Nucleotide sequence and organization of *Bacillus subtilis* RNA polymerase major sigma (σ^{43}) operon

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ABSTRACT

The gene coding for <u>Bacillus subtilis</u> RNA polymerase major σ^{43} , <u>rpoD</u>, was cloned together with its neighboring genes in a 7 kb <u>Eco</u>RI fragment. The complete nucleotide sequence of a 5 kb fragment including the entire <u>rpop</u> gene revealed the presence of two other genes preceding <u>rpoD</u> in the order <u>P23-dnaE-rpoD</u>. The dnaE codes for DNA primase while the function of P23 remains unknown. The three genes reside in an operon that is similar in organization to the <u>E.coli</u> RNA polymerase major σ^{70} operon, which is composed of genes encoding small ribosome protein S21 (rpsU), DNA primase (dnaG), and RNA polymerase σ^{70} (rpoD). There is a relatively high degree of base and amino acid homology between the DNA primase and σ genes. The most significant differences between the two operons are observed in the molecular size of the first genes (P23 and rpsU), the complete lack of amino acid homology between P23 and S21, the molecular weights of the two rpoD genes, the size of the intercistronic region between the first two genes, and the regulatory elements of the operon.

INTRODUCTION

The existence of multiple RNA polymerase σ factors in <u>B. subtilis</u> has been well documented (1,2), but little is known about their genetic properties, the regulation of their synthesis, and the factors that govern their interactions with the RNA polymerase core. An analysis of their molecular organization and the parameters which regulate their genetic expression should provide a initial basis for determining their roles in the physiology of this Gram positive sporulating bacterium.

Our laboratory has been particularly interested in the study of the σ^{43} gene (<u>rpoD</u>), whose product is known to play a major role during vegetative growth, and the early stationary and sporulation phases (3). We have been able to clone (4), genetically map (5), and sequence (6) the σ^{43} gene (<u>rpop</u>), and show that its derived amino acid sequence had a very high degree of homology with that of the <u>E.coli</u> major σ^{70} (7). By genetic mapping (5) and DNA sequencing (8), we also showed that immediately upstream of the <u>rpop</u> gene was located the <u>dnaE</u> gene, which encodes the <u>B. subtilis</u> DNA primase, whose product is very homologous to the <u>E.coli</u> <u>dnaG</u> DNA primase (9,10). No promoter region was observed in the intercistronic region between <u>rpop</u> and <u>dnaE</u>, nor in the region immediately upstream of <u>dnaE</u> (6,8).

Recently, we have determined the nucleotide sequence of the region upstream of <u>dnaE</u> including the operon regulatory region, which provided support for our previous suggestion (6,8) that <u>dnaE</u> and <u>rpoD</u> were coordinately regulated with one or more unknown genes in an operon. The DNA sequence analysis of the region upstream of <u>dnaE</u> revealed an open reading frame capable of coding for a protein of molecular weight 22,540. The function of this protein is unknown, and hence the designation <u>P23</u> is being used for this gene until a physiological role can be assigned to it.

In this paper we will discuss the similarities and differences of the structure and organization of the major sigma operons of <u>B. subtilis</u> and <u>E. coli</u>, the transcriptional and translational regulatory features of the operon, and the codon usage frequency encountered in the operon.

MATERIALS AND METHODS

Strains, Phages and Plasmids

<u>E. coli</u> JM101 was used as host for the sequencing phage vectors M13mp8, M13mp9, M13mp10, and M13mp11 (11,12), and the plasmid pCPS1 (5). <u>E. coli</u> BNN45 (13) was used to prepare the phage lysate of λ gtWES- σ 82 (4). Plasmid pSB was provided by Sui-Lam Wong (unpublished data).

DNA Manipulations

Standard procedures of Maniatis <u>et al</u>. (14) were followed exactly as described.

