



Research article

Twin-arginine signal peptide of *Bacillus licheniformis* GlmU efficiently mediated secretory expression of protein glutaminase

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ABSTRACT

Background: Protein glutaminase specifically deamidates glutamine residue in protein and therefore significantly improves protein solubility and colloidal stability of protein solution. In order to improve its preparation efficiency, we exploited the possibility for its secretory expression mediated by twin-arginine translocation (Tat) pathway in *Bacillus licheniformis*.

Results: The *B. licheniformis* genome-wide twin-arginine signal peptides were analyzed. Of which, eleven candidates were cloned for construction of expression vectors to mediate the expression of *Chryseobacterium proteolyticum* protein glutaminase (PGA). The signal peptide of GlmU was confirmed that it significantly mediated PGA secretion into media with the maximum activity of 0.16 U/ml in *Bacillus subtilis* WB600. A mutant GlmU-R, being replaced the third residue aspartic acid of GlmU twin-arginine signal peptide with arginine by site-directed mutagenesis, mediated the improved secretion of PGA with about 40% increased (0.23 U/ml). In *B. licheniformis* CBB302, GlmU-R mediated PGA expression in active form with the maximum yield of 6.8 U/ml in a 25-l bioreactor.

Conclusions: PGA can be produced and secreted efficiently in active form via Tat pathway of *B. licheniformis*, an alternative expression system for the industrial-scale production of PGA.

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1. Introduction

Protein glutaminases (EC 3.5.1.44) are a kind of enzymes that specifically hydrolyse the amido group on glutamine residues located at the surface of proteins and therefore improve protein solubility and colloidal stability of protein solution [1]. Their representative is *Chryseobacterium proteolyticum* protein glutaminase (PGA), which was discovered and biochemically identified in 2000 [2,3]. The PGA was encoded by a 1053 bp open reading frame and synthesized as a prepro-form, using a 21-amino-acid signal peptide, a 114-amino-acid pro-region (MW = 12,848.2), and a 185-amino-acid mature enzyme

(MW = 19,856.6). The PGA with its pro-region had no enzymatic activity [3].

Since the wild strain of *C. proteolyticum* produced too small amount of PGA to be suitable for industrial applications [3], its heterologous over-expression is an undoubtedly way to be exploited. For this purpose, in previous researches, PGA over-expressed using the Tat pathway in *Corynebacterium glutamicum* has been investigated [4]. It was found that PGA could be produced and secreted efficiently in an inactive pro-enzyme form in *C. glutamicum* mediated by a Tat-dependent signal peptide from *Arthrobacter globiformis* IMD [5,6] or *Escherichia coli* TorA [6,7]. This formed inactive pro-PGA could subsequently be converted to an active form by a subtilisin-like serine protease, SAM-P45 [8]. The maximal accumulation of pro-PGA in *C. glutamicum* reached up to 183 mg/l (about 4.8 U/ml) [4]. These results clearly indicated that the Tat pathway in bacteria could be an alternative for the industrial scale preparation of PGA [4,6,9].

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Remarkably, some strains from *Bacillus licheniformis* have been used for industrial scale enzyme production for over 50 years because of their dramatically high capacity of enzyme synthesis and secretion (more than 25 g/l) [10]. However, their Tat pathways were not well investigated and exploited for the production of enzymes.

In this report, we examined the possibility for high efficiency production of PGA in *B. licheniformis*. The putative twin-arginine signal peptides in *B. licheniformis* were genome-wide selected and functionally identified. The twin-arginine signal peptide of GlmU was found to efficiently mediate the secretion of PGA, and the PGA in active form could be efficiently produced in *B. licheniformis*.

2. Materials and methods

2.1. Strains, plasmids, and culture medium

The plasmids used in this study are listed in Table 1. The nucleotide sequences used for amplification of the relative signal peptide regions are listed in Table 2. *E. coli* JM109 was used for plasmid construction. *B. licheniformis* ATCC 14580 was used as a source for the selection of twin-arginine signal peptides. *Bacillus subtilis* WB600 [11] and *B. licheniformis* CBBD302 [10] were used as the expression hosts. All strains were grown in Luria–Bertani (LB) broth [12] except other mentioned. Kanamycin (20 µg/ml for *E. coli* and *Bacillus* sp.) or tetracycline (25 µg/ml for *Bacillus* sp.) was added to culture media when required. For PGA production, LB broth supplemented with 2% lactose was used as the basic broth. When necessary, another 8% lactose was fed during fermentation in 25-l bioreactor.

Table 2
Primers used in this study.

