Molecular Cloning, Expression, and Properties of an Unusual Aldo-Keto Reductase Family Enzyme, Pyridoxal 4-Dehydrogenase, That Catalyzes Irreversible Oxidation of Pyridoxal* 

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Microbacterium luteolum YK-1 has pyridoxine degradation pathway I. We have cloned the structural gene for the second step enzyme, pyridoxal 4-dehydrogenase. The gene consists of 1,026-bp nucleotides and encodes 342 amino acids. The enzyme was overexpressed under cold shock conditions with a coexpression system and chaperonin GroEL/ES. The recombinant enzyme showed the same properties as the M. luteolum enzyme. The primary sequence of the enzyme was 54% identical with that of d-threo-aldose 1-dehydrogenase from Agrobacterium tumefaciens, a probable aldo-keto reductase (AKR). Upon multiple alignment with enzymes belonging to the 14 AKR families so far reported, pyridoxal 4-dehydrogenase was found to form a new AKR superfamily (AKR15) together with A. tumefaciens d-threo-aldose 1-dehydrogenase and Pseudomonas sp. l-fucose dehydrogenase. These enzymes belong to a distinct branch from the two main ones found in the phylogenic tree of AKR proteins. The enzymes on the new branch are characterized by their inability to reduce the corresponding lactones, which are produced from pyridoxal or sugars. Furthermore, pyridoxal 4-dehydrogenase prefers NAD+ to NADP+ as a cofactor, although AKRs generally show higher affinities for the latter.

Pyridoxal 4-dehydrogenase (EC 1.1.1.107) catalyzes the oxidation of pyridoxal (pyridoxal hemiacetal) to 4-pyridoxolactone with NAD+ (Fig. 1) and is the second enzyme in degradation pathway I for pyridoxine, one of the free forms of vitamin B6, in which pyridoxine is degraded through eight enzyme-catalyzed steps to succinic semialdehyde, ammonia, and carbon dioxide (1). Pyridoxal 4-dehydrogenase has been purified partially and homogeneously from Pseudomonas MA-1 (2) and Microbacterium luteolum (3), respectively. The enzyme only catalyzes the dehydrogenation (oxidation) reaction on pyridoxal, and no reverse reduction of 4-pyridoxolactone has been observed (2, 3), although due to the low purity of the enzyme preparation (2) and the limited usage of the purified preparation (3), reexamination of the results is required. The reaction specificity suggests that the enzyme is a member of the aldehyde dehydrogenase family (4). However, the amino-terminal amino acid sequence of the enzyme protein showed no homology to the proteins in this family (3).

The aldo-keto reductases (AKRs)3 comprise one of the three enzyme superfamilies that encompass NAD(P)(H)-dependent oxidoreductases (5), which can use aldehydes as substrates. They catalyze the reduction of aldehydes, ketones, monosaccharides, ketosteroids, and prostaglandins. Although AKRs also catalyze the oxidation of hydroxysteroids and trans-dehydrodiols of polycyclic aromatic hydrocarbons, they generally show much higher activity as to the reduction of aldehydes than the dehydrogenation (oxidation) of the corresponding alcohols (6). This superfamily contains more than 100 enzymes, which are divided into 14 families (AKR1–AKR14). They possess the (αβ) eight-barrel motif characteristic of triose-phosphate isomerases and contain ~320 amino acids/monomer. The majority of known AKRs are monomeric. However, multimeric forms are found in the AKR2, AKR6, and AKR7 families.

Here we have cloned the gene encoding pyridoxal 4-dehydrogenase from M. luteolum YK-1, developed an overexpression system for the recombinant pyridoxal 4-dehydrogenase in Escherichia coli, and examined the properties of the purified recombinant enzyme. M. luteolum pyridoxal 4-dehydrogenase exhibits 54% amino acid sequence identity with the probable AKR protein d-threo-aldose 1-dehydrogenase (EC 1.1.1.122) (7) and less than 40% similarity with other AKRs. Pyridoxal 4-dehydrogenase forms a new AKR family (AKR15) and belongs to a new branch, distinct from the two main branches, in the phylogenetic tree of AKR proteins (5).

