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**Author Manuscript** 

Bioorg Chem. Author manuscript; available in PMC 2013 February 1.

# Published in final edited form as:

Bioorg Chem. 2012 February ; 40(1): 79-86. doi:10.1016/j.bioorg.2011.09.002.

# Structure and characterization of the 3-deoxy---*arabino*-heptulosonate 7-phosphate synthase from *Aeropyrum pernix*

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# Abstract

The first enzyme in the shikimic acid biosynthetic pathway, 3-deoxy-D-*arabino*heptulosonate 7phosphate synthase (DAH7PS), varies significantly in size and complexity in the bacteria and plants that express it. The DAH7PS from the archaebacterium *Aeropyrum pernix* (DAH7PS<sup>Ap</sup>) is among the smallest and least complex of the DAH7PS enzymes, leading to the hypothesis that DAH7PS<sup>Ap</sup> would not be subject to feedback regulation by shikimic acid pathway products. We overexpressed DAH7PS<sup>Ap</sup> in *Escherichia coli*, purified it, and characterized its enzymatic activity. We then solved its X-ray crystal structure with a divalent manganese ion and phosphoenolpyruvate bound (PDBID: 1VS1). DAH7PS<sup>Ap</sup> is a homodimeric metalloenzyme in solution. Its enzymatic activity increases dramatically above 60°C, with optimum activity at 95°C. Its pH optimum at 60°C is 5.7. DAH7PS<sup>Ap</sup> follows Michaelis-Menten kinetics at 60° C, with a K<sub>M</sub> for erythrose 4-phosphate of 280 µM, a K<sub>M</sub> for phosphoenolpyruvate of 891 µM, and a  $k_{cat}$  of 1.0 s<sup>-1</sup>. None of the downstream products of the shikimate biosynthetic pathway we tested inhibited the activity of DAH7PS<sup>Ap</sup>. The structure of DAH7PS<sup>Ap</sup> is similar to the structures of DAH7PS from *Thermatoga maritima* (PDB ID: 3PG8) and *Pyrococcus furiosus* (PDB ID: 1ZCO), and is consistent with its designation as an unregulated DAH7PS.

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*Aeropyrum pernix*; 3-deoxy-D-*arabino*-heptulosonate 7-phosphate synthase; shikimate pathway; crystal structure; feedback regulation

# **1. INTRODUCTION**

The shikimate pathway produces chorismate in seven steps, starting with D-erythrose 4phosphate (E4P) and phosphoenolpyruvate (PEP) (1, 2). Chorismate is the common precursor for the biosynthesis of the aromatic amino acids phenylalanine, tyrosine and tryptophan, and it is the precursor to the important molecules ubiquinone, and para-amino benzoic acid. The shikimate pathway is essential in plants and microorganisms but absent in mammals, making it a potential target for the development of antibiotics and herbicides. Glyphosate, for example, is a commercially important herbicide that inhibits 5enolpyruvlyshikimate-3-phosphate synthase (also known as EPSP synthase), which catalyzes the sixth step in the pathway (3).

The first enzyme in the shikimate pathway, DAH7PS (EC 4.1.2.15), catalyzes the condensation of E4P and PEP to form 3-deoxy-D-arabino-heptulosonate 7-phosphate and inorganic phosphate. The DAH7PS enzymes are diverse, with some organisms, including Escherichia coli, containing more than one DAH7PS. The structures of all DAH7P enzymes share a core  $(\alpha/\beta)_{8}$  catalytic core, but there are many additions to this core that subject the enzymes to a variety of different regulatory schemes. The DAH7PS from B. subtilis, for example, has an active chorismate mutase domain fused to the N-terminus of the  $(\alpha/\beta)_{R}$ catalytic core. This domain confers feedback inhibition by chorismate and prephenate (4). The DAH7PS from *Thermatoga maritima* (DAH7PS<sup>Tm</sup>, (5)) also has a domain fused to the N-terminus of the  $(\alpha/\beta)_8$  catalytic core that is involved in its regulation by tyrosine and phenylalanine (6). This domain is similar to the ACT domain (7) that regulates the activity of many amino acid metabolizing enzymes (8). The DAH7PS from Mycobacterium tuberculosis presents a much more complex regulatory mechanism. This enzyme has both an N-terminal extension and an added internal loop that provide independent binding sites for regulatory amino acids. Enzyme activity is modulated by combinations of phenylalnine and tryptophan, or tyrosine and tryptophan, but not by the individual amino acids alone. We originally chose to study the DAH7PS<sup>Ap</sup> because it is among the shortest of the bacterial DAH7Ps enzymes, and A. pernyx is a very primitive microbial species. The enzyme's small size (276 amino acids) and lack of N-terminal, C-terminal or internal additions to the basic  $(\alpha/\beta)_8$  catalytic core suggested to us that it would be an unregulated enzyme. Its primitive origin suggested to us that its structure would help unravel the mechanisms by which other enzymes are regulated. We therefore cloned DAH7PS<sup>Ap</sup>, expressed it in *E. coli*, and characterized the purified enzyme. We then solved its X-ray crystal structure with divalent Mn ion and PEP bound. The structure was deposited (PDB ID: 1VS1) in 2006. This paper describes the cloning, expression, and characterization of the enzyme, and discusses its structure in relation to those of other DAH7PS enzymes.

