Improvement of α-Amylase Production by Modulation of Ribosomal Component Protein S12 in *Bacillus subtilis* 168

Kazuhiko Kurosawa, Takeshi Hosaka, Norimasa Tamehiro, Takashi Inaoka, and Kozo Ochi*

National Food Research Institute, Tsukuba, Ibaraki 305-8642, Japan

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The capacity of ribosomal modification to improve antibiotic production by Streptomyces spp. has already been demonstrated. Here we show that introduction of mutations that produce streptomycin resistance (str) also enhances α-amylase (and protease) production by a strain of *Bacillus subtilis* as estimated by measuring the enzyme activity. The str mutations are point mutations within rpsL, the gene encoding the ribosomal protein S12. In vivo as well as in vitro poly(U)-directed cell-free translation systems showed that among the various rpsL mutations K56R (which corresponds to position 42 in *E. coli*) was particularly effective at enhancing α-amylase production. Cells harboring the K56R mutant ribosome exhibited enhanced translational activity during the stationary phase of cell growth. In addition, the K56R mutant ribosome exhibited increased 70S complex stability in the presence of low Mg\(^{2+}\) concentrations. We therefore conclude that the observed increase in protein synthesis activity by the K56R mutant ribosome reflects increased stability of the 70S complex and is responsible for the increase in α-amylase production seen in the affected strain.

Of the various starch-hydrolyzing enzymes, the α-amylases (1,4-α-D-glucan glucanohydrolase; EC 3.2.1.1) are of particular importance, as they are responsible for the solubilization of starch. As such, these enzymes are currently among the most widely utilized in biotechnology. Although α-amylases can be derived from plants and animals, it is the enzymes from microbial sources (typically *Bacillus* spp.) that are generally used to meet the expanding industrial demands. In addition to well-established applications in starch saccharification and in the textile, food, brewing, and distilling industries, bacterial α-amylases are now also used in areas of clinical, medicinal, and analytical chemistry (for a review, see reference 23). Contributing to the appeal of bacterial α-amylases is their high degree of optimization (e.g., highly thermostable or alkali tolerant), conferred by means of protein engineering or discovered through investigation of novel microorganisms (7; also reviewed in references 2 and 19).

α-Amylase is an end-type enzyme that hydrolyzes α-1,4-glucosidic linkages from starch and various other types of oligosaccharides, and several of the enzymes from *Bacillus subtilis* have been cloned, sequenced, and subjected to three-dimensional structural analysis (5, 18). The production of extracellular α-amylose by *B. subtilis* is known to be controlled by several genes, including *amyR* and *pap*, and to vary with changes in the environmental conditions as well as with changes in the structure and function of the cell envelope (1). Mutagenesis induced by chemicals such as N-methyl-N'-nitro-N-nitrosoguanidine or by UV radiation has been employed to obtain hyperproducing strains, in which α-amylose synthesis can often be doubled or even tripled (32; for a review, see reference 23). Current developments in gene engineering also make possible improvements in amylase production through molecular breeding (13).

We previously showed that a certain streptomycin resistance-producing mutation (str) in *rpsL*, which encodes the ribosomal protein S12, also gives rise to antibiotic production in *Streptomyces lividans* and *Streptomyces coelicolor* (8, 26). Later, we used other bacterial genera to demonstrate that introducing a specific str mutation together with a gentamicin resistance-producing mutation (gen) gives rise to a marked increase in antibiotic production and to shed light on the mechanism by which antibiotic is produced in *S. coelicolor* (10). It was also demonstrated that by introducing various combinations of drug resistance-producing mutations, we could increase the production of an antibiotic in a stepwise manner (11).

The finding that certain *rpsL* mutations induce dramatic activation of antibiotic production prompted us to hypothesize that bacterial gene expression may be altered dramatically by modifying ribosomal proteins or rRNA. Thus, our ultimate aim has been to develop “ribosome engineering” (20) as a rational approach to taking full advantage of bacterial capabilities. Because certain extracellular enzymes (such as α-amylase and protease) are known to be produced during the late growth phase, it seemed plausible that synthesis of these enzymes might be enhanced by introducing certain streptomycin resistance-producing (*rpsL*) mutations. The aim of the present study, therefore, was to assess the efficacy with which ribosome engineering could be used to enhance enzyme synthesis in *B. subtilis* by examining its effect on α-amylase (and protease) production.