**EXPERIMENTAL PROCEDURES**

**Bacterial Strains, Plasmids, and Cultivation—**Microbacterium luteolum YK-1 (IFO 16738) isolated from soil was aerobically cultured at 30 °C in pyridoxine medium as described previously (3) and used as a DNA donor and for preparation of M. luteolum pyridoxal 4-dehydrogenase. E. coli JM109 (TaKaRa Bio) was used for cloning of the enzyme-coding gene or for expression of the recombinant enzyme, and E. coli BL21(DE3) (Novagen) was used for overexpression of the enzyme. E. coli cells were cultured in LB medium (1% polypeptone, 0.5% yeast extract, and 1% sodium chloride (w/v)) containing an adequate antibiotic or combination of antibiotics. Plasmids p3T (Mo Bi Tec), pET21a (Novagen), and pTrc99A (Amersham Biosciences) were used as the cloning and expression vectors, respectively. Plasmid pKY206 carrying groEL/ES genes (8) was obtained from Dr. M. Ashiuchi (Kochi University) and Dr. Y. Kawata (Tottori University). This plasmid was used as a coexpression vector.

**Determination of Internal Amino Acid Sequence of Purified M. luteolum Pyridoxal 4-Dehydrogenase—**The M. luteolum enzyme was purified as described previously (3). The purified enzyme or the recomb-
nant enzyme prepared as described below (74 µg) was digested at 25 °C for 26 h with V-8 protease in a reaction mixture comprising 0.1 M Tris-HCl, pH 7.8, containing 2 M urea. The peptides were separated on an ODS-AM (YMC, 100 × 4.6 mm, 120 Å) reverse-phase column. Amino acid sequencing of the peptides was performed with an Applied Biosystems 492 protein sequencer.

Cloning of the Pyridoxal 4-Dehydrogenase Gene—Nearly half of the gene (pld1) encoding pyridoxal 4-dehydrogenase, which encodes the amino-terminal part of the enzyme protein, was amplified by PCR. Chromosomal DNA of M. luteolum YK-1 prepared by the method of Saito and Miura (9) was used as the template. The degenerate PCR primers designed based on the amino-terminal and internal sequences of the enzyme shown in Fig. 1 were 5'-GA/A/GA/A/GC/GG/GT/C/G(G/A/GT/C/GT/C/G/AC/GC)-3' and 5'-AA/A/G/G/A/GC/GG/GT/C/GC/G/CT/G/C/GT/C/GT/GT/C/GT/GT/CT/3', respectively. The PCR conditions, with TaKaRa La Taq polymerase and GC buffer, were essentially the same as those described previously (10). The nucleotide sequence of the amplified DNA fragment (507 bp from the 5'-end of pld1) was determined with an Applied Biosystems 373 DNA sequencer.

The inverse PCR method was used for cloning the residual region of the pld1 gene. The chromosome DNA of M. luteolum was digested with EcoRI, and then EcoRI-digested fragments were self-ligated to produce circular fragments. The latter were used as templates for PCR with primers 5'-AGGGCAGGTTCCATCGGGA-3' (complementary to the antisense strand at nucleotide 484) and 5'-GGCATTTCCGACCAAGGCGC-3' (complementary to the sense strand at nucleotide 77).

The 3'-end of the pld1 gene was cloned by cassette ligation-mediated PCR with a TaKaRa LA PCR in vitro cloning kit. PCR was performed essentially in the same way as described previously (10). The DNA fragment (about 3 kb) containing the 3'-end was amplified. Nucleotide sequencing of the DNA fragment clarified the entire sequence of the pld1 gene.

