [F', proAB, lacIqZ $\Delta$ M15, traD36], mcrA<sup>+</sup>) and Bacillus subtilis MI112 (ayg-15, leuB8, thy-5, recE4) were used as hosts for cloning and nucleotide sequencing. Plasmid pUH1 has been previously described (Hara *et al.*, 1983). Plasmid pUB110 was used as a vector for B. subtilis host, and plasmids pATE1 (Hara *et al.*, 1991) was for E. coli.

Media

LB broth and Panassary broth (Difco) for both *B. subtilis* and *E. coli*, Spizizen minimal medium for *B. subtilis*, and M9 minimal medium for *E. coli* were the same as described previously (Hara *et al.*, 1991). The cells carrying Km' plasmids were grown in AA medium (Tanaka and Sakaguchi, 1978) containing kanamycin (50  $\mu g/ml$ ).

DNA manipulations

The plasmid pUH1 from *B. subtilis* (*natto*) strain Asahikawa and its derived plasmids from Km' transformants were prepared and purified as previously described (Hara *et al.*, 1983) was digested with *Bst*EII, and then the ends were filled in with Klenow fragment to generate blunt ends. The DNA fragments were ligated with *Bam*HI liker, inserted at the *Bam*HI site of pUB110 with T4 ligase and then added to *B. subtilis* MI112 protoplasts. Restriction enzymes, T4 DNA ligase and bacterial alkaline phosphatase were purchased from Takara Shuzo Co., Ltd. (Kyoto, Japan), and used as recommended by the manufacture. Degradation of DNA with exonuclease Ba131 (Takara Shuzo Co., Ltd.) followed by procedure of Legerski *et al.* (1978).

Transformation procedure

*E. coli* JM101 was transformed by the method of Morrison (1977), and *B. subtilis MI112* was transformed by using protoplasted cells (Hara *et al.*, 1991).

Assessment of y-glutamyltranspeptidase assay

After cultivation at time indicated in the medium described by Fujii (1963), cells were harvested, suspended in 0.9% NaCl, and disrupted with a sonic oscillator (Branson Sonifer 185). The sonicated cells were centrifuged and the supernatant solution was used as intracellular enzyme preparation. The extracellular enzyme activity was measured in the culture fluid removed the cells by centrifugation.  $\gamma$ -GTP activity was assayed as previously described (Aumayr *et al.*, 1981). One unit of  $\gamma$ -GTP activity is defined as the amount of enzyme which liberates  $1\mu$ mol of a-naphthylamine per min at  $37^{\circ}$ C.

DNA sequencing

DNA fragments were subcloned into plasmids pUC18 and pUC19, and then, if need,

the Bal31-deleted derivatives were obtained by the stepwise deletion method of McCutchan et **al.** (1984). DNA sequencing was carried out by means of the dideoxy chain termination method (Sanger et **al.**, 1977) with Sequenase Version 2.0 DNA Sequencing Kit (United States Biochemical Corporation, Cleveland, Ohio). Nucleotide and amino acid sequences were analyzed by the Hitachi DNASIS system.

#### Chemicals

Restriction enzymes, **T4 DNA** ligase and bacterial alkaline phosphatase were purchased from Takara Shuzo Co., Ltd., and used as recommended by the manufacture. Degradation of DNA with exonuclease Ba131 (Takara Shuzo Co., Ltd.) followed by procedure of Legerski et **al.** (1978).

#### **RESULTS AND DISCUSSION**

#### Cloning of *psf* gene in B. subtilis

As reported previously (Hara et **al.**, 1981), the *psf* gene is encoded on an endogenous plasmid, pUH1 from *B.* **subtilis** (*natto*). We constructed a set of derived plasmids using pUB110 as a vector, in which a series of pUH1 fragments were inserted. Figure 1 shows the subclones of *B.* **subtilis** (*natto*) plasmid pUH1 and their expression of *psf* in *B.* **subtilis** MI112. The *psf* activity was detected in several Km' transformants, which carried plasmid pPB1 (column BsL in Fig. 1) with molecular size of 8.2 kb, including 3.7-kb BstEII fragment of pUH1. To define the boundaries of a functional unit of inserted DNA, a 3.7-kb BstEII fragment of pUH1 was digested with selected restriction endonucleases to obtain a set of overlapping DNA fragments. The plasmid DNA preparations containing each generated fragments were tested for the enzyme activity in *B.* **subtilis**. The deletion experiments revealed that a 1.7-kb



Fig. 1. Subclones of *B. subtilis* (*natto*) plasmid pUH1 and their expression of  $\gamma$ -GTP in *B. subtilis M1112*.

