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Regulation and function of transaldolase isoenzymes involved in sugar and one-carbon metabolism in the ribulose monophosphate cycle methylotroph Arthrobacter P1

P. R. Levering and L. Dijkhuizen
Department of Microbiology, University of Groningen, Kerklaan 30, NL-9751 NN Haren, The Netherlands

Abstract. In the facultative methylotroph Arthrobacter P1 the enzyme transaldolase plays an important role in both the pentose phosphate pathway and in the ribulose monophosphate cycle of formaldehyde fixation.

Among gluconate-negative mutants of Arthrobacter P1 strains occurred which also were unable to grow on xylose because they had lost the ability to synthesize transaldolase. Furthermore, this loss of transaldolase activity resulted in decreased growth rates on a number of other “hetero- trophic” substrates. Contrary to expectation, these mutants still grew on methyamine and were even able to use gluconate as a carbon source at normal rates provided methyamine was supplied as a nitrogen source. Under these conditions high levels of transaldolase were observed.

Partial purification of the transaldolases synthesized by gluconate-grown cells of wild type Arthrobacter P1 and methyamine-grown cells of one of these mutants, strain Art 98, revealed the presence of transaldolase isoenzymes. These enzymes displayed similar kinetics but were very different in heat sensitivity. Functionally these isoenzymes are apparently very similar but their synthesis is regulated differently. One of the enzymes is synthesized constitutively whereas the other is specifically induced during growth on C1 compounds. Strain Art 98 has lost the ability to synthesize the constitutive transaldolase. It is postulated that the C1-induced transaldolase serves to ensure a sufficiently high rate of regeneration of ribulose-5-phosphate during growth on C1 compounds.

Key words: Arthrobacter P1 - Transaldolase - Isoenzymes - Methylotrophy - RuMP cycle of formaldehyde fixation - Regulation - Gluconate - Methylamine - Xylose - Pentose phosphate pathway

The presence of the enzyme transaldolase (δ-sedoheptulose-7-phosphate: δ-glyceraldehyde-3-phosphate dihydroxyacetone transferase, EC 2.2.1.2) has been reported in a large variety of microorganisms, prokaryotes as well as eukaryotes. Its function is the interconversion, together with transketolase, of C3 to C7 sugar phosphates in the non-oxidative branch of the pentose phosphate pathway (Tsolas and Horecker 1972). In these reactions the precursors for the biosynthesis of nucleic acids, ribose-5-phosphate, and aromatic amino acids, erythrose-4-phosphate, are formed (Fig. 1).

In addition to this anabolic function, transaldolase also plays a role in the metabolism of those carbon- and energy sources which are converted via the pentose phosphate pathway to fructose-6-phosphate and glyceraldehyde-3-phosphate. Examples of this are the metabolism of gluconate in a number of Arthrobacter species (Zagallo and Wang 1962; Levering et al. 1982) and the metabolism of glucose and xylose in the yeast Candida utilis (Tsolas and Horecker 1972).

A somewhat different function is fulfilled by transaldolase in autotrophic and methylotrophic microorganisms that employ either the Calvin cycle, the ribulose monophosphate (RuMP) cycle or the xylulose monophosphate cycle as the primary carbon assimilation pathway. In these organisms the enzyme may be involved in the regeneration of the C1-acceptor molecule during autotrophic or methylotrophic growth (Quayle 1980; Anthony 1982). This has been demonstrated for instance in Arthrobacter P1, a facultative methylotroph able to grow rapidly on methyamine (Levering et al. 1981a). During growth of Arthrobacter P1 on C1 compounds, formaldehyde is assimilated via the RuMP cycle which in this organism characteristically involves transaldolase and transketolase in the rearrangement reactions (Levering et al. 1982; Fig. 1).