The pld1 gene was amplified by PCR with the chromosomal DNA of M. luteolum as a template. Primers 5'-CATATGCGATTGAGCATCGGAG-3' and 5'-GGATCTCTGTGGGCTGGA-3' with a NdeI site (underlined in the former) and a BamHI site (underlined in the latter), respectively, were used. The amplified DNA fragment was ligated into pET3 for TA cloning. The constructed plasmid, pETPLD, was introduced into and extracted from E. coli JM109. After the sequence of the introduced fragment had been verified, pETPLD was digested with Ndel and BamHI, and then the digested DNA fragment was inserted in the Ndel/BamHI sites of pET21a. The constructed plasmid, designated as pETPLD, was introduced into E. coli BL21(DE3). pETPLD was digested with XbaI and HindIII to obtain a DNA fragment containing the Shine-Dalgarno sequence in pET21a and the pld1 gene. This fragment was inserted into the XbaI/HindIII sites of pTrc9A to construct pTrPLD, which was then introduced into E. coli JM109.

Expression of Pyridoxal 4-Dehydrogenase and Preparation of Crude Extracts—BL21(DE3)/pETPLD and JM109/pTrPLD cells were cotransformed with plasmid pTri206 carrying the GroEL/ES gene that encodes chaperonins, GroEL and ES, and increases the amount of the active form of the recombinant enzyme in the host cells (8). The cotransformants were grown in LB broth containing ampicillin (50 µg/ml) plus tetracycline (25 µg/ml) for 48 h at 37 °C. To examine the effect of induction with isopropyl-1-thio-β-D-galactopyranoside, 1 mM isopropyl-1-thio-β-D-galactopyranoside was added to the culture when the absorbance of 600 nm reached 0.6–0.8.

Cells were harvested and washed with 50 mM potassium phosphate buffer, pH 8.0, and then suspended in 20 mM potassium phosphate buffer, pH 8.0, containing 0.1% (v/v) 2-mercaptoethanol, 10% (w/v) glycerol, 1 mM EDTA, and 1 mM phenylmethylsulfonyl fluoride (Buffer A containing phenylmethylsulfonyl fluoride). The cell suspension was thoroughly sonicated at 4–15 °C with a Heat Systems-Ultrasonics sonicator W-220. The precipitate containing cell debris and insoluble materials was removed by centrifugation at 10,000 × g for 10 min at 4 °C, and the supernatant was used as the soluble fraction. The precipitate was mixed with Buffer A containing 6 M urea, and then the mixture was kept on ice for 2 h and centrifuged to remove insoluble materials. The supernatant containing the dissolved inclusion proteins was used as the dissolved fraction.

Purification of Recombinant Pyridoxal 4-Dehydrogenase—The cotransformant cells (4.7 g), BL21(DE3)/pETPLD/pKY206, were suspended in 25 ml of Buffer A containing 1 mM phenylmethylsulfonyl fluoride. The suspension was sonicated on ice for 15 min with a Heat Systems-Ultrasonicator W-220. The supernatant (40 ml) obtained on centrifugation at 10,000 × g for 20 min at 4 °C was used as the crude extract. The crude extract was fractionated with ammonium sulfate, and the precipitate obtained upon centrifugation of the 40–70% saturated solution was dissolved in 14 ml of Buffer A. This solution, after dialysis against Buffer A, was applied to a Blue A (Amicon) column (1.8 × 8.0 cm) equilibrated with Buffer A. The column was washed with Buffer A and then Buffer A containing 0.1 M NaCl until the elution volume of the blue band became lower than 0.1. The enzyme was eluted with Buffer A containing 0.2 M NaCl as a single peak.

Enzyme and Protein Assays—Pyridoxal 4-dehydrogenase was assayed during its purification by measuring the increase in fluorescence (excitation at 356 nm and emission at 432 nm) of 4-pyridoxolactone produced as described previously (3), for which 2–100 milliunits of the enzyme was routinely used. One unit of enzyme was defined as the amount that catalyzed the formation of 1 nmol of 4-pyridoxolactone per min (3). The reverse reaction was assayed by the phenylhydrazine method, in which pyridoxal produced was measured as phenylhydrazine (11), because a high concentration of 4-pyridoxolactone, which shows maximum absorbance at 356 nm, interfered with measurement of the decrease in the absorption of NADH. The reaction was performed in a reaction mixture (0.4 ml) consisting of 50 mM sodium phosphate buffer (pH 8.0), 50 mM 4-pyridoxolactone, 10 mM NADH, and 5 units of enzyme at 30 °C for 1 h. The reaction was stopped by the addition of 66 µl of 9 M sulfuric acid. The reaction mixture (0.3 ml) after centrifugation was mixed with 0.6 ml of 1 M sulfuric acid and 0.1 ml of 2% (w/v) phenylhydrazine (dissolved in 5 mM sulfuric acid), and then the mixture was incubated at 60 °C for 20 min. The A410 of the mixture was measured. Protein was measured by the dye-binding method with bovine serum albumin as a standard (12).

