

β -Hydroxylation of the Aspartyl Residue in the Phytotoxin Syringomycin E: Characterization of Two Candidate Hydroxylases AspH and SyrP in *Pseudomonas syringae*[†]

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ABSTRACT: The pseudomonal phytotoxin syringomycin E and related nonribosomal peptides contain an L-threo- β -hydroxyaspartyl residue at the eighth position of the lipodepsipeptide backbone as part of a conserved nonproteinogenic tripeptide motif. Informatic analysis of the *P. syringae* genome suggests only one putative non-heme iron hydroxylase, AspH. On heterologous expression in *Escherichia coli* AspH shows robust catalytic activity with free L-Asp and L-Asp thioesters to make β -OH-Asp but yields the erythro diastereomer rather than the threo configuration that is found in syringomycin. Further analysis of the Syr gene cluster indicated that SyrP, previously annotated as the gene regulatory protein for the five-gene Syr cluster, is actually homologous to the known non-heme mononuclear iron hydroxylase TauD. Indeed, purified SyrP acts on Asp tethered as the protein-bound S-pantetheinyl thioester on the eighth module of the SyrE megasynthetase. The hydroxylation gives the anticipated L-threo-3-OH-Asp diastereomer found in syringomycin. The knockout of *syrP* abolishes the production of the mature syringomycin E, while knockout of *aspH* has no effect on syringomycin production.

The pseudomonad genus of Gram-negative bacteria includes prolific producers of nonribosomal peptide (NRP) natural products. Phytopathogenic *Pseudomonas syringae* strains produce a number of such plant toxins, including coronatine, syringomycins, phaseolotoxin, tabtoxin, and tagetitoxin (1–5). Syringomycins are members of a family of related cyclic lipodepsipeptides that act as plant toxins by aggregating to form pores in plant cell membranes to facilitate exit of nutrients. One hallmark of nonribosomal peptides is the content of nonproteinogenic amino acid residues. In the five known pseudomonal lipopeptides, syringomycin E, cormycin A, pseudomycin B, syringostatin A, and syringotoxin B, there is a conserved C-terminal tripeptide moiety in the scaffold, composed of dehydrobutyrine₇, β -OH-Asp₈, and 4-Cl-Thr₉ (Figure 1A) (3, 6–9). This nonproteinogenic tripeptide motif, comprising residues 7–9 of syringomycin, may be consequential for the biological function of these toxins.

The syringomycin gene cluster contains six ORFs, *syrD*, *syrP*, *syrB1*, *syrB2*, *syrC*, and *syrE*. The sequence of this biosynthetic gene cluster (10) for liponapeptidolactone syringomycin (Figure 1B) suggested that residues 1–8 are assembled on a multimodular megasynthetase protein SyrE. Residue 9 is the 4-Cl-threonine, inserted at the end of the assembly line by the tandem action of three enzymes, SyrB1, SyrB2, and SyrC (11–13). Threonine is activated by the

adenylation (A)¹ domain of SyrB1, installed as the S-pantetheinyl thioester on the adjacent T (thiolation) domain (11), and then chlorinated by the non-heme mononuclear iron-containing halogenase SyrB2 (11, 12). At that juncture SyrC acts as an acyltransferase for the 4-Cl-Thr moiety, moving it to the T₉ domain of the SyrE assembly line (13). SyrD is annotated as an ABC-type export protein and SyrP as a regulatory phosphoryl transferase that is an analogue to a sensor kinase (14, 15).

Having worked out the tailoring and insertion of threonine as 4-Cl-Thr₉ (11, 13), we have now turned our attention to genesis of the L-threo- β -OH-Asp₈ residue in syringomycin and the related pseudomonal lipodepsipeptides. Many of the proteinogenic amino acid monomers, including Asp, Asn, Glu, Pro, and Lys, undergo enzymatic β -hydroxylation, both in the posttranslational modification of ribosomal proteins (16, 17) and in the biosynthesis of nonribosomal peptides (18–22). The known hydroxylases use iron as the redox catalyst in the active site, either Fe³⁺ in a cytochrome P450 heme environment (19) or Fe²⁺ in a non-heme mononuclear state at the active site (22).

Informatic analysis of the *P. syringae* pv *syringae* B728A genome (23) turns up one ORF at the primary locus

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¹ Abbreviations: A, adenylation domain; α -KG, α -ketoglutarate; Ala, alanine; Arg, arginine; Asn, asparagine; Asp, aspartate; C, condensation domain; CoA, coenzyme A thioester; Cys, cysteine; Da, dalton; Dap, 2,3-diaminopropionic acid; DMSO, dimethyl sulfoxide; DMF, dimethylformamide; DTT, dithiothreitol; EDTA, ethylenediaminetetraacetic acid; Glu, glutamate; HPLC, high-pressure liquid chromatography; IPTG, isopropylthiogalactopyranoside; LC-MS, liquid chromatography–mass spectrometry; MS-MS, tandem mass spectrometry; NMR, nuclear magnetic resonance; Ser, serine; SNAC, N-acetylcysteamine; T, thiolation domain; TCA, trichloroacetic acid; TFA, trifluoroacetic acid; TFE, trifluoroethanol; TE, thioesterase domain.