Tat signal peptide candidates and others	Primers (5'→3') ^a
GlmU	P _{GlmU} -1: ATGGATAATAGGGATAATGGAGGC P _{GlmU} -2: ctcggatccAGGATGAAGAACTTTATATAGCTT TGA
YxaJ	P _{YxaJ} -1: ATGAAAACCAATAGAGTAAGTGTAAATGATCT P _{YxaJ} -2: ctcggatccGAGCAAAACGGTTCGGATTG
YbxG	P _{YbxG} -1: ATGGCGAACAAGAATTGAAGA P _{YbxG} -2: atcggatccGATAAAAAATCCGCAATGGC
YbgF	P _{YbgF} -1: ATGGAACAGTCAAACCAATCGCC P _{YbgF} -2: gtcggatccGACGATCAGAGCCCCGGC
PhoD	P _{PhoD} -1: ATGAAAAAACTGAGCGAGGAAAGCC P _{PhoD} -2: gatggatccATCTCCCAGTCAACGCCAAG
YcgH	P _{YcgH} -1: ATGAAACGACAAGCGAAAAAGGAG P _{YcgH} -2: actggatccAGTCCAAAGGGCAGCAAGAG
YdbS	P _{YdbS} -1: ATGAGAAGCGAGCCGAAAAATCAAA P _{YdbS} -2: gtcggatccCGGTCAAAAATGCCATCCA
YhdH	P _{YhdH} -1: ATGAGCGACCGAGAATGACAG P _{YhdH} -2: ataggatccTATCGTAAAT AGGACAAACA CAAG GAAAAA
YhjN	P _{YhjN} -1: ATGATATTGATCAGCAGTTTGGGG P _{YhjN} -2: ctaggatccTAGCATTGGCCGAAGCTAAGC
AbnA	P _{AbnA} -1: ATGTTAAAGACATCGAAATTTGAAAGGAG P _{AbnA} -2: attggatccAATATGTTCTCTTTTGTATCCCA
YesL	P _{YesL} -1: ATGATTACACAGTTGGCAAATGG P _{YesL} -2: gtcggatccGACATCCGTTTTTCCAGCGCT
pHY-WZX	Primer A: GATTCTCTCCCTTTCAATG Primer B: tctggatccAGAATTCGAGCTCCCGGTTACCAT
pro-PGA	Pga1: cgcggatccTTGGAGCGTTATCCCGGA Pga2: TTAGAAGCGCGAGCTGCTAACG
GlmU-K	P _{GlmU-K} : ATGGATAAAAGGGATAATGGAGGC
GlmU-R	P _{GlmU-R} : ATGGATAGGAGGGATAATGGAGGC

^a The single underlines were *Bam*HI site added; the double underlines were mutation sites.

Table 1
Plasmids used in this study.

Plasmid	Tat signal peptide	Genotypes/Tat signal peptide nucleotide sequence and cloning site ^a	Reference
pHY-WZX	/	<i>amyL</i> promoter and signal peptide, Km ^R , Tet ^R	[13]
pUC-PGA	/	pUC18 containing synthesized pro-PGA encoded gene	This study
pHY-TAT1	GlmU	ATGGATAATAGGGATAATGGAGCCAATACATGATGAAGCGGTTTGCAGTTGTAGCAGCTGGTCAAGGAACAAGAATGAAA TCAAAGCTATATAAAGTTCTTATCTTggatcc	This study
pHY-TAT2	YxaJ	ATGAAAACCAATAGAGTAAGTGTAAATGATCTGACTTTAAATCTCCGCTTTTCTGTTTTGTTCTATGATAGTCGAGCTTCACTTTCTCC GCTCGCTCATTACAGGCCCTCATGCTAATGAGTTTGGCACTTTGGGGATGTTGGCGGCAATCGGAACCGTTTTGCTggatcc	This study
pHY-TAT3	YbxG	ATGGCGAACAAGAATTGAAGAGAGGCGCTGGGCGCGCCACATCAAAATGATTGCGCTTGGCGGCAATCGCGCTCGGTTTATT TATGGGATCTCCAGCACGATAAAGTGACAGGACCTCCGTCCTGCTTATGCAATTTGCGGAATTTTTATCggatcc	This study
pHY-TAT4	YbgF	ATGGAACAGTCAAACCAATCGCCAGAACTTCAAAGAAAAATGCAACGAGACACCTTATCATGCTTCTTAGGAGCGGTTAT CGGCACGGGCTTTTCTAAGTTCCGGTTATACGATCTCACGGCTGTCTGCCGGAACGATCTAGCCTACTTGGCCGGGGCT CTGATCTCggatcc	This study
pHY-TAT5	PhoD	ATGAAAAAACTGAGCGAGGAAAGCCTCAAGGACAATACGTTTGACCGCCCGCTTTATCAAGGGGCGGCAAAATAGCCGGG CTTTCGCTCGACTTTCGATCGCGCAATCGATGGGGCAATGAAAGTCAATGCAGCACCGAGTTCTCCGAATATCCGTTTACACTT GGCGTTGCATCGGGAGATggatcc	This study
pHY-TAT6	YcgH	ATGAAACGACAAGCGAAAAAGGAGATTTAAATGTTGGCAGCTGTCCATGATCGGGGTCGGCTGCACAATGGAACGGGGTTT TTCTCCGGCTCAAGCATCGCGATTAAAGAAAAGCGGCTTTCCGTATTAGCCGCTTCTCTTGTCTGCCCTTGGCACTggatcc	This study
pHY-TAT7	YdbS	aTGAGAAGCGAGCCGAAAAATCAAAATCAGCCGTGATGGCGTTAAAGTCTGGAGAATCACGGCTTGTATCACATCATCTGTTTTAATG CTGGCTGCAGCAGGCTGATTGCCCCGGGTGATATTTAAGTGGCCGATGGATCGGCAATTTTTCAGCGggatcc	This study
pHY-TAT8	YhdH	ATGAGCGACCGCAGAATGACAGCAAAGTGGGATCCAAACTTGGATTGTTCTGCCCGCCGGGGTCCGCAATCGGCTGGGAGC GATTTGGAAGTTTCCCTATGTGGCGGAACGAGCGGGGAGGCGCATTTTCTTGTGTTTTCCTATTTACGATAggatcc	This study
pHY-TAT9	YhjN	aTGATATTGATCAGCAGTTTGGGGGATTTCTTCTCATTGACCGCATGACAATCGGCTGGATGGTGGGAACAATGACCGCCGCGC GCTTGTCTGCTGCTGTTCCCGGTCACCGTTGAAAGCGCGGTGACAGAAAACGGATCCACCATGGATGCTTTCAGTTCCGGC AAATGCTAggatcc	This study
pHY-TAT10	AbnA	ATGTTAAAGACATCGAAATTTGAAAGGAGAACAACCTGTGGCTAAGACTATCATTCTGGTTTTATTTATTTTTTCTGCTATTTTCTC GGCGCTGAACCAACATCAGCGGCTTTTTGGGATACAAAAGGAGACAACATTTggatcc	This study
pHY-TAT11	YesL	ATGATTACACAGTTGGCAAATGGATTTTACCCTTTTGGAGTGGGTGATGGCGCTTGGCTATTGAACTGCTGTGGATCGGCTT TACGCTGCCGGAGCGGTATCTTCGTTTACGCGCGGCGACCCCGCGATGTTCTGCTGTACTAGACAGTGGACCTGGGAAAA ACGGATCTCggatcc	This study
pHY-TAT1r	GlmU-R	Derived from pHY-TAT1 in which the third codon for aspartic acid was changed for arginine	This study
pHY-TAT1k	GlmU-K	Derived from pHY-TAT1 in which the third codon for aspartic acid was changed for lysine	This study
pHY-PGA1	GlmU	PGA-encoded gene in pHY-TAT1	This study
pHY-PGA1r	GlmU-R	PGA-encoded gene in pHY-TAT1r	This study
pHY-PGA1k	GlmU-K	PGA-encoded gene in pHY-TAT1k	This study