DNA Sequencing

DNA sequencing was conducted by the dideoxy chain

termination method of Sanger <u>et al</u>. (15) using the sequencing kit purchased from Amersham Corporation.

Computer Analysis

Routine analysis of DNA or protein sequences were carried out using either the Delaney (16) or the Pustell (17) program, while the homology search against the NBRF Data Bank was made using the Microgenie Sequence Analysis Program developed by Queen and Korn (18).

RESULTS

Nucleotide sequence of the Entire Operon

The nucleotide sequences and the sequencing strategies of <u>dnaE</u> and <u>rpoD</u> genes have been reported previously (6,8). The sequencing strategy for the upstream 1.5 kb fragment is shown in Fig. 1 (bottom) along with the physical map of the σ^{43} operon (upper). As indicated, the nucleotide sequence has been determined for both strands of virtually the entire region except for the 100 bp at the extreme 5' end. The sequence was determined across the junctions of all the restriction sites used for subcloning during sequencing, as well as for the EcoRI site between the <u>dnaE</u> and <u>rpoD</u> genes (not shown here). In our previous reports, the sequences for these two genes were determined separately (6,8). Although unlikely, the possibility existed that a small EcoRI fragment may have been left out during the subcloning of the EcoRI fragments into plasmids from the original phage λ gtWES- σ 82 (4,5). Therefore we sequenced the 0.9 kb HindIII fragment containing the EcoRI junction region, which was subcloned into M13mp10 directly from λ qtWES- σ 82, and the possibility mentioned above has been experimentally excluded. Now, the entire EcoRI-SphI fragment has been sequenced, including all the junctions of restriction sites used for sequencing. The nucleotide sequence of the entire operon and its flanking regions, and the deduced amino acid sequence of each gene are given in Fig. 2 with the first base of the 5' end EcoRI site labeled as number 1.

Features of the First Gene of the Operon

When the sequence of the region upstream of <u>dnaE</u> was analyzed by computer, only one large open reading frame was

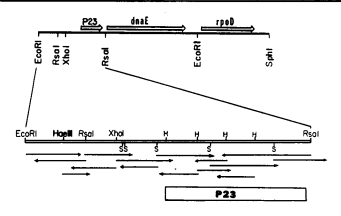


Figure 1. Sequencing strategy for the first gene and regulatory region of the σ^{43} operon.

The upper part represents the physical location of the genes in the operon. The lower part indicates the restriction sites used for subcloning and sequencing. The bar indicates the location of the cryptic P23 protein. Abbreviations: H, <u>Hpa</u>II; S, <u>Sau</u>3A.

discovered. But unlike the case for the other two genes in the operon, we could not identify any strong ribosomal binding site by sequence analysis within the open reading frame. We found instead several weak ones preceeding the potential initiation codons ATG (855), TTG (930), ATG (951), and GTG (1,200), which were able to code for proteins of molecular weights 22,540, 19,734, 18,934 and 9,312, respectively. However, our previous maxicell data showed that a protein of molecular weight around 23,000 was encoded within the upstream region of <u>dnaE</u> (4,5), which corresponded very well with the largest open reading frame identified here by sequencing, i.e., the open reading frame starting from the initiation codon ATG at nt 855.

Although the ribosomal binding site was very weak as predicted from its calculated free energy of binding ($\Delta G = -9.2$ kcal/mol), our assignment was further strengthened by our recent protein fusion studies. When the N-terminal two thirds of the P23 was fused to the <u>E. coli</u> β -galactosidase in frame, a functional hybrid protein was expressed in both <u>E. coli</u> and <u>B.</u> <u>subtilis</u> with the expected size as determined by Western blot analysis using anti- β -galactosidase antibody (data not shown). Functional P23- β -galactosidase (P23-gal) fusion protein was expressed even when only the first 8 amino acid residues of P23