Psyr_1584, *aspH*, predictively annotated as a non-heme, iron-dependent aspartyl hydroxylase. To evaluate the catalytic function of AspH, we have expressed AspH in *Escherichia coli* and report on its activity here. Our particular interest was (1) in the capacity of AspH to hydroxylate free Asp vs Asp in a thioester linkage and (2) determination of the chirality of β -hydroxylation. Our unanticipated finding, that AspH generates *L*-erythro- rather than *L*-threo-3-OH-Asp, has led us in this work to reexamine the remaining ORF in the syringomycin cluster, *syrP* (Figure 1B). Although SyrP has been annotated as a regulatory protein for the pathway (15), we report that it is actually an aspartyl hydroxylase that generates only the *threo*-3-OH-Asp product and that it is involved in syringomycin maturation.

EXPERIMENTAL PROCEDURES

Materials and General Methods. All radiolabeled chemicals were obtained from American Radiolabeled Chemicals Inc. (ARC). *L*-threo-3-OH-Asp was obtained from Tocris Pharmaceuticals, and *L*-erythro-3-OH-Asp was obtained from Wako Pure Chemicals, Ltd. All other chemicals used were from Sigma, unless otherwise specified. The TOP10 and BL21(DE3) competent *E. coli* strains were purchased from Stratagene. The *Pfu*Turbo DNA polymerase used in PCR was purchased from Stratagene. During protein purification, cells were lysed using an Avestin EmuLsiflex-C5 cell disruptor. SDS-polyacrylamide gels were obtained from Bio-Rad, and autoradiography was performed on a Typhoon 9400 scanner (GE Healthcare). HPLC was carried out on a Beckman Coulter System Gold using a Vydac C18 column or a Phenomenex Chirex column. Radio-HPLC was performed on an identical system equipped with a β -Ram radioisotope detector (INUS). LC-MS analysis was performed on an LCMS-QP8000a spectrometer (Shimadzu) with a Vydac C18 LC-MS column.

Cloning, Overproduction, and Purification of SyrE-A₈T₈, AspH, and SyrP. The SyrE-A₈T₈ construct was obtained by PCR amplification from a pET-28a plasmid containing the SyrE-A₈T₈C₉T₉TE fragment (13). The following oligonucleotide pairs were used in PCR amplification: 5' CGG GGA TCC CAT ATG CTT GAG CAG GAT CCG 3' and 5' CGG ATA GAG CTC CTA ACG GCC AGA GCG ATT 3'. The *aspH* and *syrP* genes were obtained from GeneArt in synthetic form with codons optimized for expression in *E. coli*. All constructs were digested with *Nde*I and *Xho*I and ligated into similarly digested pET28a vectors to generate N-terminally His₆-tagged constructs.

The pET-28a expression vectors containing the constructs described above were transformed into BL21(DE3) competent cells. Cultures were grown in Luria-Bertani medium supplemented with 30 μ g/mL kanamycin at 37 °C until the OD₆₀₀ reached ~0.3, at which time the cultures were cooled to 25 °C, and grown until the OD₆₀₀ reached ~0.6. The cultures were induced with 0.1 mM IPTG and grown at 15 °C overnight. Cells were harvested by centrifugation at 6000 rpm for 30 min, flash frozen in liquid N₂, and stored at -80 °C until further purification.

Cells were thawed, resuspended in buffer A (300 mM NaCl, 5 mM imidazole, 20 mM Hepes, pH 8.0), and lysed by cell disruption. Cell debris was removed from the lysate by centrifugation at 15000 rpm for 30 min, and the

supernatant was removed and bound to Ni-NTA resin by rocking at 4 °C for 2 h. The resin was added to a Bio-Rad Econo-Sphere column and washed with buffer A. Protein was eluted with buffer B (300 mM NaCl, 30 mM imidazole, 20 mM Hepes, pH 8.0) and buffer C (100 mM NaCl, 200 mM imidazole, 20 mM Hepes, pH 8.0). Protein-containing fractions were identified by SDS-PAGE, combined, and dialyzed overnight in 100 mM NaCl, 1 mM EDTA, and 50 mM Hepes, pH 8.0, with 10% glycerol. The dialyzed protein was concentrated, flash frozen in liquid N₂, and stored at -80 °C.

ATP-PP_i Exchange Assay for SyrE-A₈. A typical reaction contained 10 mM amino acid, 10 mM MgCl₂, 1 mM ATP, 1 mM DTT, 2 μ M SyrE_{8,9}, and 5 mM sodium [³²P]pyrophosphate in a total volume of 500 μ L with 50 mM Hepes, pH 7.5. Reactions were carried out with *L*-Asp, *L*-erythro-3-OH-Asp, and *L*-threo-3-OH-Asp. In tandem, reactions containing no enzyme were carried out as negative controls. Reactions were initiated by the addition of enzyme, and aliquots were quenched at 0, 1, 2, 5, 10, 20, 30, and 60 min by the addition of 750 μ L of a solution containing 1.6% (w/v) activated charcoal, 200 mM sodium pyrophosphate, and 3.5% (v/v) perchloric acid. For each time point, the charcoal was pelleted by centrifugation and washed twice with 750 μ L of a solution containing 200 mM sodium pyrophosphate with 3.5% perchloric acid (wash buffer). The charcoal pellet was then resuspended in 750 μ L of wash buffer and mixed with liquid scintillation fluid. Radioactivity bound to the charcoal was measured by liquid scintillation counting.