^a Underlined were *Bam*HI site added.

2.2. DNA manipulations

DNA manipulations were performed using conventional techniques [12]. Polymerase chain reactions (PCR), with Pyrobest DNA polymerase (TakaraBio, Dalian, China), were performed in 50 μ l working volumes and DNA products were recovered from the agarose gels with the QIAquick gel extraction kit (Qiagen, Germany). Nucleotide sequences were determined using a BigDye terminator cycle sequencing kit (Applied Biosystems) and a DNA Sequencer (model 3200; Applied Biosystems). Plasmid transformations into *E. coli*, *B. subtilis* or *B. licheniformis* were carried out according to the methods as described previously [13].

2.3. Construction of a set of plasmids for pro-PGA secretion using various signal peptides from *B. licheniformis*

When constructing a set of plasmids for pro-PGA secretion, the Tat signal peptide sequence was recovered by PCR with primers listed in Table 2 from *B. licheniformis* ATCC 14580 genome. The signal peptide region in pHY-WZX [13] was then replaced with each of putative twin-arginine signal peptide sequence by BamHI-digested PCR product of pHY-WZX(B-) using Primer A and Primer B (Table 2) and BamHI-digested Tat signal peptide PCR products to yield expression plasmids, pHY-TATs (Fig. 1). For pro-PGA expression, the codon

Table 3
Amino acid residues around putative SPase I cleavage sites^a

-3		-1		+1	
Residue	Frequency (% of total)	Residue	Frequency (% of total)	Residue	Frequency (% of total)
A	39.7	A	83.8	A	38.2
S	10.3	K	8.8	K	10.3
V	7.4	G, F, P, T, M	1.5*5	G	8.8
R	5.9			N, L	7.4 \times 2
L, Q, E, G	4.4 \times 4			I	4.4
T, I, K, M, P	2.9 \times 5			S, V, Y, P, R	2.9 \times 5
Y, N, F	1.5 \times 3			E, Q, F, H, T, W	1.5 \times 6

^a The frequency of a particular amino acid at the indicated positions around SPases I cleavage sites is given as the percentage of the total number of predicted signal peptides in which it appears.

optimized encoding sequence for the pro-PG based on its published amino acid sequence with GenBank accession no. AB046594 was synthesized and cloned into pUC18 plasmid to yield pUC-PGA. The synthesized gene encoding pro-PGA was then recovered by PCR with primers Pga1 and Pga2 (Table 2). It was then digested with BamHI and cloned into each of pHY-TAT plasmids to yield pro-PGA expression vectors, pHY-PGAs.