During our studies on the regulation of C1 metabolism in Arthrobacter P1 we tried to obtain mutant evidence for the suggested involvement of the so-called dissimilatory RuMP cycle of formaldehyde oxidation (Anthony 1982). This cycle involves glucose-6-phosphate and 6-phosphogluconate dehydrogenases as catalysts in energy-generating reactions during methylotrophic growth (Levering et al. 1981a; Fig. 1). As shown in Fig. 1 the metabolism of gluconate in Arthrobacter P1 is by way of the pentose phosphate pathway and, subsequently, the Embden-Meyerhof route (Levering et al. 1982). We therefore decided to try and isolate gluconate-negative mutants of Arthrobacter P1 deficient in 6-phosphogluconate dehydrogenase synthesis. Some of the gluconate-minus mutants that were obtained, however, were also unable to grow on xylose indicating that the metabolic blocks in these strains were in one of the enzymes of the non-oxidative branch of the pentose phosphate pathway. Contrary to our expectation, these mutants were still able to grow on methyamine. Subsequent studies revealed the presence of transaldolase isoenzymes in Arthrobacter P1.
enzymes listed in Table 2. Abbreviations: DHAP, dihydroxyacetone phosphate; GAP, glyceraldehyde-3-phosphate; E4P, erythrose-4-phosphate; Ru5P, ribulose-5-phosphate; R5P, ribose-5-phosphate; X5P, xylulose-5-phosphate; H6P, hexulose-6-phosphate; F6P, fructose-6-phosphate; FBP, fructose-1,6-bisphosphate; 6PG, 6-phosphogluconate; G6P, glucose-6-phosphate; S7P, sedoheptulose-7-phosphate.

In this paper we report the result of studies on the regulation and function of transaldolase isoenzymes in wild type and mutant strains of Arthrobacter P1.

Materials and methods

Organisms

Arthrobacter P1 NCIB 11625 has been described previously (Levering et al. 1981a). Mutants of Arthrobacter P1 were isolated following UV light (254 nm) irradiation as described by Dijkhuizen et al. (1981). Among the survivors of this UV treatment strains were selected that had lost the ability to grow on gluconate. Both wild type and mutant strains were maintained on plates containing 1% (w/v) yeast extract and 1.2% (w/v) agar.

Medium and cultivation

Arthrobacter P1 wild type and the gluconate-negative strain Art 98 were grown at 30 °C in 1-l conical flasks containing 250 ml of the mineral salts medium described by Levering et al. (1981a). Heat-sterilized solutions of carbon sources were added to the following final concentrations: glucose, 10 mM; potassium gluconate, 10 mM; sodium succinate, 15 mM; glycerol, 20 mM; and methylammonium chloride, 40 mM. Ammonia-free mineral salts medium was prepared as described by Levering and Dijkhuizen (1985). This medium was supplemented with methylammonium chloride (25 mM) as the nitrogen source and potassium gluconate as the carbon source. Incubation was on a rotary shaker at 200 rev./min.

Growth of the organism was monitored by measuring the absorbance of the cultures at 433 nm in a Vitatron 280 colorimeter (Vitatron, Dieren, The Netherlands). Cells for enzyme analysis were harvested, before they had reached the stationary growth phase by centrifugation at 6,000 × g for 10 min at 4 °C, washed once with 50 mM potassium phosphate buffer pH 7.0 containing 5 mM MgSO4 and resuspended in this buffer to a concentration of 5–10 mg dry weight/ml. These suspensions were used immediately for enzyme assays or stored at −20 °C until required.

Substrate transition experiments

Cells for the substrate transition experiments were pregrown on methylamine as described above, harvested aseptically in the late-exponential growth phase by centrifugation at 6,000 × g for 10 min at 20 °C, washed once with mineral salts medium and resuspended in a small volume of this medium. These suspensions were subsequently used as inocula for 1-l fresh mineral salts medium, containing potassium gluconate (10 mM), in 3-l conical flasks. Incubation was on a rotary shaker at 250 rev./min and 30 °C. Growth was monitored as described above and at appropriate time intervals samples were withdrawn from the cultures for the assay of enzyme activities. In the experiment with mutant strain Art 98 the culture was analysed for revertants by plating periodically on gluconate (10 mM) mineral salts-agar (1.2%, w/v).

