Carbon–Phosphorus Hydrolase: Some Properties of the Enzyme in Cell Extracts of Enterobacter aerogenes

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Cells of Enterobacter aerogenes IFO 12010 could grow on a medium containing alkylphosphonic acids as sole sources of phosphorus. The formation of the enzyme responsible for the liberation of inorganic phosphate (Pi) from alkylphosphonic acids was induced when Pi-grown cells were transferred to a medium containing no added Pi. The extracts prepared from cells after incubation in the absence of Pi showed high carbon–phosphorus (C–P) bond cleavage activity and catalyzed the liberation of Pi from alkylphosphonic acids most efficiently in the alkaline pH region. We named the enzyme C–P hydrolase, tentatively.

Despite extensive studies on the distribution of alkylphosphonic acids in living systems and their utilization by microorganisms as phosphorus sources,1~7) the activity of a carbon–phosphorus (C–P) bond cleavage enzyme had not been detected hitherto in a cell-free system, and further progress in the biochemistry of alkylphosphonic acids is greatly dependent on the detection of cell-free activity.

In order to obtain microbial strains with high C–P bond cleavage activity, we screened bacterial strains that are capable of accumulating inorganic phosphate (Pi) in culture medium containing phosphonoacetic acid.9) Among forty bacterial strains tested, two (Bacillus roseus AKU 0208 and Enterobacter aerogenes IFO 12010) were found to accumulate substantial amounts of Pi in the culture medium, possibly due to the cleavage of the C–P bond in phosphonoacetic acid, and we detected, for the first time, C–P bond cleavage activity in cell-free extracts of Enterobacter aerogenes IFO 12010.9)

The present study was undertaken to obtain further information on the C–P bond cleavage enzyme in Enterobacter aerogenes IFO 12010.

Materials and Methods

Cultivation and preparation of cell extracts. Cells of Enterobacter aerogenes IFO 12010 were grown in 100 ml of medium D [0.5% glucose, 0.1% (NH₄)₂SO₄, 0.01% MgSO₄·7H₂O, 0.005% yeast extract (Pi-free) and 30 mM N-(2-hydroxyethyl)piperazine-N'-2-ethanesulfonic acid (HEPES), pH 7.2] supplemented with various phosphorus sources. Yeast extract (Pi-free) was prepared by magnesia treatment10) and contained no detectable Pi. Alkylphosphonic acids were filter-sterilized before use. Glassware was rinsed with nitric acid to minimize Pi contamination. Cultivation was carried out in 5 dl Sakaguchi flasks at 30°C with reciprocation (120 rpm, 3 cm stroke). When the turbidity of a culture reached 0.7 (O.D. at 610 nm), the cells were collected, washed once in 20 ml of 10 mM Tris-HCl buffer (pH 7.5) containing 0.5 mM MgCl₂ [Tris/Mg], and then disrupted ultrasonically at 90 kHz and 0°C for 5 min. The homogenate was centrifuged at 25,000 × g for 30 min and the resulting supernatant was used as an enzyme source after dialysis against Tris/Mg at 4°C overnight.

Enzyme assays. C–P bond cleavage enzyme activity was assayed in a mixture (0.5 ml) comprising 50 mM alkylphosphonic acid, 20 mM MgCl₂, 50 mM Tris–HCl buffer (pH 7.5) and cell extract (10 mg/ml protein) at 37°C for 1 hr. The reaction was started by adding the cell extract and terminated by the addition of 0.5 ml of 25% trichloroacetic acid. The mixture was centrifuged and then Pi in the clear solution was determined by the method of Fiske and Subbarow.11) Alkaline phosphatase was assayed in a mixture (1.0 ml) consisting of 10 mM p-nitrophenylphosphate,
20 mM MgCl₂ and 50 mM Tris-HCl buffer (pH 8.0). The reaction was initiated by the addition of the cell extract and then the increase in absorbance at 412 nm was recorded at 25°C. One unit of enzyme activity was defined as the amount liberating 1.0 μmol of Pi per hour. Protein was determined according to the method of Lowry et al.¹²

Cell transfer experiment. Cells were grown in Medium D (600 ml in a 2 l Sakaguchi flask) supplemented with 5.0 mM Pi at 30°C with reciprocation, as above. When the turbidity of a culture reached 0.7 (O.D. at 610 nm), the cells were collected, washed once in 20 ml of 0.85% saline solution and then transferred to a 5 dl Sakaguchi flask containing 100 ml of Medium D or Medium D supplemented with 5.0 mM Pi or 1.0 mM phosphonoacetic acid. The initial cell concentration was adjusted to 7.0 x 10⁸/ml. After incubation for several hours at 30°C with reciprocation, the cells were collected and cell extracts were prepared as above.