Substrate Specificity of the Recombinant Enzyme—The enzyme activity toward sugars and the K_{m} values were determined by measuring the initial increase in A_{340} of NADH at 30 °C in 1.0 ml of a reaction mixture consisting of 50 mM sodium phosphate buffer (pH 8.0), 250 mM each sugar, 1.0 mM NAD{+}, and the enzyme. The reactivity with aldehydes, which are typical synthetic AKR substrates, was measured in the same reaction mixture containing 1.0 mM sodium phosphate buffer (pH 8.0), 50 mM 4-pyridoxolactone, 10 mM NADH, and 5 units of enzyme at 30 °C for 1 h. The reaction was stopped by the addition of 66 µl of 9 M sulfuric acid. The reaction mixture (0.3 ml) after centrifugation was mixed with 0.6 ml of 1 M sulfuric acid and 0.1 ml of 2% (w/v) phenylhydrazine (dissolved in 5 mM sulfuric acid), and then the mixture was incubated at 60 °C for 20 min. The A_{410} of the mixture was measured. Protein was measured by the dye-binding method with bovine serum albumin as a standard (12).

Other Analytical Methods—The molecular weight of the recombinant enzyme was determined by gel filtration (Hiprep 16/60 Sephacyr S-300 high resolution column) with a fast protein liquid chromatograph. A calibration curve was made based on the elution pattern of catalase (Mr = 240,000), lactate dehydrogenase (Mr = 140,000), alanine aminotransferase (Mr = 110,000), malate dehydrogenase (Mr = 72,000), and cytochrome c (Mr = 12,400). The subunit molecular weight was measured by SDS-PAGE (13) with the molecular weight markers as described previously (10).
RESULTS

Cloning of the M. luteolum Pyridoxal 4-Dehydrogenase Gene—
The amino acid sequences of the amino-terminal region and three internal peptides generated on V-8 protease digestion were determined (solid underlines in Fig. 2a). Based on the amino acid sequence information, degenerate primers were
which was almost the same value as that for (Fig. 3, the formation of large amounts of insoluble enzyme proteins pETPLD cells are shown in Fig. 3. Both types of cells showed two types of transformant cells, only the results for BL21(DE3)/

terns of the protein bands on the gel were very similar for the SDS-polyacrylamide gel (Fig. 3, protein was so low that the protein band could not be seen in the pyridoxine synthetic medium. The amount of the enzyme

pld1 predicted molecular mass was 37,889, which was in good verified (Fig. 2

was amplified (Fig. 2

B

A

37380

extracts of recombinant pKY206 cells; lane D, the purified recombinant enzyme (2.2 µg). The standard proteins were applied to lane Sd.

designed for PCR, which corresponded to the amino acids indicated by dotted lines in Fig. 2A. The pld1 gene fragment comprising nucleotides 19–528 was amplified by PCR (thick line in Fig. 2B). Upon inverse PCR with the circularized EcoRI-digested fragments, a 550-bp fragment was amplified, which contains the 5’-end and the central part of the pld1 gene (solid line in Fig. 2B). Finally, the 3’-terminal part of the pld1 gene was amplified by cassette ligation-mediated PCR, as shown by the double line in Fig. 2B. The entire pld1 gene was amplified (Fig. 2B), and the nucleotide sequence was verified (Fig. 2A). The pld1 gene consisted of 1,026 bp nucleotides, which encoded 342 amino acid residues (Fig. 2A). The predicted molecular mass was 37,889, which was in good agreement with the molecular mass (38,000) determined previously by SDS-PAGE (3).

