Biodegradative Threonine Deaminase from *Proteus morganii*

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Biodegradative threonine deaminase was purified and crystallized from cells of *Proteus morganii* grown aerobically in the medium containing high concentrations of amino acids. The enzyme was homogeneous by the criteria of ultracentrifugation and SDS disc gel electrophoresis. The molecular weight of the enzyme was determined to be approximately 120,000 by the gel filtration method. The molecular weight of subunit of the enzyme was estimated to be 32,000 by SDS disc gel electrophoresis. The enzyme seems to have a tetrameric structure consisting of identical subunits. The enzyme has a marked yellow color with an absorption maximum at 415 nm and contains 4 moles of pyridoxal phosphate per mole of enzyme protein. The threonine deaminase was activated by AMP and ADP. It catalyzed the deamination of L-threonine, L-serine and β-chloro-L-alanine. The optimal pH was about 9.0 in the absence of AMP. In the presence of 1 mM AMP, the optimal pH became in a range between 8.0 and 9.5. The enzyme was not inhibited by L-isoleucine. Its Km values for L-threonine were found to be 1.5 mM and 24 mM, in the presence and absence of 1 mM AMP, respectively. The Ka value for AMP was found to be 0.55 mM at the threonine concentration of 5 mM. However, the value was varied depending on the concentration of L-threonine.

The existence of two distinctly different L-threonine deaminases (L-threonine hydro-lyase (deaminating) EC 4.2.1.16), biosynthetic and biodegradative ones, has been reported in *E. coli*. Both enzymes contain pyridoxal phosphate as a cofactor and catalyze α,β-elimination reaction of L-threonine to generate α-ketobutyrate and ammonia. Biosynthetic threonine deaminase participates in the biosynthesis of L-isoleucine from L-threonine and is characterized by end product inhibition by L-isoleucine. On the other hand, biodegradative one participates in the catabolism of L-threonine and is characterized by marked activation by AMP. In the bacterium, the biosynthetic one is formed in the cells constitutively, while the biodegradative threonine deaminase is formed inducibly under anaerobic conditions.

In our recent studies on threonine metabolism by bacteria, it was found that *Proteus morganii* produced both biosynthetic and biodegradative threonine deaminases simultaneously, under aerobic culture conditions. This paper will report the purification, crystallization and some properties of biodegradative threonine deaminase from *P. morganii*.

MATERIALS AND METHODS

Materials. L-Threonine was kindly provided by Ajinomoto Company, Ltd., Tokyo, and pyridoxal phosphate was by Dainippon Pharmaceutical Company, Ltd., Osaka. The other chemicals used were best available commercial products.

Microorganism. *Proteus morganii* (Faculty of Agriculture, Kyoto University, AKU 0084) was chosen for the enzyme source, since the bacterium had shown the highest threonine deaminase activity of all the 35 bacteria subjected to the screening.

Culturing procedures. The medium consisted of 0.5% polypepton, 0.5% meat extract, 0.65% yeast extract, 0.2% NaCl and 0.3% succinic acid in tap water. The pH of the medium was adjusted to 7.0. The strain was inoculated into a seed culture, 5 ml of
the medium in a test tube, which was incubated at 30°C for 24 hr with reciprocal shaking. The seed culture was then inoculated into a subculture, 500 ml of the same medium in a 2-liter shaking flask. After incubated at 30°C for 24 hr with reciprocal shaking, the subculture was inoculated into a 50-liter fermentor jar (MSJ-U501 Marubishi Co. Ltd., Tokyo, Japan) containing 40 liters of the same medium, which was then incubated at 30°C for 16 hr with aeration (40 liters/hr). Approximately 9 g of cells (wet weight) were obtained per liter of the medium.

