The formation and degradation of α-peptide bonds are basic functions in living organisms. Both activities are indispensable for cells to proliferate, survive, and adapt to environmental change. Peptide degradation depends on a number of enzymes, such as proteases and peptidases, many of which have broad substrate specificities. By contrast, only one system, the ribosome system, is responsible for the formation of peptide bonds. All cellular organisms are equipped with the ribosome system, in which an amino acid is first ligated to the corresponding tRNA by aminoacyl-tRNA synthetase via aminoacyl-AMP, an active intermediate, and next transferred to an elongating peptide by the ribosome (21). The mRNA encodes the amino acid sequence of the peptide to be synthesized; thus this system can form any peptide depending on the information in the mRNA.

While the ribosome system is universal, other machineries specific for certain peptides are also known, including nonribosomal peptide synthetase (NRPS) (4, 5, 26), folypolyglutamate synthase (23), and cyanophycin synthetase (1), D-alanine:D-alanine ligase (DDL) (2). All of these enzymes are categorized into an ATP-dependent carboxylate-amine/thiol grasp motif and take acyl-phosphate as an active intermediate. The latter synthesizes polymers composed of aspartate and arginine in Escherichia coli (23). Peptidoglycan biosynthetic enzymes (in Cyanobacteriaciae (1)). All of these enzymes are categorized into an ATP-dependent carboxylate-amine/thiol ligase superfamily (9, 20), whose members contain an ATP-grasp motif and take acyl-phosphate as an active intermediate.

Despite the versatility of these peptide-forming activities, an enzyme catalyzing the synthesis of α-dipeptide from l-amino acids has not been identified. The enzymes referred to above synthesize a polymer, oligomer, dipeptide of D-amino acids, or a dipeptide with a γ-peptide bond but do not form a dipeptide of t-amino acids with an α-peptide bond. A few biochemical studies with partially purified cell extracts have suggested the existence of a dipeptide-synthesizing enzyme (18, 24), but no gene or purified enzyme for the activity has been reported. We were interested in this point, especially in the curious fact that the ATP-dependent carboxylate-amine/thiol ligase superfamily does not contain such activity though DDL is a member of the family (7). No reason can be given for the absence of an enzyme synthesizing dipeptides from t-amino acids because the ligation reaction between t-amino acids seems to be possible via the same mechanism as that of DDL. We attempted to find such a novel enzyme.

The ATP-dependent carboxylate-amine/thiol ligase superfamily is known to contain enzymes catalyzing the formation of various types of peptide, such as D-alanyl-D-alanine, polyglutamate, and γ-peptide, but, curiously, no enzyme synthesizing α-dipeptides of l-amino acids is known. We attempted to find such an enzyme. By in silico screening based on the consensus sequence of the superfamily followed by an in vitro assay with purified enzyme to avoid the degradation of the peptide(s) synthesized, ywfE of Bacillus subtilis was found to code for the activity forming l-alanyl-l-glutamine from l-alanine and l-glutamine with hydrolysis of ATP to ADP. No AMP was formed, supporting the idea that the enzyme belongs to the superfamily. Surprisingly, the enzyme accepted a wide variety of l-amino acids. Among 231 combinations of l-amino acids tested, reaction products were obtained for 111 combinations and 44 kinds of α-dipeptides were confirmed by high-performance liquid chromatography analyses, while no tripeptide or longer peptide was detected and the D-amino acids were inert. From these results, we propose that ywfE encodes a new member of the superfamily, l-amino acid ligase.
to find this enzyme. In this report, we describe the screening and finding of an enzyme synthesizing dipeptides from L-amino acids. We also present some of the characteristics of the enzyme.

MATERIALS AND METHODS

Materials. Escherichia coli strain DH5α was purchased from Takara (Kyoto, Japan). Bacillus subtilis (ATCC 15245) was obtained from the American Type Culture Collection. Plasmids pQE-60, pET-2a, and pREP4 were purchased from QiAGEN (Hilden, Germany). Restriction and modification enzymes, DNA ligase, and the PCR kit (LA PCR) were purchased from Takara. His Trap was obtained from Amersham Pharmacia Biotech (Buckinghamshire, England). Dipeptides for standards were purchased from Sigma-Aldrich (St. Louis, MO) or Bachem Feinchemikalien AG (Bubendorf, Switzerland).