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SGA CC	317 GA	5	NGA CA	ATG G1 Het V	NGA CA	11 NG	AC TC	44 53	AA AA	11C 66	CGG AA		TAATA	ATTTA	ATANA	-	-
CTC Val	0 000	VIE	ATC 1	E.F.	617 1	Arg CGC	GAT O	ALA J	GLUG	GAG G	Leu		654 NTG QJ	TIA N	NTA GI	IGA A1	NG TC
TAT 0	0 111	t Ly	6 166 P 1rp	C GTT 0 Val	C TTA P Leu	A GTA • Val	A CAG	P E C	T CGT P Arg	A GAA	A GCG	< 7	GATAT	6TGGT	ATACA	AAGTG	ATTAG
A CO	W SH	eu He	94 94	55	54 G	24 25	11 11 12	52	57 57	63 56	Le ty	MA TA	14 V 14 V	11	11 07	TA TA	5 TE
SUS ALA	3998 564 AAC ATG 66C C	CT/		4118 C00		1238 757 757	4298 ATC	202 E	₿₿	4478 GGC CGT ACA AGA ACA TTA GAA GAG GTC GLY Arg Thr Arg Thr Leu Glu Glu Val	4538 GAA (4598 AAA GAT TTC CTT GAA TAA Lys Asp Phe Leu Glu End	4634 4644 4654 4654 4654 <u>Copyc</u> tte titttiaaa aagatatate gataatatee	4634 4634 4694 4734 4734 4734 4734 4734 4734 4734 47	4734 4744 4754 4754 4764 4774 4784 4784 4784 4784 4784 478	4814 4824 4834 4834 4844 Tectatit cantitigta taaagtgiga ataaaaact	4874 4884 49 94 4904 Titic taa aastaag ta gaattagaag t giti acata
T ATC	C ATG	n Het	TAT 5	G ATC	A 11 1 10	4 ACG	500 F	7 7CA	T ACG	t ACA	A ATC	T TTC P Phe	34 TC TT	10 CV	S4 GT AT	3	71 M
TC AG	CA AA	IY As	GT AC	IGA AC	50 CF	11 11 12 14	N N	24 51	TT CA	17 AC	ភិថិ ភូទ	AA GA' Ye Aei	erret.	TCATT	TAAGO	TCTAT	19111
CTT C	GAA G	clu	e s	A14 A14	ទីទី		ÊĴ	GAA G GLU A	550	GAC Asp	Ę.	trev ∧ Lev ∧	A TC0	1 ATC	1	5	E F
G CTT GTT g Leu Val	CAT	H	T MA	P CAG	CGT GTG Arg Val	GCG GAA GAT ATG Ala giu Abp Het	5 CT	CAA GAA	Asp Asp	GAT	CGT Årg	AAA CGT TTG	4614 4624 6ATG <u>0ACGG GTCTT9AAGA TC</u>	469	5 EE	4794 4794 4804 Gatattagag atttagggat	1111GTATAGC AATCCATTTA CT
CTG CGG	77 Q	Leu Ile	667 7AT 617 771	GCC GAT Ala Asp	TC CC 1e Are	18 GL	CCC GTA Pro Val	GAA GAC Glu Asp	CTG GAA Leu Glu	GGT CTT Gly Leu	AGA GAG Arg Glu	it Mi	513	CLG	ATAT	ELV :	, MT
ANC And And And And And And And And And And	SAT C	Asp L	50 50 50 50	ATT 60	TTA ATC Leu 11e	ATT G	GAG CC GLU PC	ATT 61 11e 61	CAG CI 611 Le	Phe G	ACG AG	AGA AGT Afg Sef	461-	C ST	CTTCC	TAGAC	485. IATAG
Ala C	5	Let	4 200 A 19	900 N	¥3	Gut G	5 G	THC A	GAG C	CGA T	GTA A Val 1	AGC A Ser A	GATG	CHI	ATCTA	GATA1	1116