**Enzyme assay.** The activity of threonine deaminase was determined spectrophotometrically by measuring the amount of α-ketobutyrate formed from L-threonine. The standard assay system contained 10 μmoles of L-threonine, 0.4 μmole of NH₄OH-NH₄Cl buffer, pH 9.5, and the enzyme in a total volume of 4.0 ml, unless otherwise specified. The reaction mixture was incubated at 30°C for 20 min and the reaction was then terminated by the addition of 1.0 ml of 30% trichloroacetic acid. The amount of the keto acid formed was determined as the 2, 4-dinitrophenylhydrazone at 450 nm by the method of Friedemann and Haugen,4) or as N-methylbenzothiazolonehydrazide at 320 nm by the modified method of Soda. (MBTH method).5) One unit of the enzyme activity was defined as the amount of enzyme producing 1 μmole of α-ketobutyrate per minute under the standard assay conditions. Specific activity was expressed as units per mg of protein.

**Protein determination.** Protein concentration was determined spectrophotometrically by measuring the absorbance at 280 nm. An E value of 12.7 for 10 mg per ml and for 1 cm light path, which was used throughout, was obtained by absorbance and dry weight determinations.

**Spectrophotometric determination.** Spectrophotometric determinations were carried out with a Beckman Spectra-20 spectrophotometer. Absorption spectra of the enzymes were measured with a Shimazu Multi-purpose Recording Spectrophotometer MPS-5000.

**Ultracentrifugal analysis.** Sedimentation velocity experiments were carried out with a Hitachi Analytical Ultracentrifuge operated at 51,200 rpm at 20°C. Diffusion coefficient was measured with the same apparatus operated at 9690 rpm at 20°C, with the boundary condition at the meniscus in a sector shaped centrifuge cell.6)

**Electrophoresis.** Disc gel electrophoresis was carried out with acrylamide gel in tris (hydroxymethyl) aminomethane-glycine buffer, pH 8.3, according to Ornstein7) and Davis.8) The disc gel electrophoresis in 0.1% SDS was carried out using the method of Weber and Osborn.9) After the run the gel was stained with 0.25% Coomassie brilliant blue, destained with diffusion, and stored in 7.5% acetic acid and 5% methanol.

**Determination of pyridoxal phosphate.** Pyridoxal 5'-monophosphate was determined with a Hitachi Fluorescence Spectrophotometer MPF-4 by the fluorometric method of Adams,10)

**RESULTS**

**Purification and crystallization of the enzyme**

All subsequent extraction and purification procedures were carried out at 0~5°C, unless otherwise specified.

**Step 1: Preparation of cell extract.** A total of 720 g (wet weight) of cells were suspended with 3 liters of a 0.05 M dilution buffer. This buffer consisted of various concentrations of K₂HPO₄-KH₂PO₄ buffer, pH 7.0, 1 mM 2-mercaptoethanol, 0.01 mM EDTA and 0.01 mM pyridoxal phosphate. The suspended cells were disrupted with a Dyno-mill KDL (Willy A Bachofen, Basel, Switzerland).

**Step 2: Streptomycin sulfate fractionation.** Seven hundred milliliters of 10% solution of streptomycin sulfate was added to the cell suspension treated with Dyno-mill, under constant stirring. After 30 min, the cell debris and precipitate formed were centrifuged off.

**Step 3: First ammonium sulfate fractionation.** The protein in the supernatant solution (4.2 liters) was precipitated by the addition of solid ammonium sulfate (70% saturation). The precipitate was dissolved in 1.1 liters of the 0.05 M dilution buffer described in step 1, and was dialyzed against 14 liters of the buffer for 36 hr. The buffer was changed 3 times at intervals of 9 hr. Insoluble materials formed during the dialysis were removed by centrifugation.

**Step 4: DEAE-Sephadex column chromatography.** The dialyzed solution (1.6 liters) was then applied to a column of DEAE-Sephadex A-50 (13.5 × 45 cm) equilibrated with the 0.05M dilution buffer. After washing the column with 20 liters of the 0.05 M dilution buffer and subsequent 20 liters of the 0.1 M dilution buffer, the elution of the enzyme was carried out with
Biodegradative Threonine Deaminase

Crude enzyme solution (protein content, 51 g, 13,430 units) was applied to a column (13.5 × 45 cm),
equilibrated with 0.05 M potassium phosphate buffer, pH 7.0, containing 1 mM 2-mercaptoethanol,
0.01 mM EDTA and 0.01 mM pyridoxal phosphate. Open circles (○–○) represent the absorbance of
protein at 280 nm and closed circles (●—●) represent the activity (units/ml).