# 2. Experimental Procedures

#### 2.1. Materials

*A. pernix* genomic DNA (ATCC 700893D) was purchased from ATCC (Manassas, VA). The pCR®T7 TOPO®TA expression kit, isopropyl-β-D-thiogalactoside, and the PCR/ sequencing primers were purchased from Invitrogen (Carlsbad, CA). FailSafe® PCR premix selection kit was purchased from Epicentre Technologies (Madison, WI). Plasmid preparations were performed utilizing a Promega (Madison, WI) Wizard DNA purification

kit. XL1-Blue and BL21-CodonPlus®(DE3)-RIL E. coli competent cells were purchased from Stratagene (La Jolla, CA). Restriction enzymes, Vent DNA polymerase, and T4 ligase were purchased from New England Biolabs (Beverly, MA). Enzymatic reactions were performed in an MJ Research PTC-200 Peltier Thermal Cycler. The University of Michigan Biomedical Resources Core Facility performed the DNA sequencing. PEP mono (cyclohexylammonium) salt, E4P sodium salt, chorismate, shikimate, prephenate, and 1,3bis[tris(hydroxymethyl)methylamino]propane (BTP) were obtained from Sigma (St. Louis, MO). KCl and (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> were purchased from Fisher Scientific (Fairlawn, NJ). EDTA disodium salt was obtained from Mallinckrodt (Phillipburg, NJ). Tris base was purchased from Research Organics (Cleveland, OH). High grade Spectra/Por® 7 dialysis tubing (10,000 molecular weight cut-off and metal-free) was obtained from VWR Scientific (Chicago, IL). Centriprep YM-10 concentrators were purchased from Millipore (Billerica, MA). Phenyl Superose (HR10/10) and UNO® Q6 chromatography column were purchased from Amersham Pharmacia Biotech (New York, NY) and Bio-Rad (Richmond, CA), respectively. Bio-Rad Protein Assay Reagent was used to determine the concentration of the protein, with bovine serum albumin (from Sigma) as the standard. The Mini-PROTEAN II electrophoresis unit from Bio-Rad (Richmond, CA) was used to perform SDS-PAGE. The samples were resolved on a 12% polyacrylamide gel and dyed with 0.25% Coomassie brilliant blue R from Aldrich Chemical Company (St. Louis, MO). Optical spectroscopy was performed using an HP 8453 UV-visible spectrometer. All preparations and dialyses were performed at room temperature unless otherwise indicated. All reaction buffers were adjusted to pH 7.0 at the specified reaction temperature, unless otherwise noted.

#### 2.2. Plasmid construction

The gene encoding DAH7PS (gi:14600379) (9) was amplified from *A. pernix* genomic DNA using standard PCR methodologies. The forward primer was 5'-

<u>CATATG</u>TGGAGGTGGCTCCCA GTGGCCG-3', and the reverse primer was 5'-GAATTCAAGCTT<u>GGATCC</u>TACAGCAGCCTGTGCCACC-3'. The PCR product was isolated and ligated into the linear pCR®T7/CT-TOPO® vector according to the manufacturer's specifications. This ligation mixture was used to transform *E. coli* TOP10F' competent cells. Plasmids isolated from these cells were sequenced and one plasmid with the correct sequence was restricted with endonucleases *NdeI* and *Bam*HI (restriction sites underlined above). The restricted product was ligated into a similarly restricted pT7-7 vector that had also been treated with calf alkaline phosphatase. The ligation mixture was used to transform *E. coli* XL1-Blue competent cells. A plasmid isolated from one transformant, containing the correct sequence was used to transform *E. coli* BL21-CodonPlus®(DE3)-RIL competent cells.

#### 2.3. Protein expression and purification

*E. coli* cells harboring the DAH7PS<sup>Ap</sup> plasmid were grown in 2xYT media (100 mg/L ampicillin) or M9 minimal media (100 mg/L ampicillin) with shaking (300 rpm) at 37°C. The cultures were grown until  $A_{600} = 1.4$  and expression was induced by addition of isopropyl- $\beta$ -D-thiogalactoside to a final concentration of 0.4 mM. The *E. coli* in 2xYT media were allowed to grow for an additional 4 hours at 37°C while the *E. coli* in M9 minimal media were allowed to grow for an additional 36 hours at room temperature. *E. coli* cells were pelleted by centrifugation (6000 rpm, 6 min, 4° C). The cell pellet was resuspended in buffer A (20 mM Tris-HCl, pH 8.0 for DAH7PS<sup>Ap</sup>) and sonicated at 4°C in an ice bath (four 30 s pulses with a 2 min rest in between pulses). Cell debris was removed by centrifugation (18000 rpm, 30 min, 4°C). Solid NaCl was added to the supernatant to a final concentration of 0.1 M, and the supernatant was rapidly heated to 80°C in a boiling water bath. The supernatant was heated for an additional 10 minutes in an 80°C water bath with gentle swirling. The denatured proteins were removed by centrifugation (18000 rpm,