Genetic manipulation. DNA manipulations were basically performed according to Sambrook et al. (19). The ywfE gene was amplified from the B. subtilis (ATCC 15245) genome by PCR using the primer 5′-TTAACATGTTAGGAGAAAACAGTATTG-3′ to introduce an NcoI site and 5′-ATATGGATTCTACGTGCAGACCATCCTTG-3′ to introduce a BamHI site. The 1.4-kb PCR product was ligated with NcoI and BamHI and inserted between the NcoI and BamHI sites of pQE-60. The resulting plasmid, designated as ywfE with a histidine (His) tag at its C terminal under the control of the T5 promoter and lac operator, was named pQE60ywfE.

In silico screening. Searches for proteins containing an ATP-grasp motif and homology searches were conducted by using functions provided by EzPasy (expert protein analysis system) available at http://kr.expasy.org/. The Blast search was done with the blastp program against the complete database of the EzPasy server.

Biochemical procedures. DH5α/pQE60ywfE cells cultivated in Luria-Bertani medium (0.5 g/liter Bacto tryptone, 5 g/liter yeast extract, and 5 g/liter NaCl) containing 50 mg/liter of ampicillin at 30°C for 16 h were transferred to TB medium (0.25 g/liter Bacto tryptone, 25 g/liter yeast extract, 4 ml/liter glycerol, 17 mM KH2PO4, 72 mM K2HPO4) and incubated at 30°C with vigorous shaking. Three hours after the transfer, 1 mM of IPTG (isopropyl-β-D-thiogalactopyranoside; final concentration) was added, and cultivation was continued a further 4 h. Cells were harvested by centrifugation, resuspended with buffer A (100 mM of Tris-HCl [pH 8.0]) and disrupted by sonication at 4°C. Cellular debris was removed by centrifugation (12,000 × g for 40 min), and the supernatant was subjected to further purification using His Trap (Amersham Pharmacia Biotech) following the instructions of the manufacturer.

The dipeptide-synthesizing activity of YwfE protein was assayed as follows unless otherwise specified. The standard reaction mixture (total volume was 1 ml) contained 50 mg/liter of purified His-tagged YwfE protein, 60 mM ATP, 30 mM Mops buffer, and 30 mM substrate amino acid(s) in 50 mM Tris-HCl buffer (pH 9.0). The reaction was carried out at 37°C. A portion of the reaction mixture was taken at each sampling time and immediately subjected to derivatization with fluorescamine/methyl chloroformate (FMOC-Cl) (see “Analyses”). For matrix-assisted laser desorption ionization–time of flight mass spectrometry (MALDI-TOFMS) analysis, the reaction mixture was diluted 50-fold with water and spread on a target plate coated with 2,5-dihydroxybenzoic acid (DHB) as a matrix. The plate was dried and loaded into a MALDI-TOF mass spectrometer (REFLEX mass spectrometer; Bruker Daltonics).

(ii) MALDI-TOFMS analysis. The reaction mixture was diluted 50-fold with water and spread on a target plate coated with 2,5-dihydroxybenzoic acid (DHB) as a matrix. The plate was dried and loaded into a MALDI-TOF mass spectrometer (REFLEX mass spectrometer; Bruker Daltonics).

Phosphate. The concentration of phosphate was determined using a spectrophotometric assay utilizing the formation of phosphomolybdate acid (10) with a kit (Phospha-C Test Wako; Wako, Osaka, Japan).

RESULTS

Screening of L-amino acid ligase. An activity-based screening of dipeptide-forming enzymes using living cells or cell extracts is practically impossible because all living organisms possess dipeptide-degrading activities. Therefore in silico screening was conducted. Three keys were set up to screen for enzymes catalyzing the formation of dipeptides from L-amino acids. First, the enzyme would have an ATP-grasp motif, which is a signature of the ATP-dependent carboxylate-amine/thiol ligase family. Second, the open reading frame of the gene would be unassigned; that is, the function of the gene would be unknown. Third, the enzyme would have slight homology with DDL. The screening was carried out using a public database and its functions.