Table 1. Amino Acid	Compos	ition (mo	ol%) Analy	ysis
Group	<u>P23</u>	dnaE	rpoD	Ave. B.s. Proteins*
Small aliphatic (A+G)	10.2	13.3	11.2	15.0
Hydroxyl (S+T)	12.2	11.0	9.6	13.4
Acidic (D+E)	14.3	14.3	21.1	14.2
Acidic + acid amide (D+E+N+Q)	22.4	22.3	28.0	23.1
Basic (K+R+H)	14.3	16.7	16.0	14.8
Hydrophobic (L+I+V+M)	22.9	22.8	24.9	26.3
Aromatic (F+Y+W)	8.7	9.5	6.0	7.7
Charged (D+E+K+R+H)	28.6	31.0	37.1	29.0

Table 1. Amino Acid Composition (mol%) Analysis

* The amino acid composition of average <u>B. subtilis</u> proteins is calculated from 35 sequenced genes published up to 1985.

were fused to the 8th amino acid residue of β -galactosidase, which led us to the conclusion that ATG (855) was functionally active <u>in vivo</u>. Hence, we designated the first gene of the operon as <u>P23</u> from these data and for the reason that its physiological function is still unknown.

The deduced amino acid sequence of P23 was examined for homology against the NBRF Protein Data Bank using the Microgenie Program (18), but no significant homology was found to any of the known proteins in the bank, indicating to us that P23 was not homologous with <u>E. coli</u> S21, the first gene in the σ^{70} operon (also their size difference is significant), and that P23 might be unique to <u>B. subtilis</u>, or that its counterpart in <u>E.</u> coli has not been characterized as yet. The latter case is a possibility, since a reasonable degree of homology has been found between many <u>B. subtilis</u> and <u>E. coli</u> proteins.

The deduced amino acid composition of P23 is shown in Table 1, together with those of DNA primase, σ^{43} and an average of <u>B. subtilis</u> proteins for comparison. One difference noted from σ^{43} , which is a highly acidic protein typical of most transcription factors (7,19,20), is that P23 is more like the average composition of <u>B. subtilis</u> proteins. Thus it is difficult to categorize this protein based on its amino acid composition. We are currently raising antibody against P23 using a P23-gal fusion protein as antigen, hoping that this will provide us a tool to determine the location and possibly the function of P23 in <u>B. subtilis</u>.

Regulatory Features

Previously, we reported that there was no promoter activity detected within the intercistronic regions of the operon except for a weak heat shock promoter activity located at the C-termianl end of the <u>dnaE</u> gene (8). So we concluded that a promoter(s) should exist in front of P23 if the operon was composed of three genes as in the case of <u>E.coli</u> σ^{70} operon (10). By sequence analysis we did find at least two potential promoters with significant homology to the consensus sequence of B. subtilis σ^{43} promoters (1,2), which were then confirmed to function in vivo by fusing the 211 bp Sau3A fragment (609-821) to the subtilisin gene (aprA) in a promoter-probe plasmid pSB (Wong and Doi, unpublished data). These sequences are underlined in Fig. 2, and designated as Pl and P2. To our surprise, one additional promoter activity was detected when the 316 bp Sau3A fragment (829-1136) downstream of Pl and P2 was cloned in pSB. This promoter (P3) was temporally regulated in that it was not expressed until the culture reached the sporulation phase, while Pl and P2 were expressed efficiently mainly during growth. More detailed mapping and functional characterization of these promoters are in progress, and will be published elsewhere.

Earlier sequence analysis (6) allowed us to identify a sequence typical of rho independent terminator (21) immediately following the TAA stop codon of <u>rpoD</u> gene (underlined in Fig. 2). Recently, we confirmed its termination activity <u>in vivo</u> by subcloning the <u>PvuII-Aha</u>III fragment (4394-4641) into a <u>B. subtilis</u> terminator-probe plasmid pST19 constructed in our laboratory (Wang and Doi, unpublished data). We were able to show that introduction of this 247 bp fragment reduced the activity of the indicator enzyme in the terminator probe (subtilisin, in this case) by more than 90% compared to the control (vector alone), indicating that this was a relatively strong terminator (data not shown).