Expression of the pld1 Gene in E. coli—The expression of the pld1 gene in various E. coli transformants was examined with cells cultured in a small volume of medium (5.0 ml) under different conditions. BL21(DE3)/pETPLD and JM109/pTRPLD cells grown at 37 °C showed low pyridoxal 4-dehydrogenase activity (0.25 ± 0.05 and 0.26 ± 0.02 units/mg, respectively), which was almost the same value as that for M. luteolum grown in the pyridoxine synthetic medium. The amount of the enzyme protein was so low that the protein band could not be seen in the SDS-polyacrylamide gel as a solid line (Fig. 3, lane A). Because the pattern of the protein bands on the gel were very similar for the two types of transformant cells, only the results for BL21(DE3)/pETPLD cells are shown in Fig. 3. Both types of cells showed the formation of large amounts of insoluble enzyme proteins (Fig. 3, lane B). To increase the soluble and active forms of the enzyme protein, BL21(DE3)/pETPLD cells were cultured under different conditions, such as cultivation at high (44 and 42 °C) or low (20 °C) temperature and induction with isopropyl-1-thio-β-D-galactopyranoside. Only cultivation at the low temperature was effective, the specific activity being increased to 1.28 ± 0.31 units/mg. Because it has been reported that the coexpression of E. coli chaperonin proteins increases the soluble forms of several enzymes (8, 14), we examined the effect of their coexpression on pyridoxal 4-dehydrogenase in the transformant cells. The specific activity of the enzyme in BL21(DE3)/pETPLD/pKY206 cells and JM109/pTRPLD/pKY206 cells, which were cultured without isopropyl-1-thio-

β-D-galactopyranoside. Only cultivation at the low temperature was effective, the specific activity being increased to 1.28 ± 0.31 units/mg. Because it has been reported that the coexpression of E. coli chaperonin proteins increases the soluble forms of several enzymes (8, 14), we examined the effect of their coexpression on pyridoxal 4-dehydrogenase in the transformant cells. The specific activity of the enzyme in BL21(DE3)/pETPLD/pKY206 cells and JM109/pTRPLD/pKY206 cells, which were cultured without isopropyl-1-thio-

β-D-galactopyranoside at 23 °C, was 11.8 ± 3.24 and 14.6 ± 1.55 (i.e. 9.2- and 8.8-fold higher than the values for BL21(DE3)/pETPLD cells and JM109/pTRPLD cells, respectively). The amount of the soluble form of the enzyme was high enough to see it in the SDS-polyacrylamide gel as a solid line (Fig. 3, lane C). Thus, cold stress and cooperative expression of chaperonins increased the soluble and active forms of pyridoxal 4-dehydrogenase in the host cells. Both types of cells could be used for purification of the enzyme.

Purification and Properties of Recombinant Pyridoxal 4-Dehydrogenase—The recombinant pyridoxal 4-dehydrogenase was purified to homogeneity from BL21(DE3)/pETPLD/pKY206 cells by one-step column chromatography (Fig. 3, lane D, and Table I). The specific activity of the crude extract used as the starting material was lower than that observed in the transformant cells cultivated in a small volume (5 ml). For an unknown reason, the transformant cells showed lower specific activity when they were grown in a larger volume (200 ml) for preparation of the enzyme.

### Table I

| Fraction            | Total protein (mg) | Total activity (units) | Specific activity (units/mg) | Yield (%) |
|---------------------|--------------------|------------------------|------------------------------|-----------|
| Crude extract       | 414                | 2,700                  | 6.5                          | 100       |
| Ammonium sulfate    | 251                | 2,680                  | 11.6                         | 99.4      |
| Blue A              | 2.6                | 1,670                  | 635                          | 62.3      |