Proteins containing an ATP-grasp domain (accession number PS 50975) in PROSITE (http://kr.expasy.org/prosite/) were nominated from the Swiss-Prot database (first key). More than 300 proteins were listed, but most were assigned a function: acetyl-coenzyme A (CoA) carboxylase, carbamoyl-phosphate synthase, DDL, phosphoribosylamine-glycine ligase, etc. After eliminating the proteins of known function (second key), 13 proteins remained. To examine their similarity to DDL (third key), a BLAST search was done using the amino acid sequence of each protein as a query. One protein, Y4SG, has strong homology with DDL. The screening was carried out using a public database and its functions.

Proteins containing an ATP-grasp domain (accession number PS 50975) in PROSITE (http://kr.expasy.org/prosite/) were nominated from the Swiss-Prot database (first key). More than 300 proteins were listed, but most were assigned a function: acetyl-coenzyme A (CoA) carboxylase, carbamoyl-phosphate synthase, DDL, phosphoribosylamine-glycine ligase, etc. After eliminating the proteins of known function (second key), 13 proteins remained. To examine their similarity to DDL (third key), a BLAST search was done using the amino acid sequence of each protein as a query. One protein, Y4SG, has strong homology with DDL and thus was judged not to meet the third key. The other 10 had no similarity to DDL (we set the limit of similarity at an E value of <e−3)); MG011, MG012, MJ0620, MJ1001, MPN015, and MPN016 showed high levels of similarity to ribosomal protein; MJ0776, MJ0136, and yfiQ had homology to carbamoyl-phosphate synthase, phosphoribosylamine-glycine ligase, and acetyl-CoA synthetase, respectively; and MJ0815 had no homology to a function-known protein. Only two sequences, ywfE (E value, 2e−4) and Y4RH (E value, 4e−5), remained. As the latter was assumed to be a homolog of biotin carboxylase due to its strong homology to carboxylases from various organisms (8), ywfE was chosen as the most promising candidate.

ywfE, an open reading frame in the Bacillus subtilis genome, has a clear ATP-grasp fold and encodes a protein with homology with DDL (Fig. 1). Its function was unknown, but it is part of the bacilysin biosynthetic gene cluster (12). Bacilysin is an
antibiotic dipeptide, L-alanyl-anticapsin, synthesized from Ala and an unusual amino acid, anticapsin, in the presence of ATP. The protein is encoded by the gene cluster though the exact function of each gene has not been elucidated (12, 18, 29).

Confirmation of YwfE protein activity. Since anticapsin, a substrate for bacilysin-synthesizing activity, was not available, other amino acids were used for checking the activity. Ala and Gln were selected, because L-alanyl-L-alanine (Ala-Ala) was used as the substrate for bacilysin-synthesizing activity, was not available, and an unusual amino acid, anticapsin, in the presence of ATP. As expected, the extracts had strong dipeptide-degrading activity (data not shown), suggesting the possibility that the synthesizing activity of the ywfE gene product was masked.

To exclude this possibility, an assay with purified protein was conducted. Cell extract of DH5α/pQE60ywfE was applied to a nickel column, and the gene product was eluted with imidazole buffer and then dialyzed with Tris-HCl buffer. Through these procedures, the gene product was purified to apparent homogeneity (Fig. 2).

This purified fraction was incubated with Ala, Gln, and ATP for 4 h and analyzed with MALDI-TOFMS and HPLC. In the MALDI-TOFMS analysis, peaks corresponding to [Ala-Ala + Na]+ (m/z 183) and a dipeptide(s) composed of Ala and Gln ([Ala-Gln + H]+ or [Gln-Ala + H]+, m/z 218; [Ala-Gln + Na]+ or [Gln-Ala + Na]+, m/z 240; [Ala-Gln (Na) + Na]+ or [Gln-Ala (Na) + Na]+, m/z 262) were detected (Fig. 3). No peak corresponding to tripeptide (for example, [Ala-Ala-Ala + Na]+, m/z 272, or [Ala-Gln-Ala + Na]+, m/z 307) was observed. While the MS analysis could not identify a dipeptide with Ala and Gln as Ala-Gln or Gln-Ala, HPLC analyses clearly revealed that dipeptides formed were Ala-Ala and Ala-Gln but not Gln-Ala (Fig. 4a to d). No unassigned product was found in the HPLC analyses, supporting the absence of tripeptides.