Thus we have determined the presence of three genes in the

			able a	· · · · ·	ion use	ige or	D.SUDI		ordu	1a-43 U	Peron		
AA (Codor	P23	<u>dna E</u>	<u>rpoD</u>	B.s.	E.c.	AA (Codon	<u>P23</u>	dnaE	rpoD	B.s.	E.c.
		1.00	.73		.67	.37		UAU		.74		.62	.40
	UUC	.00	.27	.55	.33	.63		UAC	.14		.20	.38	.60
Leu	AUU	.24	.20	.31	.23	.07				1.00		.57	.75
Leu	UUG	.19	.12	.15	.13	.09	AMB	UAG	.00	.00	.00	.06	.08
	CUU	.29	.29	.28	.24	.07		CAU	.60	.78	.50	.68	.54
	CUC		.10	.00	.11	.07		CAC	.40	.22	.50	.32	.46
	CUA	.05	.03	.08	.05	.02		CAA	.71	.53	.65	.56	.24
Leu	CUG	.24	.26	.18	.23	.68	Gln	CAG	.29	.47	.35	.44	.76
Ile	AUU	.62	. 39	.54	.50	.36	Asn	AAU	.78	.58	.67	.53	.26
Ile	AUC	.23	.35	.40	.40	.61	Asn	AAC	.22	.42	.33	.47	.74
Ile	AUA	.15	.26	.00	.10	.03	Lys	AAA	.73	.73	.88	.75	.76
Met	AUG	1.00	1.00	1.00	1.00	1.00	Lys	AAG	.27	.27	.12	.25	.24
Val	GUU	.39	.29	.43	.20	.36	Asp	GAU	.83	.64	.58	.63	.46
Val	GUC	.23	.31	.14	.24	.15		GAC	.17	.36	.42	.37	.54
Val	GUA	.15	.26	.33	.23	.22	Glū	GAA	.81	.63	.77	.69	.73
Val	GUG	.23	.14	.10	.23	.27	Glu	GAG	.19	.37	.23	.31	.27
Ser	ບຕບ	.20	.30	.31	.25	.23	Cvs	UGU	.33	.20	.00	.50	.43
	UCC	.27	.09	.06	.12	.27	Cvs	UGC	.67	.80	.00	.50	.57
	UCA	.20	.13	.13	.14	.07		UGA	.00	.00	.00	.37	.17
	UCG	.20	.15	.06	.11	.11	Trp	UGG	1.00	1.00	1.00	1.00	1.00
Pro	CCU	.40	.53	.50	.33	.12	Ara	CGU	.15	.08	.33	.24	.56
	CCC	.20	.07	.08	.09	.07		CGC	.00	.11	.17	.18	.36
	CCA		.20	.17	.18	.16	Arq	CGA	.23	.14	.07	.09	.03
	CCG	.40	.20	.25	.40	.65		CGG	.00	.17	.17	.11	.03
Thr	ACU	.11	.10	.10	.17	.25	Ser	AGU	.07	.07	.31	.10	.06
	ACC	.00	.14	.10	.14	.50		AGC	.07	.26	.13	.23	.26
	ACA	.56	.29	.30	.42	.07		AGA	.54	.28	.27	.28	.01
	ACG	.33	.47	.50	.27	.18		AGG	.08	.22	.00	.10	.01
Ala	GCU	.50	.37	.26	.26	.26	Glv	GGU	.10	.14	.53	.21	.48
Ala		.20	.26	.35	.19	.21		GGC	.30	.35	.32	.33	. 39
	GCA	.30	.22	.17	.31	.22		GGA	.40	.37	.16	.32	.05
Ala		.00	.15	.22	.24	.31		GGG	.20	.14	.00	.14	.08
	303							303					

Table 2. Codon Usage of B.subtilis Sigma-43 Operon*

* 1) B.s. codon usage frequency of average <u>B. subtilis</u> proteins, compiled from 10,919 codons of 35 sequenced genes.