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30 min, 4°C), and solid (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> was added to the supernatant to a final concentration of 20% (w/v). The sample was passed through a 0.22 µm filter and loaded onto a Phenyl Superose column (HR 10/10) pre-equilibrated with 20% (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> in buffer A. The protein was eluted at a flow rate of 1 ml/min; first utilizing a reverse gradient from 20% to 0% (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> in buffer A over 60 min followed by buffer A for 15 min. Fractions with DAH7PS activity were pooled and dialyzed overnight against 2 L of buffer A at 4°C. The dialyzed preparation was filtered using a 0.22 µm filter and applied onto a UNO® Q6 column pre-equilibrated with buffer A. The protein was eluted at a flow rate of 2 ml/min using a linear gradient from 0 to 0.5 M potassium chloride in buffer A over 60 min. SDS-PAGE was used to verify the purity of each fraction, and homogeneous fractions containing DAH7PS activity were pooled and dialyzed in 2 L of buffer A overnight at 4°C. The purified enzymes were aliquoted, quickly frozen in an ethanol and dry ice bath, and stored at -80°C.

#### 2.4. Molecular weight determination

The native molecular weight of the enzyme was determined by gel filtration on a Superose 12 column (HR10/30) as previously described (10). Cytochrome c (12.4 kDa), carbonic anhydrase (29 kDa), albumin (66 kDa), alcohol dehydrogenase (141 kDa), and  $\beta$ -amylase (200 kDa) were used as molecular weight standards. Blue dextran (2000 kDa) was used to determine the column void volume. Standards and samples of 5-10 mg/ml were injected into a 50 µl sample loop for holding before being loaded onto the Superose 12 column (HR10/30).

#### 2.5. Activity assay

Enzyme (various concentrations), 3 mM PEP, and 100 mM Tris-acetate (pH 7.0 at the desired reaction temperature) were incubated for 2 min in a thin-walled PCR tube at the desired temperature. The reaction was initiated by the addition of 3 mM E4P, with the final volume being 50  $\mu$ l. The reaction was quenched after 2 min by the addition of 50  $\mu$ l of 10% (v/v) cold trichloroacetic acid. The amount of DAH7P produced was quantified using a modified Aminoff periodate-thiobarbituric acid assay (11). The production of 1  $\mu$ mol of DAH7P per min was defined as one unit of activity.

#### 2.6. Temperature Optimum, pH dependence, and Thermostability

To determine the optimum reaction temperature of the desired enzyme, the enzyme was allowed to react with PEP and E4P in 100 mM Tris-acetate at pH 7.0 at temperatures between 30 and 100°C (10°C increments) utilizing the discontinuous colorimetric assay as described above. The buffer pH was adjusted to 7.0 at the desired temperatures to account for temperature dependent pH changes. For pH dependence studies, the enzymatic activity was measured at 60°C utilizing a series of 100 mM bistris propane buffers in the pH range of 5.0 to 9.0. The pH of the buffers was adjusted at 60°C. Assays were performed as described above. Thermostability was determined by incubating the enzyme in the presence or absence of 1 mM PEP at 90°C for DAH7PS<sup>Ap</sup>. At different time intervals, aliquots were taken. The residual activity of the enzyme was measured using the discontinuous colorimetric assay at the same temperature as its incubation temperature.

#### 2.7. Metal analysis

The identification and quantification of the metal(s) in the enzyme "as isolated" was determined by high-resolution inductively coupled plasma-mass spectrometry on a Finnigan MAT ELEMENT instrument. To determine the effect of metal chelating agents on the enzyme, a mixture of 10 mM EDTA and 1 mM dipicolinic acid (DPA) was incubated with

enzyme, 100 mM Tris-acetate (pH 7.0 at 60° C), and 3 mM PEP for 1.0 hour at room temperature. The residual activity was assayed as described above.

#### 2.8. Enzyme kinetics

Initial rates were determined at  $60^{\circ}$  C by varying the concentration of E4P (10-3000  $\mu$ M) while maintaining the concentration of PEP at 3 mM. For the second substrate, the concentration of PEP was varied over a range from 50 to 6000  $\mu$ M while the concentration of E4P was kept constant at 3 mM. The amount of DAH7P produced was quantified by the discontinuous colorimetric assay (11). The kinetic constants were determined by fitting the Michaelis-Menten equation using the KaleidaGraph program.

#### 2.9. Feedback inhibition

To identify the allosteric effectors, enzyme, 3 mM PEP, 100 mM Tris-acetate (pH 7.0 at 70°C), and 1 mM inhibitor were incubated on ice for 30 minutes. The mixture was warmed to 70°C for 2 minutes before initiating the reaction with 3 mM E4P. A control reaction was performed because tryptophan interferes with the Aminoff assay. The control reaction was identical to the standard effector-inhibition reaction except 1 mM tryptophan was added after TCA quenching but immediately before the Aminoff assay as opposed to pre-incubating the enzyme with the tryptophan.