The open *arrow* in the BstEII fragment indicates an open reading frame found in the sequence data (see Fig. 3). The activity of  $\gamma$ -GTP expressed was assayed as previously described (Aumayr *et al.*, 1981). Abbreviations for restriction sites are : Ba, *Bam*HI; Bs, BstEII; **H**, Hind111; T, *Taq*I.

Strain		<i>psf</i>	$\gamma$ -GTP activity"		
			Intracellular	Extracellular	
B. subtilis	Asahikawa	+	91.8	423	
B. subtilis	MI112		10.1	22.5	
	MI112 (pBS)	+	93.5	91.5	
E. coli	coli JM1O1		8.3	10.6	
	JMIOI (pHN-1)	+	8.0	10.5	

**Table 1.**  $\gamma$ -GTP activities in *B. subtilis* and *E. coli* strains carrying *psf* gene.

<sup>a</sup> Enzyme activities were measured under the conditions described in Materials and Methods.

TaqI fragment (column T in Fig. 1) is necessary for the expression of *psf* gene in *B*. subtilis MI112.

#### Expression of *psf* gene in B. *subtilis* and *E. coli*

The presence of *psf* gene on plasmid pUH1 was confirmed by measuring the  $\gamma$ -GTP activity in strains carrying the *psf*. These results are summarized in Table 1, which includes the enzyme productivity of *B. subtilis (natto)* strain Asahikawa. The *B. subtilis MI112* (pBS) carrying recombinant plasmid pBS expressed very high  $\gamma$ -GTP activity intracellularly at 93.5 mU/ml in the PY culture medium growth at 37°C for 24 hr, while the strain MI112 (pBS) accumulated extracellularly  $\gamma$ -GTP at level of 91.5 mU/ml for 5 days. In order to compare the productivity of  $\gamma$ -GTP in *E. coli* with that in *B. subtilis, we* subcloned the 1.7-kb *TaqI* fragment of pUH1 into pATE1 constructed from pTL12 (Tanaka and Sakaguchi, 1982) (the recombinant plasmid obtained was designated pHN1). However, *E. coli* JMIOI carrying pHN1 did not express intracellularly the  $\gamma$ -GTP activity in LB broth grown at 30°C for 24hr. Also,  $\gamma$ -GTP activity could not detect extracellularly after even 5-days cultivation.

## Nucleotide sequence of pUH1 psf gene

Figure 2 shows the physical map and sequencing strategy of 1.7-kbTaqI fragment including the *psf* gene of *B. subtilis* (*natto*) plasmid pUH1. The nucleotide sequences of both strands of the 1.7-kb TaqI fragment were determined by the dideoxynucleotide chaintermination method (Sanger et al., 1977) after successive Ba131 exonuclease deletions (Davis et al., 1986). The nucleotide sequence was determined for both strands by using numerous restriction fragments to give enough overlapping regions. Figure 3 shows the nucleotide sequence of the 1.7-kbTaqI fragment. By examination of possible open reading frames (ORFs), we found only one large frame, which consists of 1,260 bp and encodes a protein molecule with 420 amino acids and an Mr of 49,356. The *psf* ORF was found to be preceded by a putative  $\sigma^{43}$  RNA polymerase promoter of B. subtilis. The 5' upstream region of psf gene contains a 5'-AACGAG-3' sequence (indicated as SD in Fig. 3) complementary to the 3' end of 16SrRNA (3'-OH-UCUUUC-CUCCAGUAG-5') of B. subtilis (McLaughlin et al., 1981) at nucleotides 295 to 300 for translation initiation. There is a 5'-TATTAT-3' sequence (-10 in Fig. 3) resembling a Pribnow box (Moran et al., 1982) at nucleotides 257 to 262, and at a site of 23 bp upstream of this -10 sequence, there is a 5'-TTCAAA-3' sequence resembling the -35 sequence (Moran et al., 1982) of B. subtilis gene. The observed distance (17 bp)



Fig. 2. Restriction map and sequencing strategy for the *psf* gene of *B. subtilis(natto)*plasmidpUH1.