0.1 M dilution buffer containing 0.2 M KCl. The enzyme activity was recovered at the elu-
tion volume from 11.7 to 17.1 liters (Fig. 1). The protein was precipitated and dialyzed by
the same way as described in step 3.

Step 5: Second ammonium sulfate fractionation. The dialyzed enzyme solution was
fractionated with ammonium sulfate at the intervals of 5% saturation. The threonine
deaminase activity was separated into two parts by this procedure. One was precipitated
between 30 and 45% saturation, and the other, between 45 and 60%. These two enzymes
were purified separately in subsequent steps. The purification and characterization of the
latter enzyme will be described in this report.

Step 6: Hydroxylapatite column chromatography. The enzyme solution (25 ml) dialy-
zed against the 0.01 M dilution buffer without EDTA was placed on a hydroxylapatite column
(3.0 × 6.0 cm) equilibrated with the same buffer. After washing the column with 600 ml of the
same buffer and then with the 0.05 M dilution buffer, the enzyme was eluted with 0.1 M potas-
sium phosphate buffer. The enzyme was recovered at the elution volume from 91 ml to
169 ml (Fig. 2).

Step 7: Sephadex G-200 gel filtration. The enzyme solution (3.0 ml) was subjected to
Sephadex G-200 gel filtration. The Sephadex was packed into a column (2.5 × 100 cm) and
equilibrated with the 0.05 M dilution buffer. The enzyme was eluted with the same buffer
and recovered at the elution volume from 210 ml to 252 ml (Fig. 3).

Step 8: Crystallization. To the concent-
trated enzyme solution (1.0 ml), solid am-
onium sulfate was gradually added until it
became slightly turbid. The test tube contain-
ing the solution was then placed in an ice bath
and was allowed to stand for several hours,
until silky sheen was observed apparently.
The test tube was then allowed to stand at
20°C overnight. Crystals were precipitated
at the bottom of the test tube, and thin and
rhombic crystals were observed under a micro-
scope (Fig. 4). The color of the crystals was
yellow.

The summary of the purification procedures
is shown in Table I.

Properties of the purified enzyme
Homogeneity. The specific activity of
enzyme achieved after the second crystalli-

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FIG. 2. Chromatography of the Partially Purified Biodegradative Threonine Deaminase on Hydroxylapatite.

Dialyzed enzyme solution (protein content, 882 mg, 6,840 units) was applied to a column (3.0×6.0 cm), equilibrated with 0.01 M potassium phosphate buffer, pH 7.0, containing 0.01 M 2-mercaptoethanol, 0.01 mM EDTA and 0.01 mM pyridoxal phosphate. Open circles (○) represent the absorbance of protein at 280 nm and closed circles (●) represent the activity (units/ml).

FIG. 3. Filtration of the Purified Biodegradative Threonine Deaminase on Sephadex G-200.

Purified enzyme solution (protein content, 66.3 mg, 1,800 units) was applied to a column (2.5×100 cm), equilibrated with 0.05 M potassium phosphate buffer, pH 7.0, containing 1 mM 2-mercaptoethanol, 0.01 mM EDTA and 0.01 mM pyridoxal phosphate. Open circles (○) represent the absorbance of protein at 280 nm and closed circles (●) represent the activity.

zation. The crystalline enzyme preparation migrated as a homogeneous entity when subjected to sedimentation experiments in an ultracentrifuge as shown in Fig. 5. A sedimentation coefficient (S20,w) of 4.63 S and a diffusion coefficient (D20,w) of 4.70×10−7 cm²/sec were determined. Though the enzyme had shown wide and obscure band on disc gel electrophoresis carried out according to Ornstein7) and Davis,8) it gave a clear single band on SDS disc gel electrophoresis (Fig. 6).