Assays of enzyme activity

Cell-free extracts were prepared as described by Levering et al. (1981a). Spectrophotometric assays were performed with a Hitachi 100-60 spectrophotometer at 30 °C. The following enzymes were assayed according to published procedures: 6-phosphogluconate dehydrogenase (NADP-dependent) (EC 1.1.1.44) according to van Dijken and Quayle (1977), except that 50 mM Tris-maleate pH 8.0 was used as a buffer; hexulose phosphate synthase, transaldolase (EC 2.2.1.2), transketolase (EC 2.2.1.1), ribulose-5-phosphate 3-epimerase (EC 5.1.3.1) and ribose-5-phosphate isomerase (EC 5.3.1.6) according to Levering et al. (1981a, 1982).

Purification of transaldolases

All steps were carried out at 0–4 °C unless stated otherwise. Both purification steps (ion exchange chromatography and gel permeation) were performed with a high-performance liquid chromatography (HPLC) system, which consisted of a Model 6000A solvent delivery system equipped with a U6K injector and a Model 441 absorbance detector (Waters Associates, The Netherlands). The purification procedure was as follows:

1. Preparation of crude extracts. Arthrobacter P1 and mutant strain Art 98 were grown on gluconate or methylamine, harvested by centrifugation, washed with 50 mM potassium phosphate buffer pH 7.0 and subsequently resuspended in a small volume of this buffer. These suspensions (3 ml; 50–75 mg dry wt/ml) were sonified with 1 g of glass beads at 20 kHz for 15 × 15 s with a MSE 100 W ultrasonic disintegrator (MSE Ltd., Crawley, Sussex, UK), with 45 s intermitting periods. Unbroken cells and debris were removed by centrifugation at 40,000 × g for 10 min and the supernatant was again centrifuged at 40,000 × g for 30 min. The pellet was discarded and the supernatant thus obtained contained approx. 25 mg protein/ml and was subjected to ion-exchange chromatography.

2. DEAE-polyol chromatography. Protein samples (1.0 ml) were injected in a column (9.5 × 500 mm) containing
diethylaminoethyl (DEAE)-Si 100 Polyol (Serva, Heidelberg, FRG) and eluted with a non-linear sodium chloride gradient (0 – 1 M) in 25 mM potassium phosphate buffer pH 7.0 at room temperature. The flow rate was set at 3.0 ml/min and 1.5 ml fractions were collected. Enzyme activities were measured as indicated above and peak fractions were pooled for further purification. The gradient was run with the aid of an Acorn Atom microcomputer (Acorn Computers Ltd., Cambridge, UK) interfaced with a three-way solenoid valve (type LFYA, The Lee Company, Westbrook, CT, USA) as described by van der Zee and Welling (1984).

3. Protein concentration. Before being subjected to gel filtration, the pooled fractions from the DEAE column were concentrated (25 – 30 x) with a Minicon B-15 system (Amicon Co., Danvers, MA, USA).

4. Gel permeation chromatography. Samples (0.1 ml) containing 0.5 – 1.0 mg protein were applied to a Protein Pak 300-SW gel permeation column (7.5 x 300 mm; Waters Associates, The Netherlands). Several runs were made with 0.2 M potassium phosphate buffer pH 7.0, with a flow rate of 0.5 ml/min at room temperature. Fractions (0.25 ml) were collected and analyzed for transaldolase activity. The fractions containing the highest activity were pooled and frozen at – 20°C until required for further analysis.

Molecular weight determinations
Molecular weights of the purified transaldolase preparations were estimated with the gel filtration technique as described above, except that 0.05 ml samples containing 5 – 10 µg protein were injected. Chymotrypsinogen A, mw 25,000, albumin, mw 45,000 and albumin, mw 68,000 (5 µg of each; products of Boehringer, Mannheim, FRG) were used as marker proteins.

SDS-PAA gel electrophoresis
Sodium dodecyl sulphate polyacrylamide gel electrophoresis was performed according to Laemmli and Favre (1973) with the calibration proteins Combithek 104558 from Boehringer (Mannheim, FRG) as references. After electrophoresis the gels were stained with the silver staining technique as described by Wray et al. (1981).