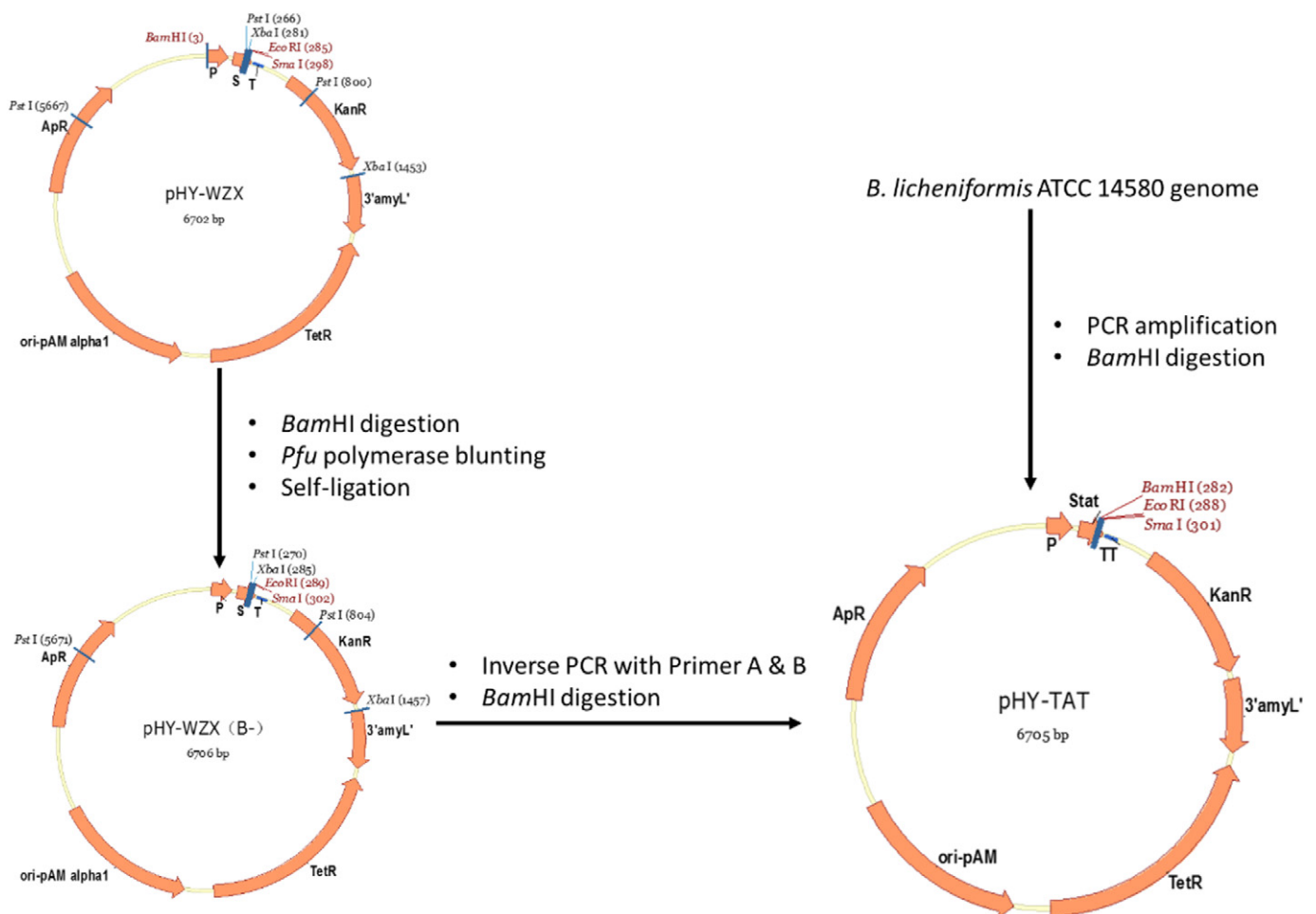


Fig. 1. The construction flowchart of pHY-TAT expression plasmids. A set of plasmids containing 11 *B. licheniformis* twin-arginine signal peptides were constructed based on pHY-WZX. The BamHI site in pHY-WZX was first removed and an intermediate plasmid pHY-WZX(B-) was obtained. The amyL signal peptide was then removed by inverse PCR and a new BamHI site was introduced with Primer B. The putative twin-arginine signal peptides were recovered from *B. licheniformis* ATCC 14580 genome by PCR and a BamHI site at 3'-terminal was introduced with primers. Both PCR products were digested with BamHI and ligated. The ligation mixture was transformed into *E. coli* JM109 and recombinant plasmids were confirmed by restriction pattern and further by nucleotide sequencing. The constructed plasmids were nominated as pHY-TAT1, pHY-TAT2, and so on.

The site-directed mutagenesis was carried out based on PCR technique as described above. The oligonucleotides primers P_{GlmU-K} and P_{GlmU-R} (Table 2) were used to mediate mutagenesis. P_{GlmU-K} and P_{GlmU-2} or P_{GlmU-R} and P_{GlmU-2} were used to amplify GlmU coding sequence using plasmid pHY-PGA1 as template. The BamHI-digested PCR products were ligated with the reverse PCR product of pHY-WZX (B-) followed by inserting the pro-PGA encoding sequence as

described above (Fig. 1) to yield the recombinant expression plasmid pHY-PGA1k and pHY-PGA1r.