FIG. 4. Photomicrograph of Crystalline Biodegradative Threonine Deaminase from Proteus morganii.

Molecular weight of the enzyme and the subunit. The molecular weight of the enzyme was determined by gel filtration through a Sephadex G-200 column. The result had shown that the enzyme has a molecular weight of approximately 120,000 (Fig. 7). The mole-
The protein concentration was 0.33% in 0.05 M potassium phosphate buffer, pH 7.0, containing 1 mM 2-mercaptoethanol, 0.01 mM EDTA and 0.01 mM pyridoxal phosphate. The photographs were taken at intervals of 6 min after reaching 51,200 rpm at 20°C. These photographs were picked up at intervals of 12 min. Sedimentation is from left to right.

(a). The SDS-treated enzyme (13.2 µg) was subjected to the electrophoresis at a current of 8.0 mA for 6 hr in 0.01 M sodium phosphate buffer, pH 7.0, containing 0.1% SDS and 0.1% 2-mercaptoethanol. The direction of migration is from the cathode to the anode.

(b). Crystallized enzyme (25 µg) was applied to the stacking gel and subjected to the electrophoresis at a current of 5.0 mA for 30 min in Tris-glycine buffer, pH 8.3. The direction of migration is from the cathode to the anode.

TABLE I. PURIFICATION OF BIODEGRADATIVE THREONINE DEAMINASE FROM P. morganii

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Total protein (mg)</th>
<th>Total activity (units)</th>
<th>Specific activity (units/mg)</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell extract</td>
<td>150,000</td>
<td>13,100</td>
<td>0.087</td>
<td>100</td>
</tr>
<tr>
<td>Streptomycin sulfate</td>
<td>81,800</td>
<td>14,700</td>
<td>0.180</td>
<td>112</td>
</tr>
<tr>
<td>1st Ammonium sulfate</td>
<td>50,700</td>
<td>13,400</td>
<td>0.269</td>
<td>102</td>
</tr>
<tr>
<td>DEAE-Sephadex</td>
<td>3,900</td>
<td>10,800</td>
<td>2.77</td>
<td>82</td>
</tr>
<tr>
<td>2nd Ammonium sulfate</td>
<td>882</td>
<td>6,840</td>
<td>7.76</td>
<td>52</td>
</tr>
<tr>
<td>Hydroxyapatite</td>
<td>66.3</td>
<td>1,800</td>
<td>27.1</td>
<td>14</td>
</tr>
<tr>
<td>Sephadex G-200</td>
<td>24.9</td>
<td>1,600</td>
<td>64.3</td>
<td>12</td>
</tr>
<tr>
<td>Crystals</td>
<td>11.2</td>
<td>925</td>
<td>82.6</td>
<td>7</td>
</tr>
</tbody>
</table>

a) The units show only the enzyme activity precipitated 45~60% ammonium sulfate saturation. The enzyme activity precipitated 30~45% saturation had shown 4,250 units in total (32.4% in yield) and was not included in the following purification steps.

The molecular weight of subunit of the enzyme was estimated to be approximately 32,000 by SDS disc gel electrophoresis (Fig. 8). These data show that the native enzyme has a tetrameric structure consisted of 4 identical subunits.
Fig. 8. Estimation of Molecular Weight of Subunit of Biodegradative Threonine Deaminase by SDS Disc Gel Electrophoresis.

The conditions of electrophoresis are the same as described in Fig. 6. β-Galactosidase (E. coli), phosphorylase a (rabbit muscle), catalase (bovine liver), aldolase (rabbit muscle), trypsin (bovin pancreas) and α-chymotrypsin were used as marker proteins.