#### 2.10. Crystallization, data collection and refinement

All solutions used for crystallization, except for the solutions of protein and MnCl<sub>2</sub>, were treated with Chelex-100 to remove metal cations. DAH7PS<sup>Ap</sup>-Mn-PEP crystals were grown at room temperature in 5 L hanging drops using the vapor diffusion method. The drops contained 8 mg/ml DAH7PS<sup>Ap</sup>, 4% (w/v) polyethylene glycol (PEG) 8000, 6.3 mM PEP, 0.8 mM MnCl<sub>2</sub>, and 100 mM LiCl in 50 mM BTP (pH 8.5). The reservoir solution contained 30% PEG 8000 in 50 mM BTP (pH 8.5). The resulting crystals were in space group P2<sub>1</sub>.

X-ray diffraction measurements were performed at the Southeast Regional Collaborative Access Team (SERCAT) beam line 22-BM at the Advanced Proton Source from a single frozen crystal in an evaporating nitrogen stream. The crystal belonged to space group  $P2_1$  with cell dimensions a = 54.8 Å, b = 130.4 Å, c = 78.4 Å,  $\beta = 110.6^{\circ}$ , and diffracted to 2.3 Å. The diffraction data were processed with HKL2000 (12) and programs of the CCP4 suite, 1994, (Table 1). Four subunits of DAH7PS<sup>Ap</sup> present in the asymmetric unit were found by molecular replacement search with MOLREP (CCP4 suite, 1994), using the  $(\beta/\alpha)_8$ -barrel core of DAH7PS<sup>Tm</sup> (PDB ID 1RZM) as a model. CNS (13) and REFMAC5 (CCP4 suite, 1994) were used for refinement and O (14) was employed for model building and adjustment. The search for bound water molecules was performed with ARP/wARP (15).

The final model of DAH7PS<sup>Ap</sup> has  $R/R_{free}$  values of 17.3/21.8% and includes 8187 protein atoms, four  $Mn^{2+}$  cations and PEP molecules, and 191 bound water molecules. The Nterminal segments (from five to nine residues) are disordered in all subunits. The model has good stereochemistry as analyzed by PROCHECK (16) with 93.0% of the residues in the most favored regions and no residues in the disallowed regions of the Ramachandran plot. The coordinates and structure factors have been deposited in the Protein Data Bank (PDB ID 1VS1). LSQMAN was used to superimpose molecules (17). Buried interface surface areas were calculated utilizing Naccess (18).

# 3. RESULTS

#### 3.1. Overexpression and purification

The gene encoding DAH7PS<sup>Ap</sup> (gi:14600379) was cloned by PCR from genomic DNA and inserted into *E. coli* expression plasmid pT7-7. Protein expression in *E. coli* BL21(DE3) was too low to be useful. We attributed this result to the rare (for *E. coli*) codons that occur at the beginning of the insert's sequence. Protein production increased when we switched to *E. coli* BL21-CodonPlus®(DE3)-RIL cells. The purification began by heating the cell lysate to denature and precipitate the *E. coli* proteins. Subsequent chromatography using a Phenyl Superose column and then an UNO® Q6 column, produced homogeneous DAH7PS<sup>Ap</sup> (Fig. 1). The typical yield of purified was approximately 20 mg per liter of culture. The enzyme "as isolated" precipitates when it is concentrated to more than 2 mg/ml, but with PEP present, the enzyme remains soluble in solution to a concentration of 10 mg/ml.

# 3.2. Quaternary structure in solution

The DAH7PS<sup>Ap</sup> monomer migrated as a 30 kDa protein on an SDS-PAGE gel, consistent with the value (29,159 Da) predicted from its sequence. We determined the native molecular weight to be 55.4 kDa, or  $1.85 \times$  the monomeric weight, using analytical gel filtration chromatography. These data indicate that DAH7PS<sup>Ap</sup> exists in solution as a homodimer.

#### 3.3. Metal content

High-resolution inductively coupled plasma mass spectrometry showed that DAH7PS<sup>Ap</sup>, as isolated, contains 0.8 molar equivalents of Zn<sup>2+</sup> per subunit and trace amounts of other metals (Table 2). To demonstrate that DAH7PS<sup>Ap</sup> requires a metal ion for activity, we attempted to remove the metal ion with chelating agents and demonstrate that the apoenzyme is inactive. Treatment of DAH7PS<sup>Ap</sup> with 10 mM EDTA and 1 mM DPA for 1 hour reduces the enzymatic activity to 21% of its original activity, but does not fully inactivate the enzyme. Attempts to fully inactivate the enzyme by treating it with higher concentrations of EDTA, or EDTA with DPA, for one hour also failed to remove all the activity (data not shown). Incubation in a buffer containing 10 mM EDTA and 1 mM DPA for 24 hours resulted in enzyme that is resistant to treatment with standard chelating agents, and that the inactive apoenzyme is unstable and precipitates.