Phosphopantetheinylation of SyrE-A₈T₈. In order to generate the active holo form of the T₈ thiolation domain in the SyrE-A₈T₈ construct, a phosphopantetheinylation reaction must be carried out. In a typical reaction, 1 nmol of enzyme is incubated with MgCl₂ (0.5 μ mol), CoA (50 nmol), and Sfp (10 nmol) in 50 mM Hepes buffer, pH 7.5, in a total reaction volume of 25 μ L for 30 min at room temperature to produce the holoenzyme.

Autoradiographic Assay for Autoaminoacylation of SyrE-A₈T₈. Holo-SyrE-A₈T₈ (0.6 nmol) was incubated with *L*-[¹⁴C]Asp (20 nmol) and ATP (100 nmol) in 50 mM Hepes buffer, pH 7.5, in a total reaction volume of 59 μ L at room temperature. The reaction was initiated by the addition of ATP. At 1, 10, and 30 min, 20 μ L aliquots of the reaction were quenched in 2 \times SDS-PAGE loading buffer. The quenched aliquots were heated to 90 °C for 10 min and then run on a 12% SDS-polyacrylamide gel. The gel was stained, destained, dried, and exposed to a phosphorimager screen for 3 days. The screen was then scanned using a Typhoon imager.

Assay for Aminoacylation of SyrE-A₈T₈ by TCA Precipitation. Holo-SyrE-A₈T₈ (17 nmol) was incubated with *L*-[¹⁴C]Asp (170 nmol) and ATP (200 nmol) in 50 mM Hepes buffer, pH 7.5, in a total reaction volume of 650 μ L at room temperature. The reaction was initiated by the addition of ATP. At various time points, 50 μ L aliquots of the reaction were quenched in 500 μ L of 10% trichloroacetic acid (TCA). To each quenched aliquot, 100 μ L of bovine serum albumin (BSA) was added, and the aliquots were incubated on ice for 5 min. To pellet the precipitated proteins, the reactions were centrifuged at 13000 rpm for 5 min, and an additional 500 μ L of 10% TCA was added to each. Centrifugation was repeated, and the supernatant was aspirated. The remaining

protein pellets were redissolved in 100 μ L of formic acid, and all samples were subjected to liquid scintillation counting.

General Assay for Hydroxylase Activity with Various Substrates. In a typical 100 μ L reaction, the recombinant hydroxylase (either AspH or SyrP) was incubated with substrate (250 μ M), $(\text{NH}_4)_2\text{Fe}(\text{SO}_4)$ (0.5 mM), α -ketoglutarate (0.5 mM), ascorbate (0.5 mM), dithiothreitol (0.5 mM), and catalase (0.5 ng/ μ L) in 50 mM Hepes, pH 7.5. Reactions were incubated at room temperature for 30 min.

Hydroxylase Assay with L-[^{14}C]Asp. Recombinant hydroxylase (either AspH or SyrP) was incubated with L-[^{14}C]Asp under the conditions described above. The reactions were quenched in 10% TCA and centrifuged at 13000 rpm for 20 min, and the supernatant was analyzed by chiral radio-HPLC on a Phenomenex 250 \times 4.6 mm 3126 Chirex column with an isocratic gradient of 95% 2 mM copper sulfate in 5% 2-propanol, monitoring ^{14}C radioactive counts. Reactions with cold L-Asp were performed in tandem for LC-MS analysis.

Hydroxylase Assay with SyrE-A $_8$ T $_8$ -S-L-[^{14}C]Asp. The SyrE-A $_8$ T $_8$ -S-L-[^{14}C]Asp substrate was generated by phosphantethinylation followed by autoaminoacylation of SyrE-A $_8$ T $_8$ with L-[^{14}C]Asp, as described above. The reaction mixture containing SyrE-A $_8$ T $_8$ -S-L-[^{14}C]Asp was washed twice with 300 μ L of 50 mM Hepes in a Biomax 10 kDa MWCO centrifugation filter to remove unbound L-[^{14}C]Asp. The SyrE-A $_8$ T $_8$ -S-L-[^{14}C]Asp substrate was then incubated with either of the recombinant hydroxylases as described, and reactions were quenched with 10% TCA and centrifuged at 13000 rpm for 20 min. The supernatant was aspirated, and the protein pellet was redissolved and thioester linkage hydrolyzed in 200 μ L of 100 mM LiOH with incubation at 80 $^\circ\text{C}$ for 20 min. The protein was then reprecipitated by addition of 40 μ L of 50% TCA followed by centrifugation at 13000 rpm for 20 min. The supernatant was analyzed by chiral radio-HPLC as described above.

Hydroxylase Assay with L-Aspartyl-S-N-acetylcysteamine. The aspartyl-S-N-acetylcysteamine (L-Asp-SNAC) substrate was synthesized as described previously (24). Recombinant hydroxylase was incubated with the L-Asp-SNAC, reactions were quenched in 10% TCA, and the thioester bond was hydrolyzed in 100 mM LiOH, as with the SyrE-A $_8$ T $_8$ -S-L-[^{14}C]Asp substrate. Reactions were analyzed by chiral radio-HPLC as described.