Absorption spectrum. The enzyme showed marked yellow color and its absorption spectrum at pH 7.0 had a peak at the wave length of 415 nm besides the absorption at 280 nm (Fig. 9). The enzyme has an $E_{1%1cm}^{1%}$ value of 12.7 at 280 nm obtained by absorbance and dry weight determinations. Absorption in the 410 to 440 nm region is characteristic of many pyridoxal phosphate enzymes and has been attributed to a hydrogen-bounded azomethine of pyridoxal phosphate.\(^{11}\)

Content of pyridoxal phosphate. The quantitative determination of pyridoxal phosphate in the enzyme was performed by the fluorometric method. After dialysis of the enzyme against 0.05 M dilution buffer, pH 7.0, the concentration of pyridoxal phosphate inside and outside the dialysis sac was determined by the method of Adams.\(^{10}\) A value of 3.4 moles per 120,000 g enzyme protein was observed, so the enzyme seems to contain about 4 moles of pyridoxal phosphate per mole of enzyme protein. This value of 4 moles of pyridoxal phosphate corresponds with the number of the subunit.

Substrate specificity. The relative rates of keto acid formation by the enzyme from various amino acids were measured and presented in Table II. L-Threonine, L-serine and β-chloro-L-alanine were degraded well, but not D-isomers of these amino acids. L-Serine showed approximately 17% reactivity in comparison with L-threonine and β-chloro-L-al-

![Figure 8: Estimation of Molecular Weight of Subunit of Biodegradative Threonine Deaminase by SDS Disc Gel Electrophoresis.](image_url)

![Figure 9: Absorption Spectrum of Biodegradative Threonine Deaminase.](image_url)

### Table II. Substrate Specificity of Biodegradative Threonine Deaminase from P. morganii

<table>
<thead>
<tr>
<th>Amino acid</th>
<th>Relative rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>L-Threonine</td>
<td>100</td>
</tr>
<tr>
<td>DL-Allothreonine</td>
<td>0</td>
</tr>
<tr>
<td>DL-α-Aminobutyrate</td>
<td>0</td>
</tr>
<tr>
<td>L-Methionine</td>
<td>0</td>
</tr>
<tr>
<td>L-Serine</td>
<td>17</td>
</tr>
<tr>
<td>L-Phenylalanine</td>
<td>0</td>
</tr>
<tr>
<td>L-Histidine</td>
<td>0</td>
</tr>
<tr>
<td>L-Alanine</td>
<td>0</td>
</tr>
<tr>
<td>S-Methyl-L-cysteine</td>
<td>0</td>
</tr>
<tr>
<td>L-Homoserine</td>
<td>0</td>
</tr>
<tr>
<td>β-Chloro-L-Alanine</td>
<td>83</td>
</tr>
</tbody>
</table>

The assay system contained 100 μmoles of amino acid, 0.4 μmoles of pyridoxal phosphate, 200 μmoles of bicine buffer, pH 9.0, and 2.0 μg of the enzyme in a total volume of 4.0 ml.
nine showed almost the same reactivity with L-threonine.

**Activation by adenosine nucleotides.** Effects of AMP, ADP, and ATP on the enzyme activity were investigated, since the activation of biodegradative threonine deaminase by these nucleotides was reported by many workers.\(^2,12,13\)

As shown in Table III, addition of AMP to the reaction mixture markedly stimulated the formation of α-ketobutyrate from L-threonine by the enzyme at pH 7.5 and 9.3. The activation of enzyme was also observed by the addition of ADP, but only at pH 7.5. It has no effect at pH 9.3. ATP did not show any activation effect at all. These activation by adenosine nucleotides strongly suggested that the enzyme from P. morganii was a so-called biodegradative threonine deaminase. Reactivity of L-serine and β-chloro-L-alanine was also stimulated in the presence of AMP around 10 to 15-fold, respectively.

The Michaelis constants for L-threonine and L-serine were determined in the presence and absence of AMP. As shown in Table IV, they were found to be 24.0 mM and 30.3 mM for L-threonine and L-serine, respectively, in the absence of AMP, and 1.5 and 8.9 mM for L-

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Km values (mM) without AMP</th>
<th>Km values (mM) with 1mM AMP</th>
</tr>
</thead>
<tbody>
<tr>
<td>L-Threonine</td>
<td>24</td>
<td>1.5</td>
</tr>
<tr>
<td>L-Serine</td>
<td>30</td>
<td>8.9</td>
</tr>
</tbody>
</table>

**Optimum pH.** Variation of enzyme activity with pH was determined with L-threonine and L-serine. The optimum pH for these substrates was around 9.0 in the absence of AMP (Fig. 10). In the presence of 1 mM AMP, the pH region at which the enzyme reacted optimally with L-threonine became much wider and it was found to be between 8.0 and 9.5 (Fig. 11).