The time course of the reaction was monitored with HPLC (Fig. 5b). As the reaction proceeded, Ala and Gln levels decreased and Ala-Gln and Ala-Ala were found to be accumulated in the reaction mixture. After 12 h, 20.5 mM of Gln and 21 mM of Ala were consumed and 19 mM of Ala-Gln and 1.8 mM of Ala-Ala were formed. The amounts of amino acids used were almost equal to the amounts of dipeptides formed, indicating that the purified fraction contained no substantial dipeptide-degrading activity or glutaminase activity. At the end of the reaction, residual ATP (28 mM), ADP (30 mM), and phosphate (29 mM) were detected but not AMP. When ATP was omitted in the reaction, no dipeptide was detected. These results clearly demonstrated that YwfE protein catalyzes the ligation reaction of Ala and Gln with the decomposition of ATP to ADP.

Characterization of YwfE protein. Using Ala and Gln as substrates, some of the characteristics of the enzyme were examined. The optimal pH and temperature were around 9.5...
and 37°C, respectively (Fig. 6). GTP did not substitute for ATP. Mg or Mn was essential for the activities. Dipeptide formation was completely abolished by the addition of 2 mM of CdSO₄ or ZnSO₄ to the standard reaction mixture, indicating strong inhibitory effects of these metal ions. No coenzyme, such as NAD(P), NAD(P)H, pyridoxal phosphate, CoA, or dihydrofolate was required. Apparent \( K_m \)s for Ala, Gln, and ATP were 0.42 (±0.07), 105.0 (±17.9), and 0.07 (±0.1) mM, respectively. The \( V_{max} \) for Ala-Gln synthesis was 764.6 nmol · min⁻¹ · mg⁻¹ protein (Fig. 7).

Next, combinations of amino acids were tested as substrates under conditions of pH 9.0 and 37°C. The reaction mixture was analyzed with three methods: by measuring the amount of phosphate formed and amounts of ATP and ADP to check whether the reaction proceeds and by analyzing the dipeptide formed by HPLC to confirm the product. Surprisingly, it was found that YwfE protein accepted a wide variety of amino acids as substrates and various dipeptides were formed. Among all combinations of 21 kinds of L-amino acids tested (231 combinations in total), reaction products were obtained in 111 reactions (see examples in Fig. 4). In some reactions, more than two dipeptides were formed. By comparing the dipeptide standard sample in the HPLC analysis, 44 kinds of dipeptides were confirmed to be produced (Fig. 8).

While the enzyme has extremely broad substrate specificity, some preferences were observed; the enzyme did not accept highly charged amino acids such as Lys, Arg, Gln, and Asp; the enzyme did not react with secondary amines such as Pro; the N-terminal residue of the dipeptide formed seemed to be limited to Ala, Gly, Ser, Thr, and Met whereas the C-terminal residue seemed to allow for a wider array of amino acids; and Ala and Ser seemed to be most preferred for the N-terminal residue while Met and Phe seemed to be preferred for the C terminus. The enzyme did not react with D-Ala, D-serine, or D-phenylalanine, confirming that it was not DDL. From these results, it was revealed that YwfE protein had the ability to synthesize various dipeptides from corresponding combinations of L-amino acids in the presence of ATP.