**MATERIALS AND METHODS**

**Bacterial strains and plasmids.** The bacterial strains and plasmids used in this study are listed in Table 1. *Bacillus subtilis rpsL* mutants (WL1, WL2, WL3, WL4, WL5, and WL9) were all derived from strain 168 (10, 14), which is a standard (Marburg) strain frequently used for studying sporulation. Mutants WL6 and WL15, which contain a K56H and a K101E substitution, respectively, were constructed by site-directed mutagenesis using the plasmid pKF19k-*rpsL* as a
template (14). The oligonucleotides 5′-AGTTCGTTTGTGGTCTAGTGT-3′ (for K56H) and 5′-GCTAGGTCTGTACACGCCTCT-3′ (for K101E), which include the mutation sites (underlined), were used to generate rpsL6 and rpsL15. The mutant WL10 (G105W) was prepared using PCR random mutagenesis on the basis of this mutant’s ability to resist streptomycin, as described previously (14). Detection of the rpsL mutations was accomplished using PCR, the products of which were directly sequenced using a sequence analyzer (14).

**Media and growth conditions.** NG medium, which was originally developed for antibiotic production by *B. subtilis* (10), was used for α-amylase production. It contained (per liter) 10 g of nutrient broth (Difco), 10 g of glucose, 2 g of NaCl, 5 mg of CuSO₄·5H₂O, 7.5 mg of FeSO₄·7H₂O, 3.6 mg of MnSO₄·5H₂O, 15 mg of CaCl₂·2H₂O, 9 mg of ZnSO₄·7H₂O, and 50 mg of tryptophan, as required (adjusted to pH 7.2 with NaOH). Strains were initially grown for 12 h in NG medium (10 ml in 100-ml flasks) at 37°C. Thereafter, aliquots (0.1 ml) of the culture broth were inoculated into 10 ml of NG medium in 100-ml flasks and cultured for the indicated times on a rotary shaker (200 rpm) at 45°C (instead of 37°C) for α-amylase production, except that cells were grown in PM medium, which contained (per liter) 5 g of peptone (Difco), 5 g of malt extract (Difco), 5 g of NaCl, 1 g of glucose, and 50 mg of tryptophan, as required (adjusted to pH 7.2 with NaOH).

**Assay for α-amylase and protein.** α-Amylase activity was assayed using the method of Fowa (6). A sample (0.2 ml) of 0.5% soluble starch in 0.05 M phosphate buffer (pH 6.0) was mixed with 0.1 ml of enzyme solution. After incubation for 15 to 45 min at 40°C, a 20-μl aliquot of the reaction mixture was added to 0.5 ml of 0.2 M Li₂K₁ solution, and the optical density at 700 nm was measured in a spectrophotometer. One unit of enzyme was defined as the amount necessary to hydrolyze 0.1 mg of soluble starch in 1 min. Protein activity was assayed as described by Shimizu et al. (27). One unit of enzyme was defined as the amount of the enzyme which solubilized a 1-μg equivalent of tyrosine in 1 min.

**Incorporation of [3H]leucine.** Strains were grown for various periods in diluted (1/8) NG medium, after which [3H]leucine (0.2 μCi, 200 μCi) was added to 10-ml samples of culture and incubated for 0, 10, 20, or 30 min. One-milliliter aliquots were then collected, mixed with 1 ml of cold 10% (wt/vol) trichloroacetic acid, and kept on ice for 30 min to precipitate the protein from the solution. The precipitated protein was collected by filtration on a nitrocellulose filter (pore size, 0.45 μm) and washed with 10 ml of 5% (vol/vol) trichloroacetic acid. The filters (containing protein) were then dried and their radioactivity measured using a liquid scintillation counter.