2.4. Fermentation test

For selecting candidate Tat signal peptide, the transformants were incubated at 37°C and 220 rpm in 250-ml shaking flask with working

Table 4
Eleven Tat signal peptide candidates from *B. licheniformis* ATCC 14580.

ORFs	Note in the genome	Signal peptide and cutting site		Signal sequence in details
GlmU	bifunctional UDP-N-acetylglucosamine pyrophosphorylase/glucosamine-1-phosphate N-acetyltransferase GlmU	MDNRDNGGQYM DKRFAV <u>VL</u> AJAG Q	1 1 61 21	ATGGATAATAGGGATAATGGAGGCCAATACATGGATAAGCGGTTTGACAGTTGTTAGCA M D N R D N G G Q Y M D K R F A V V L A 70 80 90 100 110 120 GCTGGTCAAGGAACAAGATGAAATCAAAGCTATATAAAGTTCTTCATCCTGTTTGGCGGA A G Q G T R M K S K L Y K V L H P V C G
YxaJ	putative ORF	MKTNRVSMIWT LISAFLCMIVA ASLSPLAHSGPH AINE	1 1 61 21 121 41	ATGAAAACCAATAGAGTAAGTGTAAATGATCTGGACTTAAATCTCCGCTTTTCTGTTTTGT M K T N R V S V M I W T L I S A F L F C 70 80 90 100 110 120 TCTATGATAGTCGAGCTTCACTTTCTCCGCTCGCTCATTCAGGCCCTCATGCTAATGAG S M I V A A S L S P L A H S G P H A N E 130 140 150 160 170 180 TTTGGCACTTTGGGATGTGGCGGCAATCGGAACCGTTTTCGCTTTTATATGCTGCC F G T L G M W A A I G T V L L F Y M L P
YbxG	amino acid permease	MANKELKRGLG ARHIQMIALGTTI GVGLFMGSSSTI KJWT	1 1 61 21 121 41	GTGGCGAACAAAGAATGAAGAAGGCGCTGGGCGCGCCACATCCAATGATTCGCGCTT V A N K E L K R G L G A R H I Q M I A L 70 80 90 100 110 120 GGCGCACATTCGCGCTCGGTTTATTTATGGGATCTCCAGCACGATAAAGTGGACAGGA G G T I G V G L F M G S S S T I K W T G 130 140 150 160 170 180 CCTTCGCTCTGCTTGTATGCCATTGGCGAATTTTATCTTTTATTTATGCGTGCA P S V L L A Y A I C G I F I F F I M R A
YbgF	putative amino acid permease YbgF	MEQSNTNRQNF QRKMOTRHLML SLGGVIGTGLFSL SGYTSQAIGP	1 1 61 21 121 41	ATGGAACAGTCAAACACCAATCGCCAGAACCTTCAAAGAAAATGCAACCGAGACACCTT M E Q S N T N R Q N F Q R K M Q T R H L 70 80 90 100 110 120 ATCATGCTTTCCTTAGGAGCGTTATCGGCACGGGCTTTTCTTAAGTTCCGGTTATACG I M L S L G G V I G T G L F L S S G Y T 130 140 150 160 170 180 ATCTCAGAGGCTGGTCTCGCGGAACGATTCTAGCCTACTTGGCGGGGCTGTGATCGTC I S Q A G P A G T I L A Y L A G A L I V
PhoD	phosphodiesterase/alkaline phosphatase	MKKLSEESLKDN TFDRRRFIQAG KIAGLSLGLAIAQ SMGAMEVNAJAP	1 1 61 21 121 41 181 61	ATGAAAAAAGTGGAGGAGAAAGCCCTCAAGGACAATACGTTTGACCGCCCGCTTTATT M K K L S E E S L K D N T F D R R R F I 70 80 90 100 110 120 CAAGGGCCGCAAAATAGCCGGGCTTTCGCTCGGACTTCCGATCGCGCAATCGATGGGG Q G A G K I A G L S L G L A I A Q S M G 130 140 150 160 170 180 GCAATGGAAGTCAATGCAGCACCGAGGTTCTCCGAATATCCGTTTACACTTGGCGTTGCA A M E V N A A P R F S E Y P F T L G V A 190 200 210 220 230 240 TCGGGAGATCCGCTTTCGACAGCGTGTATTGTGACAAGGCTGGCGCCGATCCGCTA S G D P L S D S V V L W T R L A P D P L
YcgH	putative amino acid transporter YcgH	MKRQAKKGLDK WWQLSMIGVC TIGTFGLSSIAI KJKS	1 1 61 21 121 41	ATGAAACGACAAAGCGAAAAAGGAGATTAAATGGTGGCAGCTGCCATGATCGGGTCC M K R Q A K K G D L K W W Q L S M I G V 70 80 90 100 110 120 GGCTGCACAATGGAACGGGTTTTTCTCCGCTCAAGCATCGCGATTAAAGAAAGCGGG G C T I G T G F F L G S S I A I K K S G 130 140 150 160 170 180 TTTTCCGATTAGCCGCTTCTTCTTGCTGCCCTTGGCACITTTCTTGTTTTTCAGCAG F S V L A A F L L A A L G T F L V F Q Q
YdbS	cytosolic protein YdbS	MRSEPKNQISRD GVKVVWITACITS FVLMIAAAGLIAI AG	1 1 61 21 121 41	TTGAGAAGCGCCGAAAAATCAAATCAGCCGTGATGGCGTTAAAGTCTGGAGAATCAGC L R S E P K N Q I S R D G V K V W R I T 70 80 90 100 110 120 GCTTGATCACATCATTCTTTAATGCTGGCTGCAGCAGGCTGATTCGCCCGGGGTG A C I T S F V L M L A A A G L I A A G V 130 140 150 160 170 180 ATATTTAAGTGGCCGATAAGGATCGGCATTTTGTACAGCGGCTTTGGCTGGGATTCA I F K W P V W I G I L S A A V W L G I S