Column chromatographies. Cells were grown on 15 l of Medium D [1500 ml in a 2 l Sakaguchi flask] supplemented with 0.3% phenylphosphonic acid at 30°C with reciprocation for 24 h and then cell extracts were prepared as before. Glycerol was added to each extract to a final concentration of 30% and then the extract was dialyzed against Tris/Mg containing 30% glycerol [Tris/Mg/Gly] at 4°C overnight. The dialysate was applied to a DEAE-cellulose column (4 cm x 32 cm) equilibrated with Tris/Mg/Gly and proteins were eluted with a linear gradient of FCCl, from 0 to 0.6 M (total volume, 1300 ml) in the same buffer system. 7 ml fractions were collected per 7 min. The Z-P bond cleavage enzyme fractions (Fraction Nos. 50~91) were pooled and then concentrated to 30 ml by ultrafiltration with an Amicon PM10 membrane. The concentrate was then loaded onto a Sephadex G-150 column (2.5 cm x 90 cm) equilibrated with Tris/Mg/Gly. Proteins were eluted with the same buffer and 4 ml fractions were collected per 25 min.

Chemicals. Methylphosphonic acid, phosphonoacetic acid and phenylphosphonic acid were purchased from Aldrich Chem. Co., Milw. Phosphonomycin was from Sigma Chemical, St. Louis, MO. Glyphosate was kindly supplied by Monsanto Japan Co., Ltd., Ryugasaki, Japan.

Results and Discussion

Utilization of alkylphosphonic acids as phosphorus sources

The utilization of alkylphosphonic acids by E. aerogenes IFO 12010 cells as sole sources of phosphorus was examined by cultivating cells in Medium D supplemented with various alkylphosphonic acids (Fig. 1). In the absence of phosphorus source, slight growth was observed, that could be explained by phosphorus contamination of Medium D. Substantial growth was observed when various alkylphosphonic acids were added to the medium, at 1.0 mM. The cells showed faster growth on Pi than on any of the alkylphosphonic acids tested. During growth on phenylphosphonic acid, the cells formed clumps and so accurate turbidimetric determination of growth was impossible, the reason for the aggregation of cells being not clear yet (Fig. 2). The poor growth of cells on phenylphosphonic acid may be partly attributable to this aggregation. The results of the growth experiment indicated that cells of E. aerogenes IFO 12010 could utilize alkylphosphonic acids as sole sources of phosphorus and the cells contained an enzyme that catalyzes the cleavage of the C-P bonds in alkylphosphonic acids.

Inducibility of the C-P bond cleavage enzyme

When cells of Enterobacter aerogenes IFO 12010 were grown on Medium D supplemented with 5.0 mM Pi, the extracts showed no detectable C-P bond cleavage enzyme activity, i.e., no liberation of Pi from phosphonoacetic acid (data not shown). This suggested that the enzyme is inducible and that its formation is dependent on the cellular Pi concentration. In

Fig. 1. Utilization of Alkylphosphonic Acids as Sole Sources of Phosphorus.

Cells were cultured in Medium D supplemented with various phosphorus sources, at 1.0 mM. -▲-, none; -●-, orthophosphate; -○-, methylphosphonic acid; -■-, phosphonoacetic acid; -□-, phenylphosphonic acid.
order to confirm the inducibility of the enzyme, a cell transfer experiment was carried out as described under Materials and Methods (Fig. 3). The C–P bond cleavage enzyme activity remarkably increased when Pi-grown cells were transferred to Medium D (Pi-free) or Medium D supplemented with 1.0 mM phosphonoacetic acid. The induction of the enzyme in medium containing phosphonoacetic acid was almost one-half that in medium containing no added Pi. This is presumably due to the repression by Pi formed from phosphonoacetic acid by the enzyme induced. On the other hand, cells transferred to Medium D containing 5.0 mM Pi showed no increase in the enzyme activity. Thus, the enzyme catalyzing the C–P bond cleavage is inducible, which supports the observation of Wackett et al. that the gene responsible for the C–P bond cleavage
enzyme is located in the Phosphate Starvation Inducible (PSI) regulon in *Escherichia coli*.\textsuperscript{13)}