#### 3.4 DAH7PS activity

Purified DAH7PS<sup>Ap</sup> exhibits maximal activity at pH 5.7 (Fig. 2) when assayed at 60°C. However, since the enzyme precipitates in an acidic buffer over a period of several hours, we performed subsequent work at pH 7.0, a pH at which the enzyme retains approximately 40% of its maximal activity at 60°C. Enzymatic activity at pH 7.0 increases sharply between 65°C and 95°C. The optimum reaction temperature, 95°C (Fig. 3), corresponds to the growth temperature of *A. pernix*, which is 90°C to 95°C (19). Incubation of the isolated enzyme at 90°C for 1 hour reduces the activity of DAH7PS<sup>Ap</sup> is reduced by 50% (Fig. 4). Addition of the substrate PEP stabilizes the enzyme significantly, increasing its half-life at 90°C to 5.8 hours. Titration of each substrate demonstrated that the activity of DAH7P<sup>Ap</sup> follows Michaelis-Menten kinetics. Apparent kinetic constants were determined for each substrate, in the presence of a saturating concentration of the other substrate, at pH 7.0 and 60°C. The apparent K<sub>m</sub> for E4P was 280 ± 20 µM and the apparent K<sub>m</sub> for PEP was 890 ± 30 µM. The apparent k<sub>cat</sub> value is 1.0 ± 0.2 s<sup>-1</sup>.

#### 3.5. Feedback inhibition

We tested several downstream products of the shikimante pathway and found that none of them inhibited the activity of DAH7PS<sup>Ap</sup> at physiologically relevant concentrations (Table 3). Tryptophan initially appeared to inhibit the enzymatic activity. However, tryptophan added after the enzymatic reaction but before the Aminoff assay also showed a reduction in pink chromophore, indicating that tryptophan interferes with the Aminoff assay. Since the same apparent inhibition occurred whether the tryptophan was added before or after the enzymatic reaction took place, we conclude that tryptophan does not inhibit the enzymatic process.

#### 3.6. Crystal structure of DAH7PS<sup>Ap</sup>

Crystals of DAH7PS<sup>Ap</sup> grown in the presence of Mn<sup>2+</sup> and PEP contained a tetramer in the asymmetric unit. The four subunits have almost identical conformations. Superposition of 267 C $\alpha$  atoms of monomers A and B resulted in a root mean squared deviation (rmsd) of 0.08 Å. Superposition of 268 C $\alpha$  atoms of monomers A and C, or A and D led to rmsds of 0.13 Å and 0.14 Å, respectively. Between five and nine N-terminal residues were disordered in each subunit, including the tryptophan residues at positions two and four. The structure refined well, with an R / R<sub>free</sub> = 17.3 / 21.8%. All  $\phi$  and  $\psi$  dihedral angles lie in either the most favored region or the additional allowed regions of the Ramachandran plot. The modes of PEP and metal binding in DAH7PS<sup>Ap</sup> were very similar to those in *T. maritima* DAH7PS (20). The plane of the carboxylate of PEP is twisted ~20° with respect to the plane of its C-C double bond, as seen in the X-ray crystal structures of *T. maritima* DAH7PS (20) and the phenylalanine-sensitive DAH7PS from *E. coli* (21). A water molecule occupies the presumed binding site for the phosphate group of E4P in the DAH7PS<sup>Ap</sup> structure.

# 4. DISCUSSION

The DAH7PS enzymes are a diverse group of proteins that have incorporated a nuanced set of regulatory mechanisms. The smaller enzymes, classified as Type I (22) and further divided into Types Ia and IB (23) using sequence-based phylogenetic algorithms, are either unregulated or regulated by small sequences appended to the N- or C-terminus. DAH7PS<sup>Ap</sup> is categorized as a Type I $\beta$  enzyme, as are DAH7PS<sup>Tm</sup> and the previously described DAH7PS from *Pyrococcus furiosus* (DAH7PS<sup>Pf</sup>, (24)). DAH7PS<sup>Ap</sup>, like DAH7PS<sup>Pf</sup> and the catalytic domain of DAH7PS<sup>Tm</sup>, has a core  $(\alpha/\beta)_8$  barrel. The DAHPS<sup>Ap</sup> has an extension of several residues at the N-terminus relative to DAH7PS<sup>Pf</sup>, but these residues are disordered in the electron density of A. pernix and do not provide a binding site for a typical feedback inhibitor. DAH7PS<sup>Ap</sup> is a dimer in solution, but the asymmetric unit in the crystal is a tightly associated tetramer (Fig. 5A). DAH7PS<sup>Pf</sup> is also a dimer in solution (25) but packs as an apparent tetramer in the crystal (24). In contrast, DAH7PS<sup>Tm</sup> is a tetramer in solution (5), but packs in the crystal as a dimer (see PDB ID: 3PG8). The buried interface surface areas between the adjacent monomers were calculated for the DAHPS<sup>Ap</sup>, DAHPS<sup>Pf</sup>, and DAHPS<sup>Tm</sup> tetramers (Table 4). The buried interface area between monomers A and D is about 122 Å<sup>2</sup> more than the buried interface area between monomers A and C; however, this difference is not significant enough to determine which pair of dimers is the functional dimer for DAHPS<sup>Ap</sup>. A closer examination of the DAHPS<sup>Ap</sup> crystal structure shows highly symmetrical residue interactions (G155, E160, R184, and F186) between monomers A and C and similar residue interactions between monomers B and D (Fig. 5B). The E160 from monomer B is in close proximity to R184 of monomer D and forms hydrogen bonds, thus, monomer A and C are more likely to be the biological dimer for DAHPS<sup>Ap</sup> than monomers A and D.