Phosphate precipitation
Since high concentrations of phosphate inhibited transaldolase activity this compound was precipitated by adding a sufficient amount of calcium acetate to the enzyme preparations. The precipitate was removed by centrifugation and the supernatant thus obtained was subsequently used for kinetic studies. Calcium acetate up to 80 mM was shown not to inhibit the activity of transaldolase when added to assay mixtures for this enzyme.

Protein determination
Protein concentrations were determined according to Lowry et al. (1951), using bovine serum albumin as a standard.

Biochemicals
Erythrose-4-phosphate was obtained from Boehringer (Mannheim, FRG) as the diethylacetel monobarium salt. Transaldolase, purified from baker’s yeast, was obtained from Sigma Chemical Co. (St. Louis, MO, USA). All other biochemicals were obtained from commercial sources and were of the highest available grade.

Results
Isolation and characterization of mutant strain Art 98
Following UV light irradiation five gluconate-negative strains of Arthrobacter P1 were isolated. A subsequent analysis of enzyme activities in cell-free extracts prepared from these mutants after growth on glucose indicated that 6-phosphogluconate dehydrogenase was present in normal activities. Thus, these mutants were either unable to take up gluconate from the medium, and/or phosphorylate it to 6-phosphogluconate, or were blocked in the non-oxidative branch of the pentose phosphate pathway (Fig. 1). Two of the gluconate-negative strains were also unable to utilize xylose as a carbon- and energy source and one of these mutants, designated strain Art 98, was subsequently studied in more detail.

Compared to wild type Arthrobacter P1, strain Art 98 exhibited significantly increased doubling times with glucose, glycerol or succinate (Table 1). From an analysis of the activities of the relevant enzymes (see above) in cell-free
Table 1

Doubling times of wild type *Arthrobacter* P1 and mutant strain Art 98 during growth on various substrates, and specific activities of transaldolase (nmol · min⁻¹ · mg⁻¹ of protein) in cell-free extracts

<table>
<thead>
<tr>
<th>Growth substrate</th>
<th>Doubling time (h)</th>
<th>Transaldolase activity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>wild type</td>
<td>Art 98</td>
</tr>
<tr>
<td>Gluconate</td>
<td>1.3</td>
<td>a</td>
</tr>
<tr>
<td>Xylose</td>
<td>1.4</td>
<td>a</td>
</tr>
<tr>
<td>Glucose</td>
<td>1.2</td>
<td>2.2</td>
</tr>
<tr>
<td>Glycerol</td>
<td>1.8</td>
<td>5.6</td>
</tr>
<tr>
<td>Succinate</td>
<td>1.3</td>
<td>2.0</td>
</tr>
<tr>
<td>Methylamine</td>
<td>2.5</td>
<td>2.5</td>
</tr>
<tr>
<td>Choline</td>
<td>2.2</td>
<td>2.6</td>
</tr>
<tr>
<td>Gluconate + methylamine minus (NH₄)₂SO₄</td>
<td>1.3</td>
<td>1.4</td>
</tr>
</tbody>
</table>

a No growth  
b Enzyme activity not detectable  
c Data of Levering et al. (1982)  
d Enzyme activity present  
nd Not determinable

Table 1

Doubling times of wild type *Arthrobacter* P1 and mutant strain Art 98 during growth on various substrates, and specific activities of transaldolase (nmol · min⁻¹ · mg⁻¹ of protein) in cell-free extracts it became clear that the metabolic block in strain Art 98 is in transaldolase synthesis. After growth of strain Art 98 on these substrates no transaldolase activities were detectable, whereas wild type cells grown under the same conditions contained high levels of this enzyme (Table 1). Strain Art 98 did grow on methylamine with a doubling time of 2.5 h, as normally observed for wild type *Arthrobacter* P1. In cell-free extracts prepared from these cells high activities of transaldolase were detected and in case of strain Art 98 it could be shown, by plating out on gluconate agar, that this was not due to the appearance of revertants. Strain Art 98 also synthesized transaldolase during growth on choline, albeit in rather low levels. Further studies demonstrated that Art 98, although unable to grow on gluconate with ammonium sulphate as a nitrogen source, can use gluconate as a carbon- and energy source when incubated in the presence of methylamine as the sole nitrogen source. In the latter situation the growth was much faster than on methylamine alone and almost equalled the growth rates of wild type *Arthrobacter* P1 on gluconate/ammonia and gluconate/methylamine (Table 1). High activities of transaldolase were detected following growth on gluconate plus methylamine and again it could be shown that this was not caused by the presence of revertants.