The open arrow in the restriction map indicates an open reading frame found in the sequence data (see Fig. 3). Below the map, arrows indicate the direction and extent of sequencing either of subcloned restriction fragments (thin arrows) or of a series of deletion fragments prepared by Ba131 (thick arrows). Abbreviations for restriction sites are: B, *Bam*HI; Ha, *Hae*III; Hd, *Hind*III; M, *Msp*I; P, *Pvu*II; T, *Taq*I.

between the -10 and -35 sequences accords well with that observed generally in B. *subtilis* genes (17–18 bp) (Moran *et al.*, 1982).

#### Homology of PSF to other y-GTP genes

The amino acid sequence of the predicted protein PSF encodes on pUH1 was compared with a number of protein sequences registered in GenBank with used of the homology search system of GENAS (Kuhara et al., 1984). As shown in Fig. 4, sequence homologous to the pUH1 PSF were found to in putative protein C403 (420 residues ; Horinouchi and Weisblum, 1982) with 48,300 daltons encoded in staphylococcal plasmid pE194. Approximately, 51.2% amino acid homologies with pUH1 PSF was observed in a segment of 86 residues of C403, but percent match of both proteins, in its entirety, was quite low value such as 11.9%. The cDNAs of rat renal (Laperche et al., 1986) and human hepatic (Sakamuro et al. 1988), hepatoma (Goodspeed et al., 1989), and placental (Mevts et al., 1988)  $\gamma$ -GTPs were cloned, and the nucleotide sequences were determined. The mammalian  $\gamma$ -GTPs, whose amino acid sequences were essentially the same, are synthesized as single polypeptides and then processed into the large and small subunits (Matsuda and Katsunuma, 1984), but did not show any similarity at all with pUH1 PSF from B. subtilis (natto). Percent match of amino acid sequence of pUH1 PSF for human hepatic y-GTP was 8.4%. Recently, Suzuki et al. (1989) performed DNA sequencing of E.coli y-GTP, and they suggested that the E.coli y-GTP might be also processed posttranslationally. The homology of amino acid