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A picture of the active site of DAH7PS<sup>Ap</sup> (Fig. 6) shows the key features involved in catalysis. The four resides that donate metal chelating ligands, H216, D253, E242, and C46, are strictly conserved within DAH7PS<sup>Ap</sup>, DAH7PS<sup>Tm</sup>, and DAH7PS<sup>Pf</sup>, and broadly conserved within the Type I DAH7PS enzymes. All of the characterized DAH7PS enzymes are metalloenzymes, and it is quite likely that every DAH7PS enzyme is a metalloenzyme. The identity of the physiologically relevant metal ion for these enzymes has not been definitively proved. Early studies (26) suggested that the phenylalanineinhibitable DAH7PS from yeast (ARO3) was an iron-containing enzyme. Subsequent experiments (27) demonstrated that apo-enzyme reconstituted with divalent zinc, copper, or iron had similar activity. Divalent cobalt ions gave elevated activity (167%), divalent cadmium or nickel ions returned modest activity (<70%), while divalent magnesium or manganese ions delivered significantly reduced activity (<20%). Structural studies of yeast DAH7PS have been done with divalent cobalt ions present and with divalent manganese ions present (28). The structures of Co<sup>2+</sup>- and Mn<sup>2+</sup>-containing enzymes, with bound PEP, are essentially the same. Both exhibit trigonal bipyramidal coordination of the metal ions, with a water molecule in an equatorial position. The metal ion in DAH7PS<sup>Ap</sup> (Fig. 6) also has a water molecule close enough to coordinate the bound Mn<sup>2+</sup>. Basic residues R181 and R130 form hydrogen bonds to the phosphate group of PEP while K75 and R70 form hydrogen bonds to the carboxyl group of PEP. This type of architecture is typical for Type IB DAH7P synthases. Fig.7 presents an overlay of the active sites of DAH7PS<sup>Ap</sup> and DAH7PS<sup>Tm</sup> that demonstrates the near identity of the two active sites.

The biochemical data presented here are also similar to those of other Type I DAH7P synthases. The temperature-activity profile noted for DAH7PS<sup>Ap</sup> is quite similar to that reported for DAH7PS<sup>Pf</sup>, which also has very low activity below 30°C, and escalating activity above  $65^{\circ}C$  (25). The thermal stabilization of DAH7PS by PEP is not surprising, since thermal stabilization of an enzyme by its substrate has been previously observed with PEP carboxykinase and PEP (29, 30), with aminotransferase and its substrates pyridoxalmonophosphate and 2-ketoglutarate (31), D-glyceraldehyde-3-phosphate dehydrogenase and coenzyme NADP+ (32), and endo-1,4-beta-xylanase and substrate xylan (33). An increased in DAHPS stability provided by the binding of PEP to the active site has also been previously reported for L-Phe sensitive *E. coli* DAH7PS (34). A more complete examination of the effect of ligands on the thermostability of an enzyme is contained in a paper by Edwards et al. (35) that describes experiments with beta-galactosidase.

The DAH7PS enzymes from *P. furiosus* and *A. pernix*, which are members of euryarchaea and crenarchea, respectively, have no allosteric modifier. Neither do the DAH7PS enzymes from several other members of the archaea, based upon the similarity of their primary sequences to those of DAH7PS<sup>Ap</sup> and DAH7PS<sup>Pf</sup>. It seems wasteful for cells to synthesize aromatic compounds and the other metabolites dependent upon the shikimate pathway in an unregulated manner. Perhaps these primitive organisms regulate this pathway at a different step in the pathway, such as the supply of E4P, or perhaps the regulation occurs at the transcriptional level. More research is necessary to resolve this conundrum. Less primitive organisms apparently evolved more complex DAH7PS enzymes that are susceptible to feedback control. This has been accomplished through additions to the primitive ( $\alpha/\beta$ )<sub>8</sub> barrel scaffold, like those seen in the structures of DAH7PS<sup>Tm</sup> and *E. coli* DAH7PS. There is evidence to support the idea that gene duplication and single point mutations within the extensions are responsible for creating DAH7PS with different inhibitor binding abilities (36).

# Acknowledgments

This work was funded by National Institutes of Health grants GM 053069 and AI061531 (to R.W.W.) and National Science Foundation grant MCB-9723633 (to R.H.K.). We thank Dr. Ted Huston (Department of Geology, University of Michigan) for performing the metal analysis studies, and Dr. Tod P. Holler of the Department of Medicinal Chemistry, University of Michigan, for assistance with writing, editing and preparing the manuscript for publication.