### Table II

| Compound          | $K_m$ (mM) | $K_{cat}$ (s⁻¹) | $K_{cat}/K_m$ (s⁻¹ mM⁻¹) |
|-------------------|------------|-----------------|------------------------|
| Pyridoxal         | 13 ± 2.5   | 59.2 ± 7.3      | 4.6                    |
| L-Fucose          | 0.41 ± 0.05| 125 ± 2.0       | 304.9                  |
| D-Arabinose       | 0.91 ± 0.08| 175 ± 1.9       | 192.3                  |
| L-Xylose          | 3.5 ± 0.19  | 214 ± 1.4       | 61.0                   |
| NAD⁺              | 0.042 ± 0.003|               |                       |
| NADP⁺             | 0.58 ± 0.14 |               |                       |
The purified recombinant enzyme showed 1.8-fold higher specific activity than the *M. luteolum* enzyme. However, the optimum pH (pH 8.0), the optimum temperature (30 °C), the thermostability (75% inactivation at 45 °C for 10 min), and the molecular weights both on gel filtration (84,000 ± 4,200) and SDS-PAGE (38,000 ± 600) were the same for the recombinant and native enzymes. Pyridoxal 4-dehydrogenase is a dimeric protein. The amino-terminal 10-amino acid sequence and the sequences of five V8-cleaved peptides derived from the recombinant enzyme were determined (the double lines in Fig. 2A). Whereas the amino-terminal methionine was removed in the recombinant enzyme, the determined amino acid sequences coincided with those deduced from the nucleotide sequences.

Activity toward Sugars and Aldehydes—Because the primary sequence of pyridoxal 4-dehydrogenase was similar to other aldose 1-dehydrogenases, we examined its activity toward a variety of carbohydrates (Fig. 4). The enzyme showed high activity toward L-fucose, glucose, fructose, and 2-deoxy-D-glucose. The rates of oxidation of these sugars were consistent with their reducing power. The enzyme was also active toward the aldehyde 2-deoxy-D-gluceraldehyde. The Michaelis constants (Km) for L-fucose, glucose, fructose, and 2-deoxy-D-glucose were 0.35, 0.30, 0.30, and 0.30 mM, respectively. The enzyme was not active toward other sugars, such as D-glucose, D-fructose, or L-fucose. The enzyme was not active toward the aldehyde 2-deoxy-D-gluceraldehyde. The Michaelis constants (Km) for L-fucose, glucose, fructose, and 2-deoxy-D-gluceraldehyde were 0.35, 0.30, 0.30, and 0.30 mM, respectively. The enzyme was not active toward other sugars, such as D-glucose, D-fructose, or L-fucose. The enzyme was not active toward the aldehyde 2-deoxy-D-gluceraldehyde.
that of a putative D-threo-aldose 1-dehydrogenase, the reactivity of the enzyme with sugars was examined. The enzyme showed high activity toward L-fucose, D-arabinose, and L-xylose (Table II). These substrates have hydroxyl groups at the second and third carbons in the threo configuration. The substrates showed Michaelis-Menten type kinetics. L-Fucose was the best substrate, having the highest specificity constant.

When L-fucose was used as the hydrogen donor, the optimum pH of the enzyme reaction was pH 9.5 (Fig. 4). In contrast, it was pH 8.0 when pyridoxal was used. Whereas carbonate buffers inhibited the dehydrogenation of pyridoxal, they did not inhibit that of L-fucose. The optimum temperature was 45°C, and the enzyme was stable up to 40°C when L-fucose was used as the substrate. Whereas no difference was observed in thermostability between the enzyme reactions with L-fucose and pyridoxal, the optimum temperature was 10°C higher with L-fucose than with pyridoxal. The \( K_m \) values for NAD\(^+\) were 0.042 ± 0.003 and 0.05 ± 0.004 mM with L-fucose and pyridoxal as the substrate, respectively. The \( K_m \) value for NADPH determined with L-fucose was 0.58 ± 0.14 mM.

Pyridoxal 4-dehydrogenase showed no measurable activity toward benzaldehyde, 2-nitrobenzaldehyde or 4-nitrobenzaldehyde although these synthetic aldehydes are good substrates for AKRs so far studied (15).