**In vitro translation assay.** poly(U)-directed cell-free synthesis of polyphenylalanine was carried out as described by Legault-Demare and Chamblis (16) with slight modifications. *Bacillus subtilis* cells grown to various growth phases in NG medium were collected by centrifugation and washed with standard buffer (10 mM Tris-HCl [pH 7.7], 10 mM magnesium acetate, 30 mM ammonium acetate, and 6 mM 2-mercaptoethanol) containing 2 mM phenylmethylsulfonyl fluoride. The cell paste was broken by grinding with aluminum oxide powder (2 g for each 1 g of cell paste; Wako) for 10 min, after which the ground paste was suspended in standard buffer containing 2 mM phenylmethylsulfonyl fluoride plus 10% (wt/vol) glycerol. The resultant lysate was treated with RNase-free DNase I (10 U/ml; Takara) for 10 min on ice and then centrifuged at 30,000 × g for 30 min to remove cell debris. The resultant supernatant was further fractionated into the S-150 fraction (supernatant) and the ribosomes (precipitant) by centrifugation at 150,000 × g for 3 h. The ribosomes were washed once more with standard buffer plus 10% (wt/vol) glycerol, after which both the ribosomes and the S-150 fraction were dialyzed against 60 volumes of standard buffer plus 10% (wt/vol) glycerol for 6 h, divided into small aliquots, frozen in liquid nitrogen, and stored at −80°C until use. The reaction mixture for polyphenylalanine synthesis (100 μl) consisted of 55 mM HEPES-KOH (pH 7.5), 1 mM dithiothreitol, 210 mM potassium acetate, 27.5 mM ammonium acetate, 10.7 mM magnesium acetate, 88 μM l-phenylalanine, 5 mM spermidine, 1.2 mM GTP, 0.8 mM GTP, 0.64 mM 3′,5′-cyclic AMP, 80 mM creatine phosphate, 0.25 mg/ml creatine kinase, 200 units/ml RNase inhibitor (recombinant solution; Wako), 0.45 mg/ml Escherichia coli total tRNA, 0.4 mM concentrations of each tRNA, and 10% (vol/vol) of ribosome fraction. The reaction was initiated by adding 75 μg of poly(U), after which the mixture was incubated at 37°C for the appropriate time; 1 ml of 10% (wt/vol) trichloroacetic acid was then added to stop the reaction, after which the mixtures were boiled for 15 min. Precipitated proteins were collected on nitrocellulose filters, and the incorporation of [3H]phenylalanine into the acid-insoluble fraction was determined using a liquid scintillation counter. To reduce levels of endogenous mRNA, ribosomes plus the S-150 were preincubated just prior to use at 37°C for 10 min in the reaction mixture as described above without the amino acids, energy-generating reagents, or poly(U).

**Sucrose density gradient centrifugation of ribosomes.** Crude ribosomes prepared from mid-exponential-phase cells (see above) were precipitated by centrifugation at 150,000 × g for 3 h and resuspended in standard buffer containing the specified concentration of magnesium acetate. The ribosomes were then laid onto a 10% to 30% (wt/vol) linear sucrose gradient prepared in buffer containing the same concentration of magnesium acetate and centrifuged in a SW41Ti rotor at 38,000 rpm for 4 h at 4°C. The profiles of the ribosomes were observed at 254 nm using an ATTO Bio-Mini UV Monitor equipped with a Biocomp Piston Gradient Fractionator (Towa Kagaku).

**RESULTS**

**Isolation of *B. subtilis* rpsL mutants.** We previously isolated various rpsL mutants that developed spontaneously on plates containing streptomycin (10, 14). In addition, to examine an even wider variety of rpsL mutations with the aim of improving α-amylase production, we also prepared three additional rpsL

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**TABLE 1. Bacterial strains and plasmids used in this study**

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Genotype and/or phenotype</th>
<th>Amino acid exchange in ribosome protein S12</th>
<th>Source and/or reference</th>
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<td><em>Bacillus subtilis</em> strains</td>
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<td>168</td>
<td>trpC2</td>
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<tr>
<td>KO272</td>
<td>trpC2-stryr(5, rpsL) Sm</td>
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<tr>
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<td>pKF19k-rpsL containing the rpsL (K101E) mutation</td>
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<td>This study</td>
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* The strain to the right of the arrow was transformed with the chromosomal DNA, PCR product, or plasmid to the left of the arrow.
mutants, WL6 (K56H), WL10 (G105W), and WL15 (K101E), which are known to confer streptomycin resistance in E. coli (30) (Table 1). Although strains WL4 and WL5 grew at a slightly slower rate than parental strain 168, all of the other strains harboring rpsL mutations grew as well as strain 168 and sporulated well, producing about 10^9 spores/ml in sporulation medium. Strains WL1, WL2, WL3, WL4, WL5, and WL6 all exhibited a high degree of resistance to streptomycin, tolerating levels of >500 μg/ml, while strains WL9, WL10, and WL15 showed an intermediate resistance, tolerating levels of 50 to 100 μg/ml. By contrast, strain 168 tolerated streptomycin at 1/100 g/ml. The rpsL mutations in B. subtilis, together with the previously reported rpsL mutations in E. coli and S. coelicolor, are illustrated in Fig. 1.