Hydroxylase Assay with N-Acyl-nonapeptide (COOH-L-Thr-L-Asp-L-Abu-L-Phe-L-Arg-L-Dab-D-Dab-D-Ser-L-Ser-N-Ac) and N-Acyl-nonapeptidyl-CoA (CoA-S-L-Thr-L-Asp-L-Abu-L-Phe-L-Arg-L-Dab-D-Dab-D-Ser-L-Ser-N-Ac). The nonapeptide and nonapeptidyl-CoA were synthesized as described previously (13). Recombinant hydroxylase was incubated with either substrate, and reactions were quenched in 10% TCA, centrifuged to remove protein, and analyzed by LC-MS and HPLC. The HPLC analysis was performed on a Vydac 250 \times 4.6 mm C18 small pore column with a 0–100% acetonitrile gradient over 40 min beginning in 0.1% TFA in H $_2$ O and monitoring absorbance at 220 nm.

Construction of the aspH Gene Disruption Cassette. The *aspH* gene was PCR-amplified from genomic DNA from *P. syringae* pv *syringae* B301D (obtained as a gift from Dennis C. Gross, Texas A&M University, College Station, TX) using the following primers: 5'-CCGGATCCATGACCCTTTCA-

TTTGTTCCTCAAGGC-3' and 3'-CCAAGCTTTTAACCAAACAACAGTAGCCCAG-5' containing the *Bam*HI and *Hind*III sites, respectively. Amplification was carried out using Finnzyme Phusion DNA polymerase (New England Biolabs), following the manufacturer's instructions. The resulting fragment was digested with *Bam*HI/*Hind*III and ligated into the similarly digested pEX18Gm gene replacement vector (25) to produce the pEX18GmA1 plasmid. A 559 bp central fragment was excised from the *aspH* gene within pEX18GmA1 using *Nru*I and *Nsi*I, and the vector fragment was purified and its 5' and 3' termini were blunted using the NEB Quick blunting kit.

The *cat* gene was PCR-amplified from pKD3 (26) using the following primers: 5'-CCGAATTCATGGAGAAAAAATCACTGGATATACCAC-3' and 3'-CCGAATTCATCGCAGTACTGTTGTATTTCATTAAGC-5', both encoding for *Eco*RI sites. The resulting fragment was *Eco*RI digested and ligated into the similarly digested pPS854 vector between the FRT-encoding sites in the multiple-cloning site of the vector to generate the pPS854-FCF vector. The *cat* gene flanked by the two FRT sites was excised from pPS854-FCF with *Sac*I, and the resulting FRT-*cat*-FRT fragment was gel purified and its 5' and 3' termini were blunted as described above. The FRT-*cat*-FRT fragment was then ligated via blunt ends into the pEX18GmA1 linear fragment described above to generate the pEX18GmAKO gene replacement vector.

Construction of the syrP Gene Disruption Cassette. The *syrP* gene was PCR-amplified from genomic DNA from *P. syringae* pv *syringae* B301D using the following primers: 5'-CCGGATCCATGAGTCTTCAAGCTAACACTGCGCC-3' and 3'-CCAAGCTTTCAGGCCGGTTGCCAAACGTCGCC-5' containing the *Bam*HI and *Hind*III sites, respectively. Amplification was carried out as described above. The resulting fragment was digested with *Bam*HI/*Hind*III and ligated into the similarly digested pET28a vector to produce the pET28aP vector. A central 300 bp segment of the *syrP* gene in pET28aP was excised with *Bst*BI, and the linear vector fragment was gel-purified and its 5' and 3' termini were blunted as described. Blunt-end ligation was carried out between this fragment and the FRT-*cat*-FRT, generated as described in the previous section to generate the pET28aPFCFP vector. Using the *Bam*HI and *Hind*III sites in this plasmid, the 1.6 kb fragment containing *syrP* interrupted by the FRT-*cat*-FRT cassette (i.e., *syrP*-FRT-*cat*-FRT-*syrP*) was excised and ligated into the pEX18Gm gene transfer vector to produce the pEX18GmPKO gene replacement vector.

Introduction of Disrupted aspH and syrP Genes into P. syringae pv syringae. The pEX18GmAKO and pEX18GmPKO gene replacement vectors were transformed into the *E. coli* S17.1 λ (27) mating strain by electroporation (28). Conjugation was performed using the filter membrane mating technique (29), using the donor strain described above and a rifampicin-resistant variant of *P. syringae* pv *syringae* B301D as the recipient strain. Briefly, the donor strain carrying the desired delivery plasmid was grown overnight at 37 $^\circ\text{C}$ in 2 mL of LB medium containing 10 $\mu\text{g}/\text{mL}$ gentamycin and 25 $\mu\text{g}/\text{mL}$ chloramphenicol. The recipient strains were grown overnight in 2 mL of LB medium without selection. The donor cells were harvested and washed twice with 2 mL of 1% NaCl to remove the excess antibiotics.

Ten microliters of the donor cells and 100 μ L of the recipient cells were added to 5 mL of 1% NaCl, vortexed, and then filtered through a 25 mm type HA 0.45 μ m Millipore membrane (Bedford, MA). The filter was placed on LB solid medium and incubated at 30 °C for 12 h. The filter was then immersed in 5 mL of 1% NaCl and vortexed to resuspend the mating culture. Primary integrants were selected by plating 10–500 μ L aliquots of the mating resuspension on LB solid medium containing 25 μ g/mL rifampicin and 25 μ g/mL chloramphenicol. Successful intergration events were confirmed by testing for ampicillin sensitivity. The second homologous crossover was selected for by dual resistance to sucrose and chloramphenicol resulting from the loss of the *sacB* counterselectable marker. Total genomic DNA was isolated from the sucrose- and chloramphenicol-resistant colonies, and PCR was used to confirm the presence of the FRT-*cat*-FRT cassette in the *aspH* or *syrP* gene within the *P. syringae* chromosome (i.e., *aspH::cat* and *syrP::cat*).