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TaqI 50 TCGATGCGACAGCGAGAATGAGAGACGCACCAGCACCGCAGGTCGCACGTCCAAATTT 100 GCCATGGCATAATTTGGTGTGTGGTGCGTTACACCAAAGATAAACTTTGTGTTACCATAACCCCTATACAGTGGTCTGAATCGGGGGGTTTTT 200 250 (SD) (-10)ARAAGCAAAAATGAAAAGTATTATCGGGGACGGGGGGCGTCCTAAATTCTGATCTGGTGAACGAGGAACCGATTAGCTATTCAAAAGCGGTTC 350 400 ATGAAAAAATTGAGGGTCGGGTCAAACGGAAGGTCCGAGCGGATGCGTGTTTTGGTCAGCGAATTTTTGATCACGGCAAGTCCTGACTAT  ${\tt MetLysLysLeuArgValGlySerAsnGlyArgSerGluArgMetArgValLeuValSerGluPheLeuIleThrAlaSerProAspTyr}$ 450 <u>ATGAAAGGGCTGAGTGATGAGGAGCAGCGGCGCTATTTTGAAACAGCGGTTGATCATTTGAAAGAGAAATACAGCGCTGAÄÄACATGCTT</u>  ${\tt MetLysGlyLeuSerAspGluGluGlnArgArgTyrPheGluThrAlaValAspHisLeuLysGluLysTyrSerAlaGluAsnMetLeu}$ 550 TATGCCACAGTCCATATGGATGAAGCGACTCCCCCATATGCATGTTGGTATTGTACCGATCACGGAGGACGGCCGACTCTCTGCGAAAGAT TyrAlaThrValHisMetAspGluAlaThrProHisMetHisValGlyIleValProIleThrGluAspGlyArgLeuSerAlaLysAsp 00 650 TTTTTTAATGGCAAATTGAAGATGAAAGCCATTCAAGATGATTTTCATCGGCACATGGTTGAAAACGGTTTTGACCTGGTGCGCGGCGAA  $\label{eq:pheAsnGlyLysLeuLysMetLysAlaIleGlnAspAspPheHisArgHisMetValGluAsnGlyPheAspLeuValArgGlyGluBelleuVa$ 700 750  $\label{eq:prosergluLysLysHisGluAsnValHisGlnTyrLysIleAsnGlnArgGlnAlaGluLeuGluArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaLeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAlaCeuArgLeuAsnAlaGluIleAsnAlaGluAsnAlaAsnAlaGluAsnAlaAsnAlaAsnAlaGluAsnAlaAsn$ 800 850 AAGGAAAAGCAGAGAGAGAGAACTGGAAAAGCAAAACAAAGCTGTTCAAGCAGTTATAGAAGTGAAAAAAGAATCACTGACAAGCTAAGGCT LysGluLysGlnArgGluGluLeuGluLysGlnAsnLysAlaValGlnAlaVallleGluValLysLysGluSerLeuThrAlaLysAla 900 950 GAAGAGTTGAAAAATGCCGACTATTGAACATGAAAAAGCGTGGCTCAAAAAGGATAAAGTCATTGTGCCGGAGCGGGAACTCCATGCTTTG  ${\tt GluGluLeuLysMetProThrIleGluHisGluLysAlaTrpLeuLysLysAspLysVallleValProGluArgGluLeuHisAlaLeu}$ 1000 10 TACGCCTATGCGGAGCAGAAAACTAAAACGACAGCCGAGCTGGCGGGCAATTGAAGTCAGAAACGCAGGAAAAGGAGCGCTGGCAGTCT TyrAlaTyrAlaGluGlnLysThrLysThrSerAlaGluLeuAlaGlyGlnLeuLysSerGluThrGlnGluLysGluArgTrpGlnSer ATCCCCCCGCAGAAGCAGATCCGCCAGATGAAAAAGACCAACGGCTTCAGGAACTGCAGAGATAAGATCCATTCAGAAGTTGAAGCGTCCA  ${\tt lleAlaArgGlnLysGlnTleGlyGlnMetLysLysThrAsnGlyPheArgAsnCysArgValArgSerIleGlnLysLeuLysArgPro$ 1150 1200 AAAAGGAAATGCGCGCCAAGCTTGCAAAGGAATTTACGGAAGAGCAAGCGTGAGGATCTTCGGCAGGAAGTGAAAGAGGAGCTGACGGCT LysArgLysCysGlyAlaSerLeuGlnArgAsnLeuArgLysSerLysArgGluAspLeuArgGlnGluValLysGluGluLeuThrAla 1250 1300 TTACGAACGGAAAATAAGGAACTGCTAGCTGAAAAATAAAGTCTTGATCATTGAGAGGAATCGTGAAGGTGAGGAAAACTTAAAACTAAAA LeuArgThrGluAsnLysGluLeuLeuAlaGluAsnLysValLeuIleIleGluArgAsnArgGluAlaGluGluAsnLeuLysLeuLys 1350 1400 CAGGAACTTGATAAGAGAGACGGGCAGTATGCTGAGGTTTTGAGTTTCGCCCAGAAGCAGAACCAAACACTTGAAAAAGTGGCTGGAGAA  ${\tt GlnGluLeuAspLysArgAspGlyGlnTyrAlaGluValLeuSerPheAlaGlnLysGlnAsnGlnThrLeuGluLysValAlaGlyGlu}$ 1450 . AsnLysAlaLeuLysLysGluAsnLysThrLeuLysGluArgValAlaValLeuGluGluTrpLysAspLysMetValGlnTrpAlaLys  $\begin{array}{c} 00 \\ \textbf{1550} \\ \textbf{GAAGGTTACCAAAGATGCGGAAATTAGCGGCATCGTTTTTCAATACAGCAGGTATCGTCGAGAAGCTGCTAAATATAAGGACAATGAAT \\ \textbf{GAAGGTTACCAAAGATGCGGAAATTAGCGGCATCGTTTTTCAATACAGCAGGTATCGTCGAGAAGCTGCTAAATATAAGGACAATGAAT$  ${\tt GluArgLeuProLysMetArgLysLeuAlaAlaSerPhePheAsnThrAlaGlyIleValGluLysLeuLeuAsnIleArgThrMetAsnI$ 1600 1650 TGGAGCGATAAATGGTTTTCTTAAGCGGTGAAAACGGGCAAGTGAATTAGTCTTCCAAAATTGCCACCACTTTTTTTGTTTAGTGGCTGC TrpSerAspLysTrpPheSer\*\*\* 1700 TagT AATCATTTTTTGTGTCTCCAGTGATTCTCGTAAAGCTTCGA

Fig. 3. Nucleotide sequence and the deduced amino acid sequence of *psf* gene of *B. subtilis* (*natto*) plasmid pUH1.

The putative -35, -10 and SD sequences were indicated by underlines and the possible terminator sequences are shown by arrows. Nucleotides are numbered from the 5' end of TaqI site of the 1.7-kb DNA strand with the same polarity as mRNA. Amino acid sequences are shown below the coding frame.