The regulation of the synthesis of transaldolase in wild type *Arthrobacter* P1 and strain Art 98 was subsequently studied in substrate-transition experiments in which methylamine-grown cells were transferred into media containing gluconate as the sole carbon- and energy source and ammonia as the nitrogen source (Fig. 2). After incubation of *Arthrobacter* P1 in this medium growth started after 1 h and a doubling time of 1.3 h was reached in a few hours (Fig. 2A). An analysis of enzyme activities in cell-free extracts showed that hexulose phosphate synthase, an enzyme induced by formaldehyde during growth on methylamine (Levering et al. 1984, 1986a, b), decreased rapidly in activity and was no longer detectable after 4.5 h. The same response was observed for transaldolase, but the strong decrease in the activity of this enzyme levelled off, at 4.5 h, to reach a value of approximately 500 nmol · min⁻¹ · mg⁻¹ protein. The same activity is normally found with methylamine-grown cells of the organism (Fig. 2). Finally, the activity of 6-phosphogluconate dehydrogenase reached a somewhat higher level.

Two possible explanations were considered for the phenotype and characteristic properties of mutant Art 98. Assuming that only one transaldolase enzyme is present in *Arthrobacter* P1, the data indicate that in Art 98 this enzyme can only be induced by methylamine and choline, but not anymore by other substrates such as glucose, gluconate, glycerol, xylose and succinate. Alternatively, the data may be interpreted to indicate that *Arthrobacter* P1 is able to synthesize two transaldolase isoenzymes and that only one of these enzymes is expressed during growth on “heterotrophic” substrates, whereas the second is specifically induced by the C₃ compound methylamine, and by choline, the oxidation of which also results in the generation of formaldehyde (Levering et al. 1984b). If this second assumption were correct it then follows that strain Art 98 must be blocked in the synthesis of the “heterotrophic” transaldolase. With the data available it was not possible to discriminate between these two possible explanations. We therefore decided to purify the transaldolases, from gluconate-grown cells of wild type *Arthrobacter* P1 and from methylamine-grown cells of strain Art 98, and to compare their properties.

**Purification of transaldolases**

The typical elution profiles obtained during DEAE-polyl chromatography of cell-free extracts of methylamine- or gluconate-grown cells of wild type *Arthrobacter* P1 and methylamine-grown cells of Art 98 are shown in Fig. 3 A - C, respectively. Interestingly, the transaldolase activity of methylamine-grown cells of *Arthrobacter* P1 was separated in two distinct peaks of activity, whereas only one activity peak was observed in the two other preparations. The data in Fig. 3 further demonstrate that the major peak of activity found with methylamine-grown cells of wild type *Arthrobacter* P1 (Fig. 3 A) was eluted from the column in
Fig. 3A–C. DEAE-polyol chromatography elution profiles of cell-free extracts of A methylamine- or B gluconate-grown cells of wild type *Arthrobacter* P1 and C methylamine-grown cells of mutant Art 98. The transaldolase specific activities in these cell-free extracts were A 1,397, B 640, C 1,000 nmol · min⁻¹ · mg⁻¹ of protein. ---, absorbance at 280 nm; ----, sodium chloride gradient; ●, transaldolase.

Fig. 4A, B. Gel permeation chromatography elution profiles of the pooled peak fractions obtained during DEAE-polyol chromatography of cell-free extracts of A gluconate-grown cells of wild type *Arthrobacter* P1 and B methylamine-grown cells of Art 98. ---, absorbance at 280 nm; ●, transaldolase.