Removal of the *cat* selection marker from the resulting *P. syringae* knockout clones was performed by transformation of the latter with the FLP recombinase-encoding plasmid pFLP2 (25). Transformants were selected for by ampicillin resistance and were grown in liquid culture. The FLP-catalyzed excision of the *cat* selection marker was confirmed by chloramphenicol sensitivity. Total genomic DNA was isolated from sensitive colonies, and PCR was used to confirm loss of the *cat* element and the presence of the truncated *syrP* and *aspH* genes.

Syringomycin Production, Extraction, and Analysis. The *syrP::cat*, Δ *syrP*, and wild-type strains of *P. syringae* pv *syringae* B301D were grown in 200 mL of PDB media in still culture at 25 °C for 6 days, and the Δ *aspH* strain was grown for 8 days due to delayed growth. Cultures were quenched with 1:1 volumes of ice-cold acetone, acidified to pH 2.5, and stored at 4 °C overnight. Cell debris was removed by centrifugation, and acetone was removed by rotary evaporation under vacuum. The water residue was then extracted three times in water-saturated 1-butanol, and the butanol phase was evaporated by rotary evaporation under vacuum. The remaining water residue was lyophilized and redissolved in 5 mL of ddH₂O, pH 2.0. The extracts were analyzed both by LC-MS and by C18 HPLC. The HPLC analysis was performed on a Vydac 250 \times 4.6 mm C18 small pore column with a 0–100% acetonitrile gradient over 50 min beginning in 0.1% TFA in H₂O and monitoring absorbance at 220 nm.

RESULTS

Overproduction and Purification of AspH and SyrE-A₈T₈. A search of the annotated genome of *P. syringae* pv *syringae* B728a led to the identification of a single aspartyl/asparaginyl β -hydroxylase at the primary locus Psyr_1584 (GenBank ID AAY36632.1), which we hereafter term *aspH*. Comparison of the protein sequence of AspH to other known bacterial iron and α -ketoglutarate-dependent aspartyl/asparaginyl hydroxylases indicates that it contains the conserved His₂-Asp facial triad responsible for iron binding as well as the two arginines required for binding α -ketoglutarate. Indeed, AspH shows significant homology to known aspartyl β -hydroxylases from *Alcanivorax* and *Ralstonia*, as well as to LpxO, an iron- and α -KG-dependent dioxygenase from *Salmonella*

typhimurium that hydroxylates lipid A (30). To circumvent difficulties in recombinant expression, *aspH* was obtained as a synthetic gene with codons optimized for expression in *E. coli*. The N-terminally His₆-tagged 35.7 kDa protein AspH was expressed in *E. coli* at 15 °C following induction with 0.1 mM IPTG and purified using nickel affinity chromatography to give a yield of 0.5 mg/L of culture (Figure 2A).

In addition, a construct comprising part of the eighth module of the SyrE megasynthetase, namely, SyrE-A₈T₈, was cloned into the pET-28a vector and expressed in *E. coli* as described above. With this 70.1 kDa construct, yields of 8–10 mg of protein/L of culture were obtained (Figure 2B).

ATP-PP_i Exchange Assays To Evaluate SyrE-A₈ Substrate Specificity. In order to obtain a first approximation of the timing of aspartyl hydroxylation in syringomycin biosynthesis, the ATP-PP_i exchange assay was performed to examine the substrate preference of the eighth adenylation domain of SyrE (SyrE-A₈). The ability of the adenylation domain of SyrE-A₈T₈ to activate L-Asp, L-erythro-3-OH-Asp, and L-threo-3-OH-Asp was assayed. It was found that A₈ preferentially activates unmodified L-Asp ($k_{\text{obs}} = 6.3 \text{ min}^{-1}$) in comparison to the erythro ($k_{\text{obs}} = 0.1 \text{ min}^{-1}$) or threo ($k_{\text{obs}} = 0.2 \text{ min}^{-1}$) 3-OH-Asp substrates (Figure 3). These results indicate that hydroxylation of Asp₈ in syringomycin biosynthesis occurs either on the assembly line or after cyclization and release of the cyclic lipopeptidolactone but is unlikely to occur on free aspartic acid.

Phosphopantetheinylation and Autoaminoacylation of SyrE-A₈T₈. While the ATP-PP_i exchange assay demonstrated the activity and substrate specificity of the adenylation domain of the SyrE-A₈T₈ construct, further work was necessary to prove the competence of the thiolation domain for covalent thioesterification with aspartic acid. Reaction of SyrE-A₈T₈ with coenzyme A and the phosphopantetheinyl transferase Sfp yielded the holo form of the thiolation domain. Further incubation with L-[¹⁴C]-Asp and ATP led to activation of the substrate and subsequent aminoacylation of the thiolation domain. Autoaminoacylation was monitored over time by TCA precipitation followed by liquid scintillation counting as well as by autoradiography (Figure 4).