		10	20	30	40	50
1	s. aureus <sub>PE194</sub> C403	MSHSILRVARVK	GSSNTNGIQRH	INQRKNKNYNN	KDINHEETYK	NYDLINA
2	pUH1 PSF					
3	PNKH PSF					
4	E. coli K-12 y-GTP	MIKPTFLRRVA	IAALLSGSCFS	AAAAPPAPPVS	SYGVEEDVFHP	VRAKQ
5	Human hepatic y-GTP	VKK	KLWLGLLAVVI	VLVIVGLCL	WLPSASKHPD	NHVYTR
	60 70	80	90	100	110	120
1	ONIKYKDKIDETIDENYSGKR	KIRSD-AIRHVD	GLVTSDKDFFI	DLSGEEIERE	FKDSLEFLEN	VEYGKEN
2	MKKLRGESN	GRSERMHVLVSE	FLITASPDYM	IGLSDEEQRRY	FETAVDHLKI	EKYSAEN
3	MKKLRGESN	GRSERMRVLVSE	FLITASPDYM	GLSDEEQRRY	EETAVDHLKE	IYYSAEN
4	GMVASVDATATOVGVDILKEG	GNAVDAAVAVGY	ALAVTHPOAG	VLGGGGFMLI-	-RSKNGNTTA	IDFR-EM
5	AAVAADAKOCSKIGRDALRDD	DSAVDAAIAALL	CVGLMNAHSM	TGGGLFLTI	NSTTRKAEVI	INAR-EV
	-					
	130 140	150	160	170	180	190
1	MY-LATVHLDE	RVPHMHFGFVPI	TEDGRLSAKE	LGNKKDFTQI	QDRFNEY	VNEKGYE
2	ML-YATVHMDE	ATPHMHVGIVPI	TEDGRLSAKD	FNGKLKMKA	QDDFHRH	IVENGED
3	ML-YATVHMDE	ATPHMHVGIVPI	TEDGRLSAKDI	FNGKLKMKA	QDDFHRHI	IVENGFD
4	APAKATRDMFLDDOGNPDSKK	SLTSHLASGTPO	TVAGFSLALD	YGTMPLNKV	QPAFKLARD	GFIVNDA
5	APRLAFATMF-NS-SE-OSOK	GG-LSVAVPO	EIRGYELAHO	RHGRLPWARL	<b>OPSIQLARQ</b>	GFPVGKG
-	<b>x</b> - <b>x</b> -		~			
	200 210	220	230	240	250	260
1	LERGTSKEVT-EREHKAMDOY	KKDTVFHKOELC	EVKDELOKAN	KOLOSGIEHM	STRPFDYEN	ERTGLFS
2	LVRGEPSEKKHENVHQYK-IN	QREPELERINAL	IALKEKÕREEI	LĒKÕNKAVQAV	JIEVKKESLT	AKAEELK
3	LVRGEPSEKKHENVHOYK-IN	OROAELERLNAE	IALKEKOREEL	LEKQNKAVQAV	/IEVKKESLT/	AKAEELK
4	LADDLKTYGSEVLPNHENSKA	ĨFŴKEGEPLKKO	DTLVQANLAK	SLEMIAENGPI	DEFYKGTIAE	QIAQEMQ
5	LAAALEN-KRTVIEOOPVLCE	VFCRDRKVLREC	ERLTLPQLAD	<b>FYETLAIEGA</b>	DAFYNGSLTA	QIVKDIQ
	270 280	290	300	310	320	330
1	GREETGRKILTADEFERLQET	ISSAERIVDDYE	NIKSTDYYTEI	QELKKRRESI	LKEVVNTWKE	<b>JYHEKSK</b>
2	MPTIEHEKAWLKKDKVIVPER	ELHALYAYAEQH	TKTAAELAGQ	-LKSETQE-KI	ERWQSIARQE	ADR-ADE
3	MPTIEHEKAWLKKDKVIVPER	ELHALYAYAEQH	TKTTAELAGQ	-LKSETQE-KI	ERWQSIARQE	ADR-ADE
4	KNGGLITKEDLAAYKAV-ERT	PISGDYGYQVYS	MPPPSSGGIH:	IVQILNILEN	DMKKYGFGS	ADAM-
5	AAGGIVTAEDLNNYRAELIEH	PLNISLGDAVLY	-MPSAPLSGP	VLALILNILK	<b>GYNFSRESVE</b> :	SPEQKGL
	340 350	360	370	380	390	400
1	EVNKLKRENDSLNEQLNVSEK	FQASTVTLYRA	RANFPGFEKG	FNRLKEKFFNI	OSKFERVGQFI	MDVVQDN
2	KDQRLQELQSRIHSEVEASKK	EMRRKLAKEFD-		QDLRQEVKEE]	LTTLRTENEE:	LSAENKV
3	KDQRLQELQSKIHSEVEASKK	EMRRKLAKEFD-		EDLRQEVKEEI	LTALRYENKE:	LLAENKV
4	-QIMAEAEKYAYADRSEYLGD	PDFVKVP-WV	ALTNKAYAKS	IADQIDINKAI	(PSSEIRPGK)	LAPYESN
5	TYHRIVEAFRFAYAKRT-LLG	DPKFVDVTEVVF	NMTSEFFAAQ	LRAQISDDTT	PISYYKPEF	YTP-DDG
	410 <i>420</i>	430	440	450	460	470
1	VQKVDRKREKQRTDDLEM					
2	LIIQRNSEAAESLKLKQELDK	RNGQYAEVLSFA	KKQNQTLEKV	AGENKALKKEN	WATLKERVAV	LEQWKDK
3	LIIERNREAEENLKLKQELDK	RDGQYAEVLSFA	QKQNQTLEKV	AGENKALKKEI	<b>KTLKERVAV</b>	LEEWKDK
4	QTTHYSVVDKDGNAVAVTYTL	NITFGTGIVAGE	SGILLNNQMD	OFSAKPGVPN	<b>YGLVGGDAN</b>	AVGPNKR
5	GTAHLSVVAEDGSAVASTSTI	NLYFGSKVRSPV	SGILFNNEMD	OFSS-PSITN	EFGVPPSPAN	FIQPGKQ
	480 490	500	510	520	530	540
2	MVQWAKFKLPKMRKLAASFFVI	RLECLEKPINTR	MN			
3	MVQWAKEKLPKVRRLAVSFFN	AGIVEKLLNIR	MNWSDKWFS			
4	PLSSMSPTIVV-KDGKTWLVT	GSPGGSRIITTV	LOWVVNSIDY	<b>JLNVAEATNA</b>	PRFHHQWLFD	LRVEKG
5	PLSSMCPTIMVGQDGQVRMVV	GAAGGTQITTAT	ALAIIYNLWF	JYDVKRAVEE1	PRLHNQLLFN	VTTVERN
	550 560	570	580	590	600	
4	FSPDTLKLLEAKGQKVALKEA-	-MGSTQSIMVGPI	GELYGASDPRS	SVDDLTAGY		
5	IDOAVTAALETRHHHTOIAST	FIAWOAIVRTAG	-WAAASDSRKO	GE-PAGY		