**DISCUSSION**

*M. luteolum* pyridoxal 4-dehydrogenase shows the highest identity (54%) with a probable D-threo-aldose 1-dehydrogenase in *Agrobacterium tumefaciens* (AAL44626, 348 residues). *Mesorhizobium loti* putative D-threo-aldose 1-dehydrogenase (BAB50246, 332 residues), *Sinorhizobium meliloti* putative protein (AAK65425, 338 residues), *S. meliloti* putative L-fucose dehydrogenase (CAC41406, 339 residues), *M. loti* unknown protein (BAB54355, 340 residues), *Bradyrhizobium japonicum* putative oxidoreductase (BAC46389, 325 residues), *Pseudomonas sp.* L-fucose dehydrogenase (BAA06803, 329 residues), and *Bacillus anthracis* str. Ames oxidoreductase (AAP27231, 336 residues) show identities of 45, 44, 42, 40, 39, 38, and 37%, respectively. Since D-threo-aldose 1-dehydrogenase is a potential member of the AKR superfamily (16), *M. luteolum* pyridoxal 4-dehydrogenase could belong to the AKR superfamily.

The amino acid sequence of pyridoxal 4-dehydrogenase was aligned with those of *A. tumefaciens* D-threo-aldose 1-dehydrogenase, *Pseudomonas sp.* L-fucose dehydrogenase, *C. tenuis* xylose reductase (AKR2B7) (18), and rat aflatoxin dialdehyde reductase (AKR7A1) (19) (Fig. 5). *C. tenuis* xylose reductase and rat aflatoxin dialdehyde reductase are novel dimeric members of the AKR superfamily with defined tertiary
structures. All of the residues in the active sites of AKRs are conserved in pyridoxal 4-dehydrogenase, namely Asp-56, Tyr-61, Lys-86, and His-152, corresponding to Asp-40, Tyr-45, Lys-73, and His-109 in the AKR7A1 protein, respectively (Fig. 5). The amino acid residues involved in NAD(P)⁺ binding are also conserved, namely Gly-250, Arg-256, Gln-315, Gln-318, and Asn-319, corresponding to Gly-198, Arg-204, Glu-290, Glu-293, and Asn-294 in the AKR7A1 protein, respectively (Fig. 5). Therefore, M. luteolum pyridoxal 4-dehydrogenase is a member of the AKR superfamily.

The AKR superfamily, so far, comprises 14 families. A representative was selected from each family and aligned with M. luteolum pyridoxal 4-dehydrogenase, A. tumefaciens threo-aldo-1-dehydrogenase, and Pseudomonas sp. l-fucose dehydrogenase. Enzymes belonging to one family exhibit less than 40% amino acid sequence identity with enzymes belonging to other families (16). No enzymes in the 14 families show more than 40% amino acid sequence identity with pyridoxal 4-dehydrogenase. When an unrooted phylogenetic tree was constructed, pyridoxal 4-dehydrogenase was found to form a distinct branch together with two aldose reductases (Fig. 6A). These two aldose reductases show reactivity with aldoses with hydroxyl groups at the 2- and 3-positions with the three configuration. From these data, we propose a new AKR family, which is composed of pyridoxal 4-dehydrogenase and other aldose reductases. The sequence of pyridoxal 4-dehydrogenase was submitted to the AKR superfamily homepage (available on the World Wide Web at www.med.upenn.edu/akr/), and the new family was designated as AKR family 15.

The phylogenetic tree of the AKR superfamily has two major branches; one branch is composed of AKR families 1–5, and the other is composed of AKR families 6–14 (5) (Fig. 6A). The pyridoxal 4-dehydrogenase family, family 15, forms a third major branch. Enzymes in family 15 are different from the others in several biochemical properties as well. They only catalyze the oxidation (dehydrogenation) reaction and have a dimeric structure, and some of them show higher affinity for NAD⁺ than for NADP⁺. Pyridoxal 4-dehydrogenase showed no reactivity with 4-pyridoxolactone even when the lactone concentration was 50 mM and a 100-fold higher amount of the enzyme was used. The effective irreversibility of the enzyme reaction has also been found for Pseudomonas pyridoxal 4-dehydrogenase (2). l-Fucose dehydrogenase from rabbit liver belonging to the AKR family 15 shows no catalytic activity toward l-fucolactonolactone (20). This irreversibility is attributed to spontaneous hydrolysis of l-fucolactonolactone at basic pH because the enzyme shows an optimum pH of 10.0. However, this is not the case for pyridoxal 4-dehydrogenase, because the enzyme shows an optimum pH of 8.0–8.5, where 4-pyridoxolactone is not easily hydrolyzed. The inability of family 15 enzymes to reduce lactones can cause controversy regarding the molecular evolution of AKRs. Further studies are required to elucidate the mechanism involved.