Characterization of AspH Dioxygenase Activity with Free Amino Acids. In order to determine the timing of hydroxylation carried out by AspH, a number of substrates were used to test AspH activity. A chiral radio-HPLC assay was developed in order to monitor product formation and determine product chirality. In a typical assay, 1 μ M AspH was incubated with 250 μ M substrate, 0.5 mM (NH₄)₂Fe(SO₄), 0.5 mM α -KG, 0.5 mM ascorbate, 0.5 mM DTT, and 0.2 ng/ μ L catalase in 50 mM Hepes buffer for 30 min. In all cases, control reactions were carried out without enzyme, without (NH₄)₂Fe(SO₄), and without α -KG. Upon incubating AspH with L-[¹⁴C]Asp under the conditions described above, an enzyme, Fe(II), and α -KG-dependent peak was observed in the chiral radio-HPLC chromatogram (Figure 5A). The peak coeluted with the L-erythro-3-OH-Asp authentic standard, and LC-MS indicated its mass coincided with that of 3-OH-Asp (3-OH-Asp 149.10 [(M + H)⁺] calculated, 149.02 observed). The generation of the L-erythro product was quite unexpected, since syringomycin contains the L-threo diastereomer. In a similar set of experiments, D-Asp, D-Glu, L-Glu, D-Asn, and L-Asn were

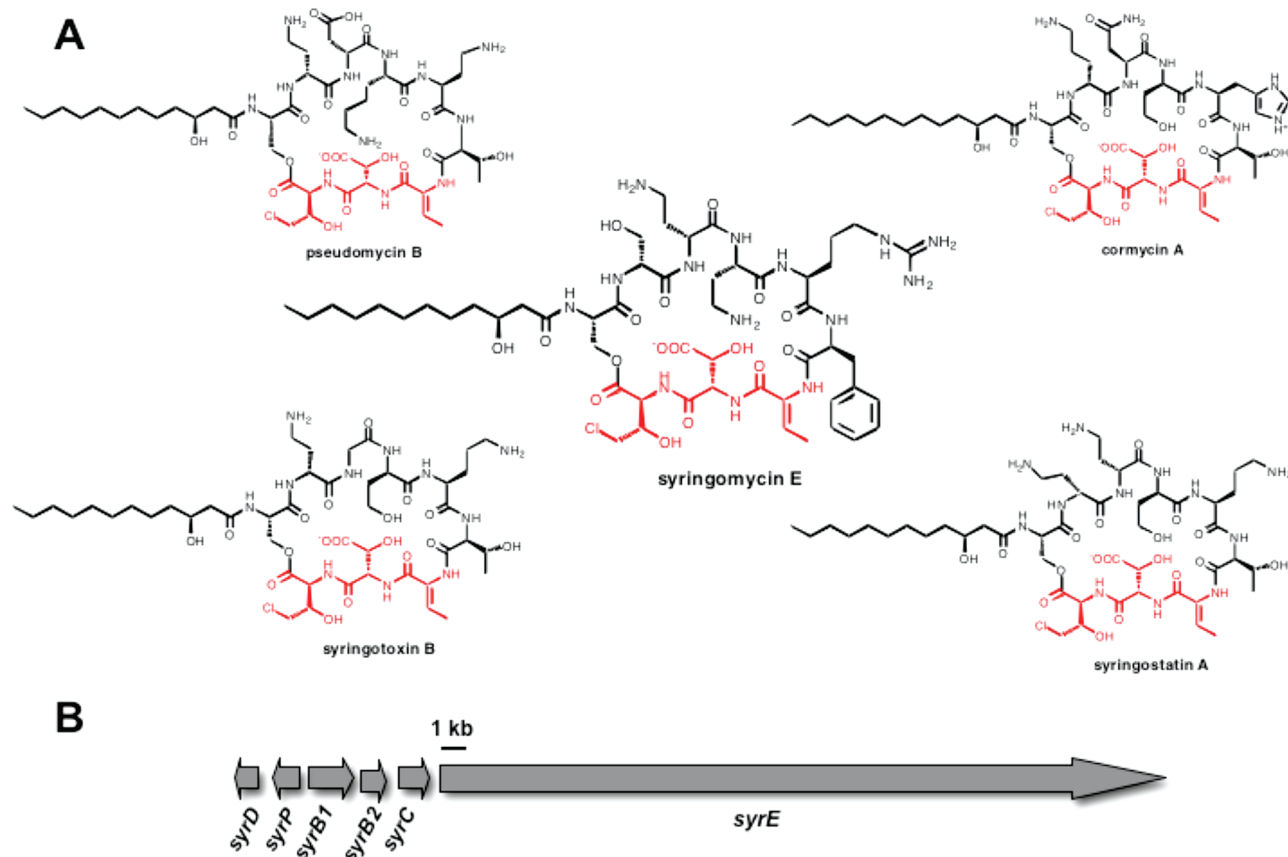


FIGURE 1: (A) Chemical structures of several pseudomonal lipopeptides containing the conserved C-terminal Z-Dhb₇-L-3-OH-Asp₈-L-4-Cl-Thr₉ tripeptide. (B) The syringomycin E gene cluster.