# Abbreviations

ВТР	1,3-bis[tris(hydroxymethyl)methylamino]propane		
DAH7PS	3-deoxy-D-arabino-heptulosonate 7-phosphate synthase		
DAH7PS <sup>Ap</sup>	the DAH7PS from Aeropyrum pernix		
DAH7PS <sup>Pf</sup>	the DAH7PS from Pyrococcus furiosus		
DAH7PS <sup>Tm</sup>	the DAH7PS from Thermatoga maritima		
DPA	dipicolinic acid		
E4P	D-erythrose 4-phosphate		
EDTA	ethylenediamminetetraacetic acid		
KDO	3-deoxy-D-manno-octulosonate		
PCR	polymerase chain reaction		
PEP	phosphoenolpyruvate		
SDS-PAGE	sodium dodecylsulfate polyacrylamide gel electrophoresis		
rmsd	root mean squared deviation		
TCA	trichloroacetic acid		
Tris	tris(hydroxyethyl)aminomethane		

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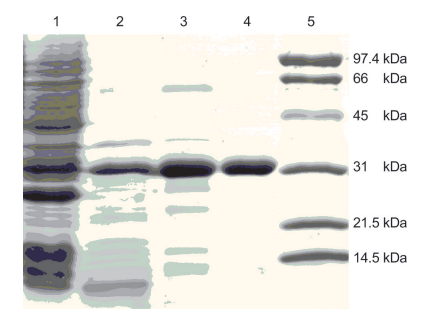
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# Highlights

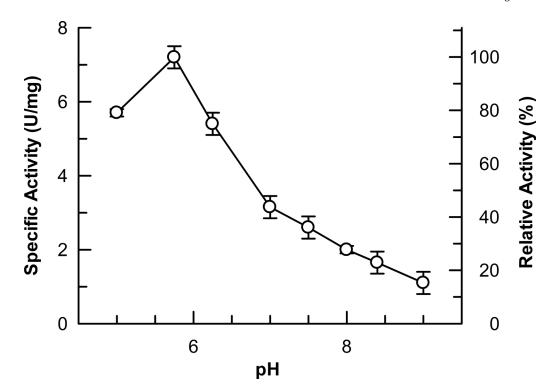
> We investigate the DAH7P synthase from *A. pernyx*, a primitive archaebacterium. > We present the biochemical characterization and crystal structure of the enzyme. > The enzyme activity is not regulated by metabolites from the shikimate pathway.

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# Figure 1.

SDS-PAGE of DAHPS<sup>Ap</sup>. Electrophoresis was performed under reducing conditions on a 12% polyacrylamide gel and stained with Coomassie Brilliant Blue R. Lane 1, supernatant after cell disruption; lane 2, heat supernatant; lane 3, phenyl superose column; lane 4, UNO<sup>TM</sup> Q6 fraction; lane 5, molecular weight standards. DAHPS has an apparent molecular weight of 30 kDa.



#### Figure 2.

Optimum pH. The optimal pH of the DAHPs were determined by measuring enzyme activities in 3 mM PEP, 3 mM E4P, and 100 mM BTP buffer with an adjusted pH of 5 - 9 at 60°C; (•- DAHPS<sup>Ap</sup>).

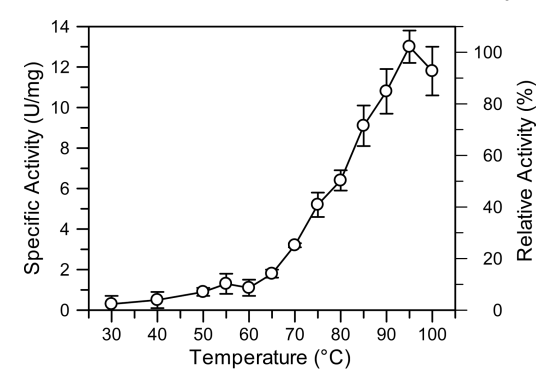
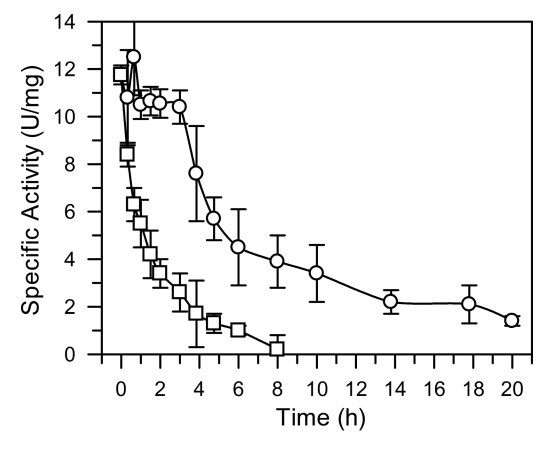


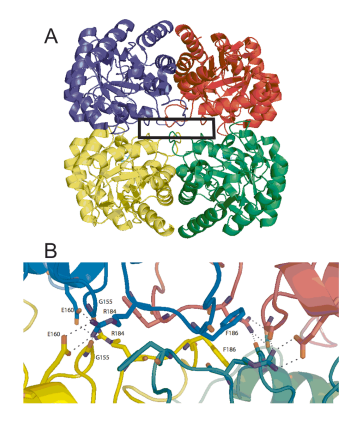
Figure 3.

Optimum reaction temperature. The enzyme activities were measured in 3 mM PEP, 3 mM E4P, and 100 mM Tris-acetate (pH 7.0 at indicated temperatures) under various reaction temperatures (•- DAHPS<sup>Ap</sup>). Error bars illustrate standard deviation of three sets of data.