Table 4 (continued)

YhdH	putative sodium-dependent transporter YhdH	MSDRRMTAKWA SKLGFVLAAGS AIGLGAIWKFPYV AIG	1	ATGAGCGACCCGAGAATGACAGCAAATGGGCATCCAACTTGGATTGTTCGCGCC
			1	M S D R R M T A K W A S K L G F V L A A
			61	70 80 90 100 110 120
			21	GCGGGTCCGCCATCGGTCTGGAGCGATTGGAGTTCCCTATGTGGCGGAACGAGC A G S A I G L G A I W K F P Y V A G T S
			121	130 140 150 160 170 180
41	GGGGAGGCGCATTTTCTTGTGTTGTCTTATTTACGATACTGCTGGGATACCCCTG G G G A G F F L V F V L F T I L L G Y P L			
YhjN	putative ammonia monooxygenase YhjN	MILISFGFLLS LTGMTIGWMVGT MTAAACLSLFRP SPLKAJAV	1	TTGATATTGATCAGCAGTTTGGGGATTCTTCTCTCATTGACCGGCATGACAATCGGC
			1	L I L I S S F G G F L L S L T G M T I G
			61	70 80 90 100 110 120
			21	TGGATGGTCGGAACAATGACCGCCGCGCTTGTCTGCTGCTGTTTCGCCGTCACCGTTG W M V G T M T A A A C L S L F R P S P L
			121	130 140 150 160 170 180
41	AAAGCGCGCTCAGACAAAACGGCATCCACCATGGATGGCTTCAGTTGGCCAAATGCTA K A A V R Q N G I H H G W L Q F G Q M L			
AbnA	arabinan endo-1,5-alpha-L-arabinosidase AbnA	MLKTSKFERRT VAKTIISGFILFFL LIFSAIAEP	1	ATGTTAAAGACATCGAAATTTGAAAGGAGAACAACTGTGGCTAAGACTATCATTTCTGGT
			1	M L K T S K F E R R T T V A K T I I S G
			61	70 80 90 100 110 120
			21	TTTATTTATTTTCTGCTTATTTCTCGGCGGCTGAACCAACATCAGCGGCTTTTGG F I L F F L L I F S A A E P T S A A F W
			121	130 140 150 160 170 180
41	GATACAAAAGGAGACACATTTATCATGATCCTCCATGATTAAGAAGAAATACGTGG D T K G D N I I H D P S M I K E G N T W			
YesL	putative membrane protein YesL	MIHTLANGFYRF CEWVMRLAYLNL LWIGFLAGAVIF GLAPATAJAM	1	ATGATTCACACGTTGGCAAATGGATTTTACCGCTTTGCGAGTGGTGATGCGGCTTGGC
			1	M I H T L A N G F Y R F C E W V M R L A
			61	70 80 90 100 110 120
			21	TATTTGAATCTGCTGGATCGGCTTTACGCTGGCGGAGCGGTCATCTTCGGTTTAGCG Y L N L L W I G F T L A G A V I F G L A
			121	130 140 150 160 170 180
41	CCGGCGACCCCGCGATGTTGCTGTGACTAGACAGTGGACGCTGGGAAAACGGATGTC P A T A A M F A V T R Q W T L G K T D V			

volume of 50 ml. When fermenting in a 25-l bioreactor (Bioflow110; New Brunswick Scientific Co., Inc., Edison, NJ), the recombinant strain was incubated at 42°C, 1 vvm aeration and 200–700 rpm agitation with the initial working volume of 10 l. When necessary, a total of 800 g of lactose in 300 g/l of lactose solution was fed.

2.5. PGA activity assay

PGA activity was assayed according to reference [2] using Cbz-Gln-Gly (Peptide Laboratory, Shanghai, China) as a substrate at pH 6.5 and 37°C. One unit of PGA is defined as the amount of enzyme that releases 1 μmol of ammonium per minute under the assay conditions specified.