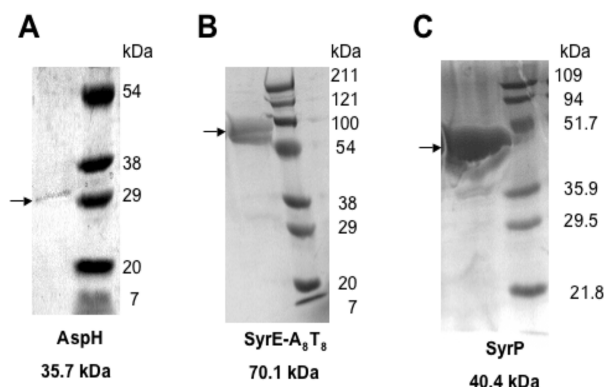


FIGURE 2: SDS-PAGE analysis of purified (A) AspH, 35.7 kDa, (B) SyrE-A₈T₈, 70.1 kDa, and (C) SyrP, 40.4 kDa.

tested as substrates for AspH, but no hydroxylation activity was noted for any of these substrates.

Characterization of AspH Dioxygenase Activity with Thioester-Linked Substrates. The kinetic parameters of AspH with L-[¹⁴C]Asp ($k_{\text{cat}} = 0.15 \text{ min}^{-1}$, $K_m = 51.9 \mu\text{M}$) suggest that the free substrate is suboptimal, so the question arose as to whether the enzyme requires a thioester-linked substrate as opposed to one with a free carboxylate. Therefore, the SyrE-A₈T₈-S-L-[¹⁴C]Asp substrate was generated by phosphopantetheinylation of SyrE-A₈T₈ via Sfp followed by autoaminoacylation of the holo construct with L-[¹⁴C]Asp as described above. The SyrE-A₈T₈-S-L-[¹⁴C]Asp substrate was incubated with AspH under standard conditions, and the product was released hydrolytically with LiOH. Chiral radio-HPLC and LC-MS analysis of the reaction products again showed formation of the L-erythro-3-OH-Asp product in an

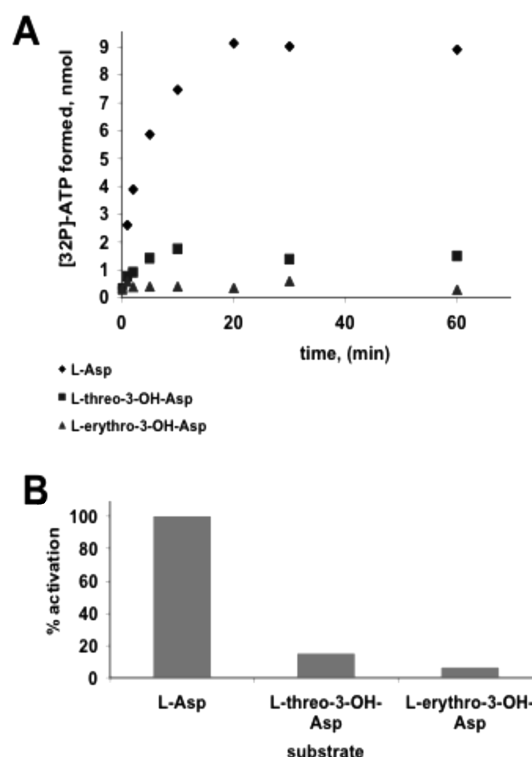


FIGURE 3: Results of the ATP-PP_i exchange assay to determine the substrate specificity of SyrE-A₈. (A) Determination of k_{obs} for three substrates: black diamonds, L-Asp, $k_{\text{obs}} = 6.3 \text{ min}^{-1}$; gray squares, L-threo-3-OH-Asp, $k_{\text{obs}} = 0.2 \text{ min}^{-1}$; white circles, L-erythro-3-OH-Asp, $k_{\text{obs}} = 0.1 \text{ min}^{-1}$. (B) Comparison of % activation by SyrE-A₈ of L-Asp, L-threo-3-OH-Asp, and L-erythro-3-OH-Asp.

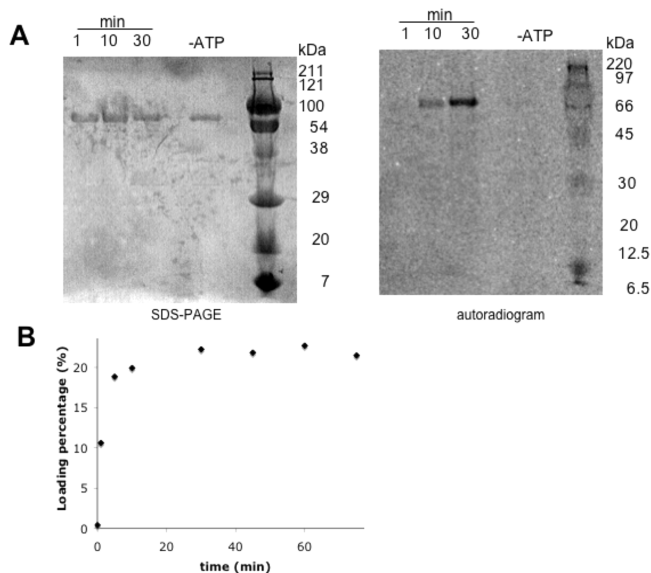


FIGURE 4: Autoaminoacylation of SyrE-A₈T₈ with L-[¹⁴C]Asp. (A) 12% SDS-PAGE and autoradiogram showing an ATP dependence of autoaminoacylation. (B) Graphical representation of time dependence of autoaminoacylation, as measured by a TCA precipitation assay.

enzyme, Fe(II), and α -KG-dependent fashion (3-OH-Asp 149.10 [(M + H)⁺] calculated, 149.02 observed) (Figure 5B).