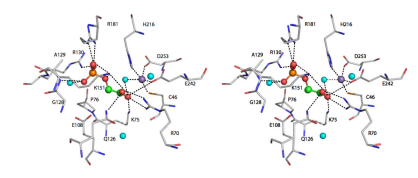
# Figure 4.

Thermostability. The DAHPS<sup>Ap</sup> in 20 mM Tris-acetate (pH 8.0 adjusted at room temperature) was incubated at 90°C in the presence ( $\Box$ ) or absence ( $\circ$ ) of 1 mM PEP. At various times, an aliquot was taken out to determine its residual activity at 90°C. Standard deviation of three sets of data is shown by the error bars.



#### Figure 5.

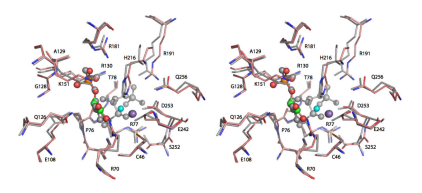
DAHPS<sup>Ap</sup> tetramer. (A). DAHPS<sup>Ap</sup> tetramer structure (monomer A-green, monomer Bcyan, monomer C-magenta, and monomer D-yellow). (B). Magnified interface shown G155, E160, R184, and F186 from each monomer interacting at the monomer interface. Interacting residues are labeled in monomers B and D, and similar interactions occur between monomer A and C. Zhou et al.



#### Figure 6.

Active site of DAHPS<sup>Ap</sup>. A stereoview showing the active site residues interacting with the manganese ion (purple sphere), the PEP substrate (sphere model: orange-phosphorus, red-oxygen, and green-carbon), and water molecules (cyan spheres). Four residues and one water molecule are shown to coordinate to manganese (C46-Mn 2.58Å, H216-Mn 2.44Å, E242-Mn 2.13Å, D253-Mn 2.39Å, H<sub>2</sub>O-Mn 2.26Å).

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#### Figure 7.

Superposition of the active sites of DAH7PS<sup>Ap</sup> and DAH7PS<sup>Tm</sup>. Stereo view of the DAHPS<sup>Ap</sup> active site residues interacting with bound the manganese ion (purple sphere) and the PEP substrate (sphere model: orange-phosphorus, red-oxygen, and green-carbon). The water molecules are shown as cyan spheres. DAH7PS<sup>Tm</sup> residues with its bound PEP, E4P, and cadmium ion are shown in gray. All active site residues shown here are conserved between the two enzymes.

Data collection and refinement statistics for  $DAH7PS^{Ap}$ .

A. Diffraction Data				
Resolution (Å)	20.0-2.3			
Unique reflections	45666			
Completeness (%)	96.5 (89.1)			
Multiplicity	7.5 (7.2)			
<i>/&lt;\sigma(I)&gt;</i>	19.2 (6.2)			
R <sub>merge</sub> (%)	9.3 (30.5)			
Outermost shell (Å)	2.38-2.30			
B. Refinement				
rmsd bond length (Å)	0.019			
rmsd bond angles (deg.)	1.67			
Average B-factor, protein (Å <sup>2</sup> )	36.1			
$R_{work}/R_{free}$ (%)	17.3/21.8			
Test set (%)	5.1			

# Metal analysis.

Metal	Molar equivalent metal per DAH7PS <sup>Ap</sup> subunit
Zinc	0.8
Magnesium	0.05
Iron	0.03
Cobalt	$ND^{a}$
Nickel	ND
Copper	ND
Manganese	ND
Cadmium	ND

<sup>a</sup>None detected (value < 0.01)

# Feedback regulation.

Compounds	DAH7PS <sup>Ap</sup> activity remaining after treatment $(\%)^a$
None	100
L-Phenylalanine	$93.9\pm6.8$
L-Tyrosine	$88.9\pm 6.8$
Chorismate	$89.0\pm6.7$
Shikimate	$99.9\pm5.0$
Prephenate	$84.0 \pm 2.5$
L-Tryptophan	$49.4\pm3.9$
L-Tryptophan <sup>b</sup>	41.8 ± 8.6

<sup>a</sup>Results show the standard deviation of triplicate assays.

 $^b$  added after quenching with 10% TCA, but before the periodate-Aminoff thiobarbiturate assay.

Interface contact areas for monomers within DAH7PS crystals.

	DAHPS <sup>Ap</sup>	DAHPS <sup>Pf</sup>	DAHPS <sup>Tm</sup>
	B C D A	A' B' B A (	B' B A' A
	Functional 4° structure	Interface contact area between	Interface contact area between
DAHPS <sup>Ap</sup>	Dimer	C and A = 1393 $Å^2$	D and A = 1515 $Å^2$
DAHPS <sup>Pf</sup>	Dimer	B' and A = 1383 $\text{\AA}^2$	B and A = 1266 $Å^2$
DAHPS <sup>Tm</sup>	Tetramer	B and A = 1469 $\text{\AA}^2$	A' and A = 1426 $\text{\AA}^2$