Fig. 5. Stability of transaldolase purified from gluconate-grown cells of *Arthrobacter* P1 (○) or methylamine-grown cells of Art 98 (●) during incubation at 50°C for various periods of time.

Exactly the same fractions as the single peak found with methylamine-grown cells of mutant Art 98 (Fig. 3C). The same is true for the minor peak of activity in Fig. 3A and the single peak found with the gluconate-grown cells of wild type *Arthrobacter* P1 (Fig. 3B). Several runs with extracts from gluconate-grown cells of *Arthrobacter* P1 and methylamine-grown cells of Art 98 were performed and the pooled peak fractions obtained were combined, concentrated and subsequently subjected to gel permeation chromatography. The typical elution profiles obtained with each of these preparations are shown in Fig. 4A and B, respectively. Again several runs were performed and the fractions containing the highest transaldolase activities were pooled and combined. The two preparations thus obtained were subsequently used for comparative studies of the two transaldolases.

Purity of the transaldolases. The specific activities of the transaldolase preparations obtained from gluconate-grown cells of *Arthrobacter* P1 and methylamine-grown cells of Art 98 were 16.0 and 27.5 μmol · min⁻¹ · mg⁻¹ protein, respectively. This means that overall a 25- and 27.5-fold purification, respectively, had been achieved (see Table 2).
Activities of the three other enzymes operative in the pentose phosphate pathway/RuMP cycle in *Arthrobacter* P1, namely transketolase, ribose-5-phosphate isomerase and ribulose-5-phosphate 3-epimerase, were also detected in these transaldolase preparations. Compared to the transaldolase activities, however, the levels of these enzymes were rather low. This is especially the case when the relative activities of these enzymes in the cell-free extracts and the transaldolase preparations are considered (Table 2). It is therefore concluded that also with respect to these functionally closely related enzymes a considerable degree of purification of the transaldolases had been achieved.

### Properties of the transaldolases

1. **pH optima.** Enzyme activities were determined over a range of pH values between 6.0 and 10.0. Both enzyme preparations reached maximum activities around pH 8.0, with very little change in values between pH 7.8 and 8.2. Similar observations have been reported for transaldolase purified from yeast (Venkataraman and Racker 1961a; Tsolas and Horecker 1972).

2. **Enzyme kinetics.** The apparent $K_m$ values of the partially purified transaldolases for fructose-6-phosphate and erythrose-4-phosphate were estimated from Lineweaver-Burk plots. It was noticed, however, that the orthophosphate present in these preparations inhibited the activities of both transaldolases. This (reversible) inhibition became especially evident when, at submaximal concentrations of one of the substrates for transaldolase, larger amounts of the enzymes, and thus of phosphate, had to be added to the assay mixtures in order to obtain sufficiently high rates of product formation. The inhibitory effect of high phosphate concentrations on transaldolase activity is a well-known phenomenon and has for instance been reported for the yeast enzyme (Tsolas and Horecker 1972). Following the removal of phosphate (see Materials and methods) the apparent $K_m$ values of the enzyme purified from strain Art 98 (methylamine-grown) for fructose-6-phosphate and erythrose-4-phosphate were estimated as 2.3 mM and 0.10 mM, respectively. Comparable values, 2.1 mM and 0.07 mM, respectively, were derived for the enzyme purified from wild type *Arthrobacter* P1 (gluconate-grown). When these kinetic studies were performed with transaldolase purified from Baker’s yeast (Sigma Chemical Co., St. Louis, MO, USA) $K_m$ values (1.2 mM and 0.08 mM, respectively) slightly higher than reported in the literature (Venkataraman and Racker 1961a) were found.