**C–P bond cleavage enzyme activity in extracts of *E. aerogenes***

Previously, we detected C–P bond cleavage enzyme activity in extracts of *E. aerogenes* IFO 12010 cells grown on a medium containing phosphonoacetic acid as a sole source of phosphorus.\textsuperscript{9)} In order to compare the catalytic properties of the enzyme induced in the absence of Pi with those of that formed in the presence of phosphonoacetic acid, Pi-grown cells were transferred to Medium D (Pi-free), to induce the C–P bond cleavage enzyme, and then incubated for 3 hr according to the procedures described for the cell transfer experiment under Materials and Methods. Extracts containing the induced enzyme were incubated with various alkylphosphonic acids, including a herbicide (glyphosate) and an antibiotic (phosphonomycin) (Fig. 4). The liberation of Pi from alkylphosphonic acids increased with both increasing protein concentration (Fig. 4A) and reaction time (Fig. 4B). The enzyme was highly active on phosphonoacetic acid and catalyzed the liberation of Pi most efficiently in the alkaline pH region (Fig. 5). Thus, the catalytic properties of the C–P bond cleavage enzyme induced in the absence of Pi were almost the same as those of that formed in the presence of phosphonoacetic acid.\textsuperscript{9)} The gene for the C–P bond cleavage enzyme in *E. aerogenes* IFO 12010 may be applicable to the breeding of herbicide-resistant crops, since the enzyme can abolish the herbicidal effect of glyphosate by cleaving the C–P bond in the compound (Fig. 4).

**An attempt to purify the C–P bond cleavage enzyme in *E. aerogenes***

Extracts of *E. aerogenes* IFO 12010 cells grown on phenylphosphonic acid as a sole source of phosphorus were fractionated on DEAE-cellulose (Fig. 6A) and Sephadex G-150 (Fig. 6B) columns as described under Materials and Methods. The C–P bond clea-

![Fig. 5. Effect of pH on the C–P Bond Cleavage Enzyme Activity.](image)

The activity was determined with phosphonoacetic acid as a substrate. Other conditions were the same as those under Materials and Methods, except that various buffers were used, at 0.1 m. The activity at pH 8.0 was taken as 100%: —, acetate buffer; — O — HEPES/NaOH buffer; — □ —, Tris/HCl buffer; — ■ —, glycine/KOH buffer.

![Fig. 6. Fractionation of the C–P Bond Cleavage Enzyme by Column Chromatographies.](image)

The operation conditions for the DEAE-cellulose [A] and Sephadex G-150 [B] columns are given under Materials and Methods. The activity of the C–P bond cleavage enzyme was determined with phosphonoacetic acid as a substrate. — O —, absorbance at 280 nm; —■ —, C–P bond cleavage enzyme activity; — ▲ —, alkaline phosphatase activity; — — , KCl concentration.
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The enzyme was co-eluted with alkaline phosphatase, at approximately 0.1 m KCl, from the DEAE-cellulose column. However, the enzyme was successfully separated from alkaline phosphatase on the Sephadex G-150 column, although there was a significant loss of its activity.

Thus, we could detect the activity of a C-P bond cleavage enzyme in cell free extracts of *E. aerogenes* IFO 12010. As to the C-P bond cleavage in phosphonoacetic acid, we first thought that phosphonoacetic acid is reduced to phosphonoacetaldehyde and then the C-P bond is cleaved by an enzyme, “phosphonatase”, as was assumed for the C-P bond cleavage in 2-aminoethylphosphonic acid. However, the enzyme we found in *E. aerogenes* IFO 12010 requires no cofactors for its catalytic function. Furthermore, “phosphonatase” was shown to be inactive on alkyl- and phenylphosphonic acids. Therefore, the C-P bond cleavage enzyme we found in *E. aerogenes* IFO 12010 is different from “phosphonatase,” although the occurrence of the latter enzyme has not been confirmed in *E. aerogenes* IFO 12010.

Wackett et al. suggested that certain microorganisms contain an enzyme that catalyzes C-P bond cleavage in various phosphonic and phosphinic acids, and they named the enzyme “C-P lyase.” The enzyme we found in *E. aerogenes* IFO 12010, however, seems to catalyze the hydrolysis of the C-P bonds in phosphonic (and possibly phosphinic) acids, and thus should be classified as a hydrolase, not a lyase. We propose to call the enzyme a “Carbon–phosphorus (C–P) hydrolase” until the reaction mechanism of the enzyme has been fully elucidated. Anyway, this is the first report of a microbial “C-P hydrolase” that hydrolyzed C–P bonds in alkyl- and phenylphosphonic acids.

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References