Fig. 4. Comparison of the amino acid sequences of the pUH1 and pNKHPSFs, the putative protein C403 of pE194 and the *E. coli* and human hepatic  $\gamma$ -GTPs.

The putative C403 encodes on staphylococcal plasmid pE194 is from Horinouchi and Weisblum (1982). The pNKH PSF is from Hara *et al.* (1994). The *E. coli* y-GTP is from Suzuki et al., (1989). The human hepatic y-GTP sequence is from Sakamuro *et al.*, (1988). Gaps have been inserted to gain maximum matching. The one-letter amino acid code has been used. Percent match of amino acid sequence of pUH1 PSF for C403 on plasmid pE194 was 11.9%, and for *E. coli* and human hepatic  $\gamma$ -GTPs were 7.8% and 8.4%, respectively.

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sequences between the *E.coli* y-GTP and pUH1 PSF is relatively low, and percent match was only 7.8%. Furthermore, the percent match of amino acid sequence among mammalian and *E.coli*  $\gamma$ -GTPs was not so high (ca. 30%). Though mammalian  $\gamma$ -GTPs are connected with metabolism of glutathione-related compounds, but the organisms cannot procedure  $\gamma$ -PGA, like B. *subtilis (natto)* strains. It is, therefore, to be assumed that there are more than one enzyme catalyzes the transfer of the y -glutamyl residue from  $\gamma$ -glutamyl compounds, and that one of them plays a important role in y-PGA synthesis in B. *subtilis (natto)*.

Recently, the B. subtilis y-GTP was cloned and sequenced by 'Katsumata et al. (1991). The sequence contains a single open reading frame encoding the signal peptide and large and small subunits, in that order, as is the case with E.coli y-GTP (Suzuki et al., 1989), but should be concerned with y-PGA production in B. subtilis (natto).

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