Enhanced production of α-amylase in some rpsL mutants. We first monitored growth and α-amylase production in strain 168 (Fig. 2A). In NG medium, the production of α-amylase commenced during the stationary phase (24 h) and reached a maximum by 54 h. We next examined the capacity of rpsL mutation to enhance α-amylase production using the rpsL mutation mutant strains listed in Table 1. It was found that the K56R mutation led to the greatest increase in α-amylase production but that increases were also seen with the K101E, K56I, and P104S mutations (Fig. 3). The other mutations (G105W, K56T, K56I, and K56Q) were without effect. The K56R mutant ribosome, was about 40% greater than that seen with the parental strain. It is notable that the K56R mutation was effective also in improving protease productivity (30% greater than that of the parental strain) (Fig. 2B). Thus, certain rpsL mutations effectively improved the production of extracellular enzymes, α-amylase, and protease.

The WL1 rpsL mutant strain exhibits enhanced protein synthesis activity. Recent work in our laboratory revealed that cells harboring the K88E (in S. lividans and E. coli) or K88R (in Streptomyces albus) rpsL mutation exhibited enhanced protein synthesis during the stationary phase, as determined by both in vivo and in vitro translation assays (9, 22, 29). We therefore reasoned that the enhanced α-amylase production by B. subtilis strain WL1 harboring the K56R rpsL mutation might reflect a similar effect on protein synthesis. To test that idea, we first monitored the abilities of wild-type and mutant cells to synthesize protein in vivo. Strains 168, WL1, and WL2, which served as a reference strain, were grown to various growth phases in diluted (1/8) NG medium (to promote the incorporation of [3H]leucine), after which [3H]leucine was added and the cells were incubated for an additional 30 min. In parental strain 168, protein synthesis was maximal during the mid-exponential phase (2 h) and then declined sharply (by 13-fold) once cells entered the transition or stationary phase (Fig. 4). By contrast, strain WL1 (K56R mutant) sustained a higher level of protein synthesis during the stationary phase than was seen with strain 168: protein synthesis by strain WL1 was only reduced eightfold during the stationary phase. Strain WL2 (K56N mutant), which did not show enhanced α-amylase production, also did not show enhanced protein synthesis during the stationary phase (Fig. 4).

K56R mutant ribosomes exhibit enhanced translational activity. To confirm the results of the in vivo experiments, we measured the in vitro translational activity of ribosomes isolated from wild-type and mutant cells at various growth stages using a poly(U)-directed cell-free translation system (polypeptide synthesis) (see Materials and Methods). For ribosomes isolated from wild-type (strain 168) cells, the rate of polypeptide synthesis was maximal when the organelles were extracted during the mid-exponential phase and was markedly lower when they were extracted during the stationary phase (Fig. 5). Notably, K56R mutant ribosomes isolated from strain WL1 cells during the stationary phase (36 h) exhibited twice as much activity as those from the wild-type strain. No such increase in ribosomal activity was seen with the strain WL2 K56N mutant, indicating the enhanced translational activity during late growth was a specific characteristic of the K56R mutant ribosome.

K56R mutant ribosomes form more stable 70S complexes. Starvation for an essential amino acid increases the spatial separation between the two ribosomal subunits (21, 34). Indeed, ribosomes from methionine-starved cells are reported to be structurally unstable in the presence of low concentrations of Mg^{2+}, which is indicative of the structural instability of open-form ribosomes (25). We therefore reasoned that K56R mutant ribosomes from strain WL1, which actively synthesize protein even under starvation conditions, such as those encountered during the late stationary phase, may maintain more stable intersubunit interactions. To test this hypothesis, we prepared ribosomes from wild-type and mutant strains grown to mid-exponential phase in NG medium and examined the stability of the 70S complex in the presence of selected con-
centrations of Mg$^{2+}$. At a high Mg$^{2+}$ concentration (10 mM), almost all ribosomes were recovered in the 70S form (Fig. 6). However, dissociation of 70S ribosomes into the 30S and 50S subunits, along with formation of an intermediate (detected as a peak between the 50S and 70S peaks), occurred when the Mg$^{2+}$ concentration was reduced to 5 mM or less. In that regard, a notably larger fraction (about 1.5-fold size) of the K56R mutant ribosomes from strain WL1 remained in the 70S form in the presence of low concentrations (5, 2, and 1 mM) of Mg$^{2+}$, as confirmed by four independent experiments. A typical result at 5 mM and 1 mM Mg$^{2+}$ concentrations is presented as Fig. 6. The K56N mutant ribosomes from strain WL2 displayed a pattern similar to that of strain 168 (data not shown). These findings suggest that the K56R (but not K56N)
ribosomes are structurally more stable under stressful conditions, which could account for the enhanced protein synthesis observed in strain WL1.