**DISCUSSION**

We report here that the protein encoded by ywfE, which was found in the \( B. subtilis \) genome by in silico screening, catalyzes the formation of \( \alpha \)-dipeptides from L-amino acids in an ATP-dependent manner. This is the first gene for an enzyme synthesizing dipeptides from L-amino acids. The most outstanding feature of the enzyme is its broad substrate specificity. Since all other peptide bond-forming enzymes except for the ribosome have very restricted substrate specificities, YwfE protein is unique not only in forming a dipeptide of L-amino acids but also in the wide spectrum of its products. Based on these characteristics, we propose to call the enzyme L-amino acid ligase.

YwfE protein apparently belongs to the ATP-dependent carboxylate-amine/thiol ligase superfamily for the following reasons. (i) It has a clear ATP-grasp sequence (Fig. 1) but shares no homology with a motif of aminoacyl-tRNA synthetase or the adenylation domain of NRPS. (ii) ADP, but not
AMP, and phosphate were formed in the reaction catalyzed by the enzyme. Since an enzyme that belongs to the ATP-dependent carboxylate-amine/thiol ligase superfamily transfers phosphate from ATP to its acyl substrate, it forms ADP and phosphate as reaction products (9, 14), whereas aminoacyl-tRNA

FIG. 4. HPLC analysis of the reaction catalyzed by YwfE protein. (a) Authentic samples of Ala (A), Gln (Q), Ala-Ala (AA), Ala-Gln (AQ), and Gln-Ala (QA). (b to d) Reaction with Ala and Gln at time zero (b) and 4 h (c) and reaction in which the sample at 4 h was coinjected with FMOC-derivatized Ala-Gln (d). (e) Authentic samples of Ser (S), Phe (F), Ser-Ser (SS), Ser-Phe (SF), and Phe-Ser (FS). (f to h) Reaction with Ser and Phe at time zero (f) and 4 h (g) and reaction in which the sample at 4 h was coinjected with FMOC-derivatized Ser-Phe (h). (i to k) Reaction with Gly (G) and Met (M) at time zero (i) and 4 h (j) and reaction in which the sample at 4 h was coinjected with FMOC-derivatized Gly-Met (GM) (k). (l to n) Reaction with Ala (A) and Leu (L) at time zero (l) and 4 h (m) and reaction in which the sample at 4 h was coinjected with FMOC-derivatized Ala-Leu (AL). Peaks indicated by single and double asterisks are derived from Tris buffer used for the enzyme reactions and FMOC reagent remaining in the reaction mixture, respectively.

FIG. 5. Time course of reactions with crude extracts (a) and with the purified enzyme fraction (b). Reaction conditions were as described in Materials and Methods except for the enzyme concentration (250 mg/liter). Open diamond; Ala; open triangle, Gln; solid circle, Ala-Gln; solid square, Ala-Ala.

FIG. 6. Effects of pH (a) and temperature (b) on YwfE protein activity. (a) Purified His-tagged YwfE protein (50 mg/liter) was incubated with 10 mM ATP, 5 mM each of Ala and Gln, and 10 mM MgSO₄ in 100 mM Tris-HCl (pH 7.0 to 10.0) or 100 mM borate-carbonate (pH 9.0 to 11.0) at 37°C for 1 h and the amount of Ala-Gln formed was measured. The amount relative to that formed at pH 9.5 with Tris-HCl was expressed as relative activity. (b) Purified His-tagged YwfE protein (50 mg/liter) was incubated with 10 mM ATP, 5 mM each of Ala and Gln, and 10 mM MgSO₄ in 100 mM Tris-HCl (pH 9.5) at various temperatures. The amount of Ala-Gln relative to that formed at 37°C was expressed.
synthetase or the adenylation domain of NRPS forms acyl-AMP as a reaction intermediate, and thus AMP and pyrophosphate are the reaction products (5). A previous report has suggested that the bacilysin-synthesizing reaction was carried out in an NRPS-like manner (29). The discrepancy with our results may arise from a difference in enzyme purity; the former study used a partially purified fraction whereas we used a fraction purified to apparent homogeneity (Fig. 2).