 E.c. codon usage frequency of average <u>E. coli</u> proteins, from reference 24.

3) The number tabulated is the fraction usage of each codon compared with total for identical amino acid.

RNA polymerase major σ^{43} operon in the order <u>P23</u>, <u>dnaE</u> and <u>rpoD</u> and have physically and functionally located the promoter and terminator regions for the operon.

<u>Codon</u> <u>Usage</u>

It has been well established in <u>E. coli</u> that there is a correlation between expression level of a gene and its codon usage pattern. The more highly expressed genes show a very nonrandom pattern of codon usage, utilizing a restricted set of codons which are recognized by major species of isoacceptor tRNAs, while genes which are expressed at very low levels show an almost random pattern of codon usage (22,23). The analysis of this kind of correlation in <u>B. subtilis</u> has been limited due to the small number of sequenced genes, and the lack of knowledge concerning the expression levels of these genes in vivo. Recently, the rapid advance of cloning and sequencing of B. subtilis genes have allowed us to compile a codon usage table for average B. subtilis proteins and compare this with the codon usage of each gene in the σ^{43} operon, and also with that of average E. coli proteins (24) (see Table 2). Comparison of the codon usages between B. subtilis and E. coli led us to the general conclusion that B. subtilis tends to more evenly or randomly distribute the codons for its amino acids than E. coli. Nevertheless, the rarely used codons in E. coli, CUA (Leu), AUA (Ile), CCC (Pro), AGG (Arg) and GGG (Gly), were also used least in B. subtilis, although the bias is not as dramatic as that in E. coli. When the usage frequencies of codons AUA, AGG and GGG in the three genes of the operon were carefully examined, striking differences were found. In rpoD, a relatively highly expressed gene in B. subtilis [2,000-10,000 molecules/cell during growth (6)] just as its counterpart in E. coli (10), these codons were not used at all, while in <u>P23</u> and <u>dnaE</u> they were used quite frequently. Especially in <u>dnaE</u>, the usage frequencies for codons AUA and AGG were 0.26 and 0.22, which was much higher than those for the average B. subtilis proteins, which were 0.10 and 0.10, respectively. This preliminary analysis suggests to us that P23 and <u>dnaE</u> are expressed at lower levels than <u>rpoD</u>, which was not unexpected for <u>dnaE</u> since the DNA primase is required only in small amounts during DNA replication (25). Also consistent with this idea were the relative strengths of the ribosome binding sites for the three genes, which were found in an increasing order of $\triangle G' = -9.2$ kcal/mol for <u>P23</u>, -13.8 kcal/mol for <u>dnaE</u> and -18.8 kcal/mol for rpoD.

Since we do not know the function of P23, we can only speculate that it may have some role in translation or act as a regulatory protein present in relatively low concentrations in the cell.

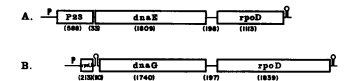


Figure 3. Schematic representation of the organization of <u>B. subtilis</u> (A) and <u>E. coli</u> (B) major σ operons. The number of base pairs is shown in parentheses for each structural unit. The organization of the <u>E. coli</u> σ^{70} operon is based on results from Burton <u>et al.</u> (10) and Lupski <u>et al.</u> (26). For simplicity, internal promoters and RNA processing sites are not shown.

Operon Organization

A comparison of the two major σ operons from <u>B.</u> subtilis and E. coli is illustrated in Fig. 3. Both operons contain three genes including the DNA primase and major σ genes, which are ordered in the same way. The operons are all under control of multiple promoters, indicating a complex transcription regulation system. However, significant differences do exist between them, of which the most striking is the first gene in the operon. P23 is more than twice as large as S21, and there is no sequence homology at all between them, while the middle gene products (DNA primases) are 31% homologous (8), and the last gene products (major σ factors) more than 50% homologous (6). Also different are the sizes of the intercistronic regions between the first and second genes, 33 bp in **B.** subtilis and 110 bp in E. coli, while that between the second and third genes are very similar in size (8,10). In E. coli, there is a mRNA processing site immediately following the dnaG gene (10), which was not found in B. subtilis.