### Table 2

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Percentage activity</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Arthrobacter</em> P1 gluconate-grown</td>
<td></td>
</tr>
<tr>
<td>A Transaldolase</td>
<td>100</td>
</tr>
<tr>
<td>B Transaldolase</td>
<td>100</td>
</tr>
<tr>
<td>(2) Transketolase</td>
<td>68</td>
</tr>
<tr>
<td>(3) Ribulose-5-phosphate 3-epimerase</td>
<td>162</td>
</tr>
<tr>
<td>(4) Ribose-5-phosphate isomerase</td>
<td>63</td>
</tr>
<tr>
<td>Hexulose phosphate synthase</td>
<td>0</td>
</tr>
<tr>
<td><em>strain Art 98 methylamine-grown</em></td>
<td></td>
</tr>
<tr>
<td>A Transaldolase</td>
<td>100</td>
</tr>
<tr>
<td>B Transaldolase</td>
<td>100</td>
</tr>
</tbody>
</table>

3. **Molecular weights.** Using the HPLC gel permeation technique (see Materials and methods) molecular weights of 47,500 (± 2,000) and 52,000 (± 2,000) were determined for the enzymes purified from Art 98 and wild type, respectively. When using the same method, baker’s yeast transaldolase appeared to have a molecular weight of about 64,000, which is in the same order of magnitude as reported for this enzyme by Venkataraman and Racker (1961b). When the transaldolase preparations of *Arthrobacter* P1 and Art 98 were subjected to SDS-polyacrylamide gel electrophoresis one major band in the 50,000 dalton region was observed in each case, indicating that both enzymes are monomeric in nature.

4. **Effects of temperature.** With both preparations transaldolase activities were found to increase with temperatures up to 40°C. When the temperature was further raised, maximal activities were observed in the range of 45–50°C with the enzyme isolated from the wild type, whereas the enzyme from Art 98 became very sensitive to inactivation. In the latter case no clear temperature optimum for the transaldolase reaction could be determined. The observed difference in temperature sensitivity was further investigated by incubating small amounts of the enzyme solutions, 20 μl containing approx. 3 μg of protein, in stoppered glass tubes at 50°C for various periods of time. After cooling on ice the remaining transaldolase activities in these samples were determined at 30°C. The results of this experiment (Fig. 5) indicated that, within 2 min of incubation at 50°C, the enzyme purified from Art 98 (methylamine-grown) lost 90% of its activity. No activity was detectable anymore after 10 min of incubation. In contrast, the enzyme isolated from *Arthrobacter* P1 (gluconate-grown) remained fully active when incubated at 50°C for prolonged periods of time. Interestingly, when a crude cell-free extract prepared from methylamine-grown cells of *Arthrobacter* P1 was subjected to this temperature treatment a rapid decrease in transaldolase activity, to approximately 40% of the initial level, was only observed in the first 2 min of incubation. No further loss of activity occurred upon prolonged incubation (data not shown).

### Discussion

In this paper the isolation and characterization of a transaldolase-minus mutant of *Arthrobacter* P1, designated Art...
98, is reported. Because of this mutation it has lost the ability to grow on gluconate and on xylose, substrates which are exclusively metabolized via the pentose phosphate pathway (Fig. 1; Levering et al. 1982). The observation that this mutant is still able to grow in mineral media with glucose, glycerol or succinate as carbon- and energy sources, albeit at lower rates compared to the wild type strain (Table 1), indicates that transaldolase is not really essential for growth of \textit{Arthrobacter P1} on these substrates. Under these conditions the intermediates of the pentose phosphate pathway required for biosynthetic reactions, i.e. ribose-5-phosphate and erythrose-4-phosphate, apparently are synthesized in strain Art 98 via 6-phosphogluconate/ribulose-5-phosphate and fructose-6-phosphate/glyceraldehyde-3-phosphate, respectively (Fig. 1). In contrast, mutants of \textit{E. coli} lacking the functionally closely related enzyme transketolase are not only unable to grow on pentose sugars, but also in media with glucose or other carbon substrates unless supplemented with aromatic amino acids or a precursor, shikimic acid (Fraenkel and Vinopal 1973; Josephson and Fraenkel 1969, 1974). More complex but essentially similar observations have been made with transketolase-minus mutants of \textit{Salmonella typhimurium} (Eidels and Osborn 1971) and \textit{Bacillus subtilis} (Sasajima and Yoneda 1974; Sasajima and Kumada 1979, 1981).