2.6. Database mining of *B. licheniformis* Tat signal peptides

The signal peptide prediction of twin-arginine signal peptides in *B. licheniformis* ATCC 14580 genome was made using TatP 1.0 (<http://www.cbs.dtu.dk/services/TatP/>).

3. Result and discussion

3.1. Analysis of *B. licheniformis* Tat signal peptides

Based on the consensus sequence of Tat signal peptides, all putative twin-arginine signal peptides in *B. licheniformis* ATCC 14580 genome were predicted using TatP 1.0 server according to the presence of an RR(FGAVML)(LITMVF) motif [14]. A total of 4156 putative ORFs in the genome sequence were analyzed. Of which, 58 ORFs containing putative twin-arginine signal peptides were finally predicted. Their hydrophobic core (H-domain) had an average length of 18 amino acid residues. Their SPases 1 recognition sequences consisted of small aliphatic residues at positions -3 and -1. The preferred alanine and

lysine residues were at the position -1 and the residue, such as alanine, lysine, serine, valine, arginine or leucine accommodated the position -3 (Table 3). The most frequent residue was alanine accounting at positions -3 and -1 for 39.7% and 83.8%, respectively (Table 3). Of total 58 putative Tat signal peptides in *B. licheniformis*, 11 putative Tat signal peptides were randomly selected for further study and their sequences are summarized in Table 4.

3.2. Tat-dependent secretion of PGA mediated by Tat signal peptides of *B. licheniformis*

To illustrate the possibility of the Tat signal peptides from *B. licheniformis* for mediating PGA secretion, the nucleotide sequences encoding for 11 selected twin-arginine signal peptides were recovered from *B. licheniformis* ATCC 14580 genome by PCR amplification. The PCR products were purified and digested with *Bam*HI, and then cloned into the plasmid pHY-WZX to replace the AmyL signal peptide [13]. A total of 11 expression plasmids, nominated as pHY-TAT₁₋₁₁, were constructed, which were further confirmed by PCR amplification, restriction pattern and nucleotide sequencing (Fig. 1).

To express the pro-PGA, the chemically synthesized pro-PGA coding sequence was recovered, digested with *Bam*HI, and cloned into each of pHY-TAT plasmids digested with *Bam*HI and *Sma*I, yielding recombinant plasmid pHY-PGAs. The transformants were obtained by transforming the recombinant plasmid pHY-PGAs into *B. subtilis* WB600. The fermentation tests were carried out and the PGA activities in the supernatant were examined. The activities are summarized in Table 5. The twin-arginine signal peptides of GlnU, YhdH and YxaJ were able to mediate the PGA secretion in *B. subtilis*. Of which, GlnU twin-arginine signal peptide mediated the highest PGA secretion and YhdH and YxaJ twin-arginine signal peptides mediated very weak but detectable PGA secretion (Table 5). GlnU in *B. licheniformis* is a

Table 5

B. licheniformis twin-arginine signal peptides mediated PGA secretion in *B. subtilis*.

Tat signal peptide	PGA activity (mU/ml)
GlmU	160 ± 25
Yxaj	N.D. ^a
YbxG	57 ± 13
YbgF	N.D.
PhoD	N.D.
YcgH	N.D.
YdbS	N.D.
YhdH	24 ± 15
YhjN	N.D.
AbnA	N.D.
YesL	N.D.
AmyL	N.D.

^a N.D.: Not detectable.

putative (yet its function not identified) bifunctional UDP-N-acetylglucosamine pyrophosphorylase/glucosamine-1-phosphate N-acetyltransferase for the synthesis of UDP-N-acetylglucosamine [15].

Several previous reports indicated that the PGA might fold too rapidly to be treated by the heterologous Sec machinery [8,16]. Different from the Sec pathway, the Tat pathway tends to translocate its substrates in a folded conformation [17,18,19]. Using Tat signal peptides from *E. coli* TorA or *A. globiformis* IMD, PGA was successfully secreted with the Tat pathway in *C. glutamicum* [4]. In this report, we found that three twin-arginine signal peptides from *B. licheniformis* were able to indicate the production of PGA in *B. subtilis* in active form and the different secretion efficiencies were also observed (Table 5). Interestingly, PGA in *B. subtilis* WB600 was produced in active form without additional processing, for example, being converted to an active form by removing the propeptide with a subtilisin-like serine protease SAM-P45 [8]. These results suggested that an unknown peptidase accurately digested pro-PGA to active one in *B. subtilis* WB600 although its six extracellular proteases (neutral protease A, subtilisin, extracellular protease, metalloprotease, bacillopeptidase F, and neutral protease B) were deleted [11]. *B. subtilis* WB600 kept 0.32% of the wild-type extracellular protease activity and no residual protease activity could be detected when it was cultured in the presence of 2 mM phenylmethylsulfonyl fluoride [11].