DISCUSSION

We determined the nucleotide sequence of a 5 kb fragment in the <u>dnaE-rpoD</u> region of the <u>B. subtilis</u> chromosome and found three open reading frames transcribed in the same counterclockwise direction, two of which were identified as genes for the DNA primase and RNA polymerase major σ^{43} factor. The function of the first gene is still unknown. Discovery of the promoters in front of <u>P23</u> and a terminator 3' to <u>rpoD</u>, and the absence of promoters in the intercistronic regions between these genes provided strong evidence that <u>P23</u>, <u>dnaE</u>, and <u>rpoD</u> comprised a three gene operon. The operon was named RNA polymerase major sigma (σ^{43}) operon in analogy to that in <u>E. coli</u>.

The structure and organization of the σ^{43} operon resemble those of the <u>E. coli</u> σ^{70} operon except for the first gene. The size of P23 and its lack of amino acid sequence homology with S21 represent the most significant differences between the two operons at the molecular level. Since a reasonably high degree of homology exists between the DNA primase and σ genes, one might have expected some homology between the first genes <u>P23</u> and <u>rpsU</u>. We are currently attempting to identify P23 by use of immunological and cell fractionation techniques to see whether P23 might be associated with the ribosome fraction of <u>B.</u> <u>subtilis</u>.

The regulation of the <u>E. coli</u> σ^{70} operon is very complex since it is an important operon controlling not only translation, but also DNA replication and transcription (10,26). One of its interesting features is the control mechanism(s) to keep the expression of the <u>dnaG</u> gene lower than its adjacent genes, <u>rpsU</u> and <u>rpoD</u>. At least four mechanisms have been proposed including an internal terminator between the first and second genes (10), a weak ribosomal binding site for dnaG (10), frequent use of rare codons (27), and a mRNA processing site between the second and third genes (10). Although no experimental data are available concerning the expression level of <u>dnaE in vivo</u>, a low expression is expected from its function, and its counterpart in E. coli. The results of our codon usage analysis and the comparison of ribosomal binding sites are also in good agreement with the notion that <u>dnaE</u> represents a weakly expressed gene, and rpoD a fairly highly expressed gene. However, besides the possible regulatory mechanisms at the translational level, it is very likely that there are also control mechanism(s) involved at the transcriptional level.

In general, <u>B. subtilis</u> requires a more stringent Shine-Dalgarno sequence for gene expression than <u>E. coli</u> (28,29). The calculated free energies of interaction of the Shine-Dalgarno regions of <u>B. subtilis</u> mRNAs with the 3' end of its 16s rRNA have an average value of -17 kcal/mol (30), contrasted with an

average of -11 kcal/mol for that in E. coli (31). However, the calculated free energy value for P23 gene, -9.2 kcal/mol, is far below that of the average for **B. subtilis.** Considering this and the codon analysis data, it is tempting to propose that P23 encodes a regulatory protein which is weakly expressed, but physiologically important. Also, since <u>B. subtilis</u> cells undergo differentiation and can form spores, it is possible that the cell may have evolved a unique regulation system that is absent in E. coli, and that P23 may be one of the members in that system. The possibility also exists that P23 encodes an unidentified component of the B. subtilis translation machinery which is absent or has not been identified as yet in E. coli, since it has been reported that sequences other than the Shine-Dalgarno region can affect the translation efficiency of a gene (29,32); it thus is possible that P23 might still be expressed efficiently in vivo. More experimental data are required before we can say anything conclusive about this cryptic gene.

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