During growth of \textit{Arthrobacter P1} on methylamine, the presence of transaldolase activity is considered essential for the regeneration of the formaldehyde-acceptor molecule ribulose-5-phosphate (Fig. 1; Levering et al. 1982). The observation that mutant Art 98, gluconate-negative because of a deficiency in transaldolase, was still able to grow on methylamine and did synthesize high levels of transaldolase under these conditions was rather puzzling at first (Table 1; Fig. 2). In order to investigate the possible synthesis of transaldolase isoenzymes in \textit{Arthrobacter P1} we decided to (partially) purify the enzymes present in gluconate-grown cells of \textit{Arthrobacter P1} and methylamine-grown cells of strain Art 98. Their subsequent characterization revealed that they have certain features in common, i.e. pH optima, kinetic parameters and inhibition by orthophosphate, but that they are clearly different in other respects. Slight but reproducible differences were found in their behaviour during the DEAE-polychromatography (Fig. 3 B, C) and in the molecular weights of the enzymes (Fig. 4 A, B). As was reported for two transaldolase isoenzymes in the yeast \textit{Candida utilis} (Tsolas and Horecker 1970), the enzymes from \textit{Arthrobacter P1} and Art 98 showed a significant difference in sensitivity towards heat treatment. When incubated at 50°C the enzyme from Art 98 (methylamine-grown) rapidly lost its activity, whereas the enzyme from \textit{Arthrobacter P1} (gluconate-grown) remained stable (Fig. 5). In addition, the transaldolase activity present in methylamine-grown cells of \textit{Arthrobacter P1} wild type could be separated in two distinct peaks of activity during DEAE-poly chromatography (Fig. 3A) and became only partially inactivated during incubation at 50°C. On the basis of these data it is concluded that wild type \textit{Arthrobacter P1} is able to synthesize two transaldolase isoenzymes. These data, and the observations made in the substrate transition experiments, also provide evidence for the conclusion that the enzyme purified from gluconate-grown cells of \textit{Arthrobacter P1} is synthesized constitutively whilst the other enzyme, purified from methylamine-grown cells of Art 98, is specifically induced in the presence of methylamine. It follows then that Art 98 has lost the ability to synthesize the transaldolase which normally becomes expressed constitutively.

Although the two transaldolases in \textit{Arthrobacter P1} are structurally apparently somewhat different, the available evidence indicates that they are functionally very similar, as is the case in \textit{C. utilis} (Tsolas and Horecker 1972). This follows most clearly from the fact that, in the presence of methylamine as the sole nitrogen source, Art 98 is still able to utilize gluconate, reaching the same doubling time as normally observed with wild type \textit{Arthrobacter P1} (Table 1). The major difference between these two transaldolase enzymes therefore lies in the regulation of their synthesis. The additional recruitment via induction of a "methylotrophic" transaldolase during growth on \textit{C}_{1} compounds may simply serve to sufficiently enhance the flow of carbon via the rearrangement reactions in order to ensure continuous availability of the \textit{C}_{1}-acceptor molecule ribulose-5-phosphate under these conditions. In view of the large differences in the activities of the enzymes of the non-oxidative pentose phosphate pathway/RuMP cycle (Tables 1 and 2; Levering et al. 1982) in methylamine- and gluconate-grown cells, it may well be that \textit{Arthrobacter P1} is able to additionally induce a complete set of these enzymes during growth on \textit{C}_{1} compounds.

The presence of isoenzymes among methylotrophic microorganisms is not a rare phenomenon (Bellion and Woodson 1975; Harder and Quayle 1971; Müller-Kraft and Babel 1983; O’Connor and Hanson 1975; this paper). As is the case in \textit{Arthrobacter P1}, most of these "methylotrophic" isoenzymes clearly differ from the enzymes synthesized under "heterotrophic" growth conditions in the way they are regulated, either at the level of their synthesis or at the level of existing enzyme activity, or both. Since during methylotrophic and heterotrophic growth these enzymes fulfill different metabolic roles, the occurrence of these isoenzymes is clearly essential and apparently serves as a further mechanism of fine control of metabolic fluxes.

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