AKRs generally show only affinity for NADP⁺ or higher affinity for NADP⁺ than for NAD⁺ when both coenzymes are used as cofactors, although a NAD-prefering xylose reductase has recently been characterized (21). Codeinone reductase (AKR4B2), morphine dehydrogenase (AKR5B), pyridoxal reductase (AKR8A1), and aryl-alcohol dehydrogenase (AKR9A3) can only use NADP⁺. ARA1p-arabinose dehydrogenase (AKR3C), shaker channel β-subunit (AKR6A1), and aldehyde reductase (AKR1A2) show 3–10-fold higher affinity for NADP⁺ than for NAD⁺, respectively. On the contrary, pyridoxal 4-dehydrogenase showed 10-fold higher affinity for NAD⁺ than for NADP⁺ as the xylose reductase (21).

Although the majority of AKRs are monomeric, recent studies have shown that xylose reductase (AKR2B5) (6) and aflatoxin-inducible aldehyde reductase (AKR7A1) (22) are dimeric. M. luteolum pyridoxal 4-dehydrogenase in AKR15 was shown to be dimeric. The amino acid residues necessary for dimerization of C. tenuis xylose reductase, Asp-178, Glu-205, Phe-206, Trp-313, and Pro-319, are well conserved as Asp-203, Glu-229, Phe-230, Trp-322, and Pro-328 of pyridoxal 4-dehydrogenase (Fig. 5). These residues in pyridoxal 4-dehydrogenase are likely to be involved in dimer formation.

l-Fucose was a better substrate for pyridoxal 4-dehydrogenase than pyridoxal. Although several l-fucose dehydrogenases (EC 1.1.1.122, d-threo-aldo-1-dehydrogenase) from animals and microorganisms have been characterized, their reactivity with pyridoxal has not been reported. l-Fucose dehydrogenase from porcine liver acts on the β-anomer of l-fucose and catalyzes hydride transfer of the C-1 hydrogen in the lowest energy chair conformation (23). Korytnyk and Singh (24) have found that pyridoxal existed exclusively in the hemiacetal form at pH 1.0–11.0 by examining its proton magnetic resonance spectra. Interestingly, the conformations of pyridoxal hemiacetal and the chair form of l-fucose show several similarities between functional groups (Fig. 6B). When the conformational structures are placed one upon another at C-1 of l-fucose and C-4 of pyridoxal hemiacetal, both of which bind the axial hydrogen to be released, the C-2 oxygen and C-3 oxygen in l-fucose and pyridoxal hemiacetal, respectively, occupy similar positions to the carbons. Furthermore, the C-3 oxygen of l-fucose is located at a similar position to the nitrogen of the pyridine ring in pyridoxal hemiacetal. Thus, it is possible that l-fucose and pyridoxal hemiacetal bind to the active site pocket of the enzyme, where some amino acid residues provide binding sites for these atoms through hydrogen bonds, and that the enzyme shows reactivity with the compounds. Indeed, the optimum pH of pyridoxal was lower than that of l-fucose, suggesting that the C-3 phenolic hydroxyl group in pyridoxal hemiacetal should be in a nondissociated form for binding to the binding site. The substrate-binding pocket of pyridoxal 4-dehydrogenase from M. luteolum may be similar to those of d-threo-aldo-1-dehydrogenases. Comparative studies on d-threo-aldo-1-dehydrogenases from different sources will provide the clues for understanding the molecular mechanism underlying the changes in the substrate specificities of AKR proteins.

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