**Activator constant.** Calculation of activator constants as performed at different substrate...
FIG. 11. Effect of pH on the Rate of Reaction of Biodegradative Threonine Deaminase in the Presence and Absence of AMP.

The conditions of determination of enzyme activities are the same as described in Fig. 10 except that the symbols (×-×) shows bicine buffer between pH 8.0 and 9.0. The enzyme, 1.8 μg was used in the presence of 1 mM AMP and 2.3 μg was used in its absence. L-Threonine was used as the substrate.

concentrations by plotting the reciprocals of the concentrations of AMP. Table V shows the \( K_a \) values for AMP at different substrate concentrations. The value decreases as the substrate concentration increases.

**TABLE V. DEPENDENCY OF \( K_a \) VALUES FOR AMP ON L-THREONINE CONCENTRATIONS**

The enzyme activities were determined under the standard assay system except that variable amounts of AMP were added to the reaction mixture.

<table>
<thead>
<tr>
<th>L-Threonine (mm)</th>
<th>( K_a ) values (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.50</td>
<td>3.13</td>
</tr>
<tr>
<td>1.25</td>
<td>1.12</td>
</tr>
<tr>
<td>2.50</td>
<td>0.95</td>
</tr>
<tr>
<td>5.00</td>
<td>0.55</td>
</tr>
</tbody>
</table>

**DISCUSSION**

*Proteus morganii* showed markedly high threonine deamination activity in its cells when it grew in a medium containing high concentration of yeast extract. During the purification of the enzyme, the enzyme activity was divided into two parts by ammonium sulfate fractionation. These activities can be ascribed to different enzymes because they can also be separated by gel filtration with Sephadex G–200 and by hydroxylapatite column chromatography. These two enzymes were purified separately and distinguished from each other. The purification and some properties of the biodegradative enzyme were described in this paper. The other enzyme, which is inhibited by L-isoleucine and should be a biosynthetic threonine deaminase, will be described elsewhere.

In *E. coli*, formation of the biodegradative threonine deaminase was observed only when the bacterium grew anaerobically in the medium containing high concentration of amino acids and no glucose. Formation of the biodegradative threonine deaminase by *P. morganii* was, however, observed under aerobic conditions. The biodegradative enzyme from *E. coli* was unstable and required AMP not only to show the maximum activity but also for stabilization and crystallization, while the enzyme from *P. morganii* was considerably stable even in the absence of AMP and it could be purified and crystallized in its absence, although it showed the maximum activity in the presence of AMP.

The molecular weight of the enzyme from *P. morganii* was approximately 120,000 and that of the subunit was about 32,000, suggesting the enzyme to consist of 4 identical subunits. Taking into consideration this fact and according to the result of fluorometric analysis, the enzyme seems to contain 4 moles of pyridoxal phosphate per mole of enzyme protein. These properties were very similar to those of biodegradative threonine deaminase from *E. coli*.

Wood and Gunsalus reported that biodegradative threonine deaminase from *E. coli* was activated by AMP and Hayashi et al. reported that the enzyme from *Clostridium tetanomorphum* was activated by ADP, while we found out that the enzyme from *P. morganii* was activated by both AMP and ADP.

The \( K_a \) value for AMP of the enzyme was found to be dependent on the concentration of L-threonine. It was 0.55 mM at threonine con-
centration of 5.0 mM, whereas the $K_a$ value for AMP of the enzyme from *E. coli* was reported to be 1.7 mM\(^{13}\) and the $K_a$ value for ADP of the enzyme from *C. tetanomorphum* was 0.006 mM\(^{15}\) at the same threonine concentration of 5.0 mM.

REFERENCES