The substrate specificity of YwfE was unexpectedly relaxed (Fig. 8). For the N-terminal residue, Ala or Ser seems to be most preferred. Following these amino acids, Gly is preferred, suggesting that nonbulky and neutral amino acids tend to be chosen for the N terminus. For the C-terminal residue, Phe and Met are most reactive. Leu is next. Bulky and neutral amino acids seem to be preferred for the C terminus. Due to the unavailability of Cys-X and X-Cys peptides except for Gly-Cys, we could not confirm whether they or part of them are synthesized by the enzyme. However, the consumption of ATP and formation of ADP and phosphate were found with various combinations of amino acids containing Cys. As the reaction with Cys alone formed ADP, Cys is apparently taken as the N-terminal amino acid and, at the very least, Cys-Cys must be formed.

The substrate specificity of an enzyme is not simply defined as a matrix of N terminus-acceptable and C terminus-acceptable amino acids, but rather is determined by the combination of substrate amino acids (Fig. 8). For example, Ser is acceptable for the N terminus and Thr is acceptable for the C terminus, but Ser-Thr was not formed. Given that the enzyme belongs to the ATP-dependent carboxylate-amine/thiol ligase superfamily, the acylphosphate derivative of the N-terminal amino acid would be formed first. Thus, the amino acid accepted as the N-terminal residue may determine the possible range of C-terminal amino acids.

While the apparent $K_m$ was lower for Ala than Gln, 10 times more Ala-Gln than Ala-Ala was formed (Fig. 5b). One possibility is that the $V_{max}$ value for the synthesis of Ala-Gln is much higher. Another possibility is that the $K_m$ value of the N-terminal substrate binding site and that of the C-terminal substrate binding site are different, and the apparent $K_m$ for Ala was the sum of these values. This is supported by the fact that DDL has two subsites for binding D-Ala and the $K_m$s of the two sites are different (16). Clarifying how the enzyme recognizes its substrate and what factor affects the reaction rate is important for further application of this enzyme. Since DDL, a related enzyme, has been studied extensively, comparing the secondary structure of YwfE with that of DDL may reveal the relevant residues for substrate recognition.
The physiological significance of YwfE protein other than the participation in bacilysin biosynthesis is unclear. The following points seem to suggest that the enzyme has no additional role in B. subtilis. First, the synthesized dipeptides, except for bacilysin, would be decomposed rapidly by cellular peptidases (see below). This means the formation of a futile cycle between YwfE protein and peptidases that is meaningless for the cells except in some specific circumstances in which ATP is in excess. Second, the expression of ywfE is strictly controlled and coordinated with that of the bacilysin biosynthetic gene cluster (15, 29). Third, a ywfE-deficient mutant was reported to grow normally (11, 28). To elucidate the function of YwfE protein in more detail, examining the effect on cell growth of each dipeptide which can be synthesized by YwfE protein would be helpful.

The distribution of this new type of enzyme in nature is interesting, but no obvious homolog has been found in public databases. As we cannot assign any specific motif to L-amino acid ligases, screening with the same procedure used for ywfE would be useful for finding the next example of this type of enzyme. We adapted in silico screening plus an assay type of enzyme. We adapted in silico screening plus an assay for helpful discussions and Aya Kubota-Hada, Reiko Kouda, and Mayumi Fukano for technical assistance. We thank Yoshiko Matsu-moto for skillful MS analysis.

ACKNOWLEDGMENTS

We thank Akio Ozaki, Makoto Yagasaki, and Yoshiyuki Yonetani for helpful discussions and Aya Kubota-Hada, Reiko Kouda, and Mayumi Fukano for technical assistance. We thank Yoshiko Matsu-moto for skillful MS analysis.

REFERENCES


FIG. 8. Product spectrum of YwfE protein. Amino acids able to be accepted at the N terminus are listed vertically. The ones able to be accepted at the C terminus are listed horizontally. Arg, Lys, Glu, Asp, Pro, and trans-4-hydroxy-L-proline were inert. A filled circle indicates that the formation of the corresponding dipeptide was confirmed by HPLC analysis. An open circle indicates that synthesis of the corresponding dipeptide was suggested by the formation of ADP and phosphate and by the appearance of a new HPLC peak but could not be confirmed due to the unavailability of standard dipeptide. Reaction conditions were as described in Materials and Methods.