**DISCUSSION**

Working with *B. subtilis rpsL* mutations, we have been able to demonstrate the capacity of several (especially K56R) to improve production of α-amylase and protease. Although the observed increase of α-amylase or protease production was 1.3- to 1.4-fold, these values seem to be significant, since the increase of enzyme production caused by a single mutation is in general at most twofold in either wild-type strains and industrial strains that had been bred to produce a high level of enzymes (32; S. Itoh, personal communication). The present results show (i) that cells harboring the K56R mutant ribosome can sustain a higher level of gross protein synthetic activity than wild-type cells during the late growth phase, as determined by measuring the incorporation of \[^{3}H\]leucine, (ii) that K56R mutant ribosomes isolated during the late stationary phase have a higher capacity for translating synthetic polynucleotide [poly(U)] than wild-type ribosomes, and (iii) that the mutant ribosomes are structurally more stable than wild-type ribosomes under stressful conditions. It is thus concluded that the enhanced protein synthesis seen with cells harboring the K56R mutant ribosome is likely due, at least in part, to the increased stability of the 70S particle. This proposal is consistent with earlier findings that the *Streptomyces* K88E and the *E. coli* K87E ribosomal mutants also show enhanced protein synthesis during the late phases of growth (9, 22, 29). Although much progress has been made toward increasing the productivity of amylase-producing strains (23), the present method is characterized by the host cell's amenability (generation of spontaneous drug resistance-producing mutations) and is thus applicable to a number of microorganisms (10, 11, 29). It should also be emphasized that *rpsL* mutations found in the present study caused no impairment of growth and sporulation under the conditions tested.

The effects of streptomycin on bacterial ribosomes have been studied in great detail (3, 4, 31). Among the numerous actions attributed to this drug, its ability to cause mRNA codons to be misread is the best characterized. In that regard, it is well known that S12 mutations that confer streptomycin resistance can increase the accuracy of protein synthesis, which was the case with *B. subtilis rpsL* mutants (14). However, working with *S. coelicolor* and *S. lividans*, it became apparent that the increased accuracy of the protein synthesis by *rpsL* mutants is unrelated to the increased production of antibiotic (22). It is therefore noteworthy that, in the present study, strain WL1 harboring the K56R mutant ribosome exhibited a higher degree of translational activity during the stationary phase (Fig. 4 and 5). This increased activity is likely the result of a more stable ribosomal structure (Fig. 6) and could explain why strain WL1 is capable of producing greater amounts of α-amylase and protease than the wild-type strain. Moreover, production of secondary metabolites, including α-amylase and protease, usually commences during the late growth phase (i.e., the transition or stationary phase); thus, the enhanced protein synthesis seen at that time would be expected to include increased production of these extracellular enzymes. Consistent with that idea, strain WL2 harboring the K56N mutant ribosome, which showed no increase in α-amylase production, did not exhibit increased translational activity during the stationary phase.
(Fig. 4 and 5). In *E. coli*, the rate of protein synthesis is reportedly reduced by more than 90% when cells are starved for amino acids (28). That the protein synthetic activity of strain WL1 persisted despite such conditions distinguishes it from previously studied wild-type strains.

The ribosomal protein S12 is a component of the 30S subunit in bacteria. Best characterized is its role in determining the efficiency with which cognate tRNAs are selected, which contributes to the accuracy achieved when decoding mRNA sequences (3, 14, 15). Recent structural analyses of the 50S and 30S subunits, as well as the intact 70S ribosome, have greatly advanced our understanding of protein synthesis (24, 33). All of the Smt mutations found in the S12 protein are limited to two conserved regions: region 1, spanning amino acid residues 55 to 59 (in numbering system for *B. subtilis*), and region 2, spanning residues 101 to 107. Both of these regions comprise loop structures in the S12 protein, with region 1 projecting into the space between the 530 loop and the 1492 to 1493 strand of the decoding site (17). Most mutations within region 1 lead to a hyperaccurate phenotype (15), though they can weaken the interactions between the tRNA-mRNA complex and the 30S A site (33). Although it is difficult at present to explain how the K56R mutation in the S12 protein mediates ribosomal stabilization, it is possible that it affects interactions between the 16S and 23S rRNAs.

Our previous studies have shown that certain mutations within S12 confer resistance to streptomycin and enhance or activate antibiotic production (reviewed in reference 20). It is worth mentioning that the K56R S12 mutation found to enhance α-amylase production in *B. subtilis* also very effectively enhanced antibiotic production (80-fold) of this organism (10). Novel α-amylase that is highly resistant to chelating reagents and chemical oxidants from the alkaliphilic *Bacillus* isolate KSM-3K8. Appl. Environ. Microbiol. 67:1744–1750.


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