3.3. Modification of N-terminal of GlmU twin-arginine signal peptide improved PGA secretion

The positive charge of the N-region is generally considered to be essential for recognition by translocase to initiate protein translocation and to interact with the negatively charged lipids at the cell membrane as well although its importance is uncertain [20]. The third residue, aspartic acid (N) of GlmU twin-arginine signal peptide, was replaced with lysine (K) or arginine (R) by site-directed mutagenesis. Two mutants, GlmU-K and GlmU-R, were obtained (Fig. 2A). Their effects of amino acid alterations on the secretory performance of PGA were evaluated. The change of the N-region's amino acid residues from N to K (GlmU-K) or to R (GlmU-R) resulted in a different level of PGA secretion in *B. subtilis* WB600. GlmU-R mediated the improved secretion of PGA with about 40% increase (230 mU/ml). GlmU-K did, however, reduced the secretion of PGA (Fig. 2B).

It was found that the N-region's positive charge density in alkaline phosphatase signal peptide increased the secretory efficiency of *E. coli* alkaline phosphatase [21]. The mutant GlmU-R developed in this study was to increase the N-region's net charge by the introduction of a positive charge in the form of arginine residue, which might also make it more susceptible by a signal peptide peptidase [22].

3.4. Over-expression of PGA in *B. licheniformis*

To examine the possibility of PGA production in *B. licheniformis*, the recombinant plasmid pHY-PGAR was transformed into *B. licheniformis* CBBD302, an industrial enzyme expression host [10]. The yield of PGA in *B. licheniformis* was determined in a 25-l fermentor (Fig. 3). With no surprise, PGA was produced and secreted in its active form in *B. licheniformis*. The maximal yield was 6.8 U/ml in the supernatant. To our knowledge, this is the highest yield ever reported.

4. Conclusion

In conclusion, PGA can be efficiently produced and secreted in active form via Tat pathway of *B. licheniformis*, an efficient alternative for the industrial-scale production of PGA.

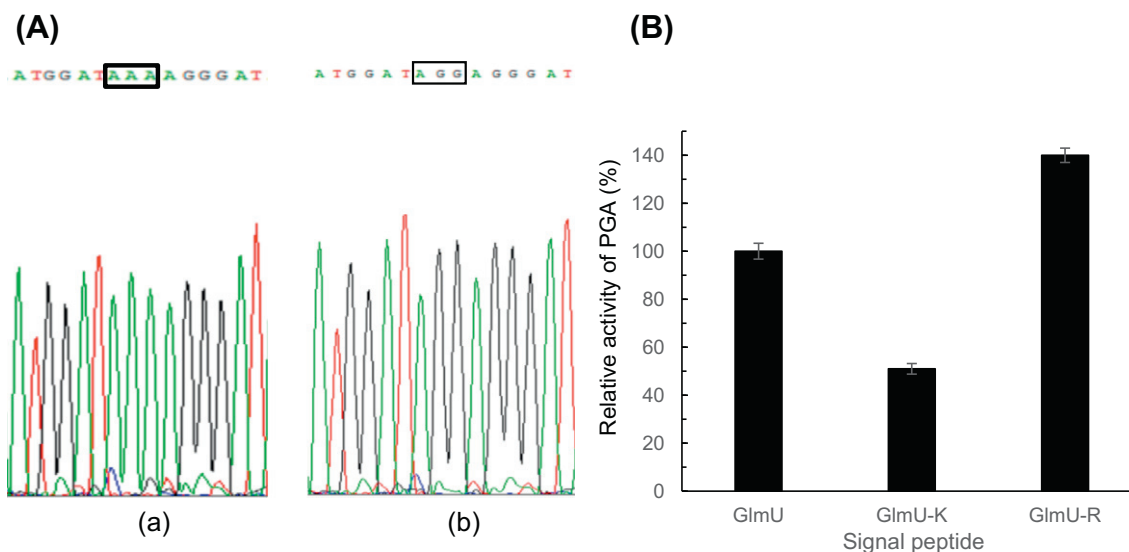


Fig. 2. Effects of mutant GlmU signal peptide on the secretion of PGA. (A) The sequencing analysis of the site-directed mutants of GlmU, GlmU-K (a) and GlmU-R (b). (B) Secretory activity of PGA mediated by GlmU mutants. The activity mediated by wild-type GlmU was taken as 100%.

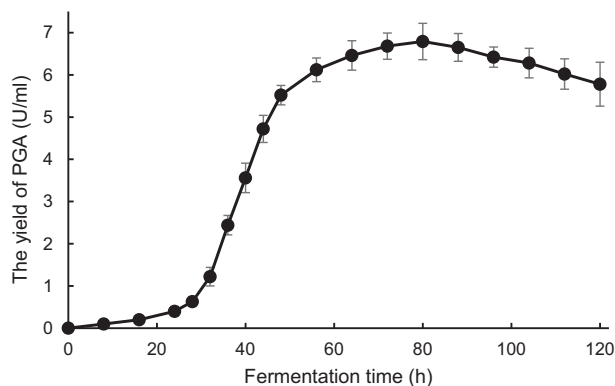


Fig. 3. The time-course of PGA production in *B. licheniformis*. The production of PGA was taken place in a 25-l bioreactor with initial working volume of 10 l. LB complemented with 40 g/l of lactose was used as medium. During the fermentation about 267 ml of 300 g/l lactose (total 800 g) was fed.

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Conflict of interest

The author(s) declare that they have no competing interests.

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