Since it is difficult to accurately determine the concentration of the SyrE-A₈T₈-S-L-[¹⁴C]Asp substrate, the kinetic parameters of AspH with this substrate could not be determined. However, an aspartyl-S-N-acetylcysteamine (Asp-SNAC) substrate was generated as a mimic of the PCP-tethered substrate such that kinetic studies of AspH with a thioester-linked substrate could be carried out. The L-Asp-SNAC substrate was incubated with AspH under standard conditions, the thioester linkage was cleaved with LiOH, and the reaction products were analyzed as described. In this case we again noted the formation of the L-erythro-3-OH-Asp product (Figure 5C) and the mass of the 3-OH-Asp product was noted by LC-MS (3-OH-Asp 149.10 [(M + H)⁺] calculated, 149.02 observed). The kinetic parameters of the hydroxylation of the L-Asp-SNAC ($k_{\text{cat}} = 42.5 \text{ min}^{-1}$, $K_m = 43.6 \mu\text{M}$) were significantly improved over those for the free L-Asp substrate, with k_{cat} increasing by about 283-fold.

Characterization of AspH Dioxygenase Activity with Peptide Substrates. To determine AspH activity on a third class of substrates, a linear N-butyryl-nonapeptide mimic of the syringomycin liponapeptide was generated by chemical synthesis. The linear nonapeptide was incubated with AspH under standard conditions, and the reaction products were characterized by C₁₈ HPLC and LC-MS. In the HPLC chromatogram, an AspH-dependent peak was observed (Figure 5D), and the mass of the hydroxylated peptide was apparent by LC-MS (1098.53 [(M + H)⁺] calculated, 1099.04 observed). However, the exact position of hydroxylation in the nonapeptide could not be determined by these methods and would have to be determined by hydrolysis of the reaction product followed by chiral HPLC. It has thus far not been possible to generate sufficient material for the hydrolytic analysis of the hydroxylated product.

Although the exact position and stereochemistry of hydroxylation in the nonapeptide remain to be determined, the kinetic parameters of AspH with the linear liponapeptide

substrate were obtained ($k_{\text{cat}} = 120 \text{ min}^{-1}$, $K_m = 26.9 \mu\text{M}$) and suggest that this substrate is the most kinetically competent substrate of those tested, with an apparent catalytic efficiency (k_{cat}/K_m) improved 1500- and 4.6-fold over that of L-Asp and L-Asp-SNAC, respectively.

In combination, these data suggest that AspH preferentially modifies substrates in which the C-terminus of L-Asp is bound in a thioester or amide linkage, which may indicate that AspH is involved in posttranslational modification of peptides and proteins (Table 1). However, in all cases where stereochemistry could be determined, AspH generates the L-erythro-3-OH-Asp diastereomer, while syringomycin contains the L-threo diastereomer. This observation, in combination with the fact that AspH is located far upstream of the syringomycin gene cluster, strongly suggests that AspH is probably not involved in tailoring Asp₈ in syringomycin biosynthesis.

Overproduction and Purification of SyrP. The fact that AspH does not appear to be involved in syringomycin tailoring led us to reexamine the members of the syringomycin gene cluster in search of a putative dioxygenase, since it is most likely that the tailoring enzyme would be in or near the gene cluster itself. A BLAST analysis of SyrP, which was previously assigned as a regulator of syringomycin production, revealed a high degree of homology to TauD, the well-studied taurine dioxygenase (31–33) (Figure 6). Indeed, SyrP contained the conserved His₂-Asp facial triad responsible for iron binding, as well as the conserved Arg residues that act as α -KG ligands. This strongly suggested that SyrP might be the aspartyl dioxygenase responsible for hydroxylation of Asp₈.

A synthetic gene for *syrP* with codons optimized for expression in *E. coli* was cloned into the N-terminal His₆-tagged vector. Then SyrP was expressed in *E. coli* at 15 °C following induction with 0.1 mM IPTG. The soluble 40.4 kDa protein was purified using nickel affinity chromatography to produce a yield of approximately 10 mg/L of culture (Figure 2C).

Examination of SyrP Dioxygenase Activity with Asp-Containing Substrates. The ability of SyrP to act as an aspartyl hydroxylase was examined with a number of Asp-containing substrates, including free L-[¹⁴C]Asp, SyrE-A₈T₈-S-L-[¹⁴C]Asp, L-Asp-SNAC, the linear N-acyl-nonapeptide, the N-acyl-nonapeptidyl-CoA, and the cyclic N-dodecanoyl-nonapeptidolactone. Assays were performed as described above, and products were characterized by HPLC and LC-MS. However, SyrP did not generate a hydroxylated product from any of the listed substrates except SyrE-A₈T₈-S-L-[¹⁴C]Asp. Furthermore, with this substrate, the stereochemistry of the 3-OH-Asp product was found to be exclusively in the L-threo configuration (Figure 7). Interestingly, the SyrP-mediated hydroxylation was found to be dependent on α -KG but was independent of exogenous (NH₄)₂Fe(SO₄), leading to the hypothesis that SyrP is purified with Fe(II) bound. The Ferrene S assay (34) revealed that, indeed, pure SyrP contains 28% Fe(II).

These data suggest that SyrP is likely to be the Fe(II)- and α -KG-dependent aspartyl hydroxylase responsible for hydroxylating Asp₈ in syringomycin biosynthesis. In addition, the fact that SyrP acts exclusively on the PCP-tethered substrate indicates that hydroxylation occurs while the

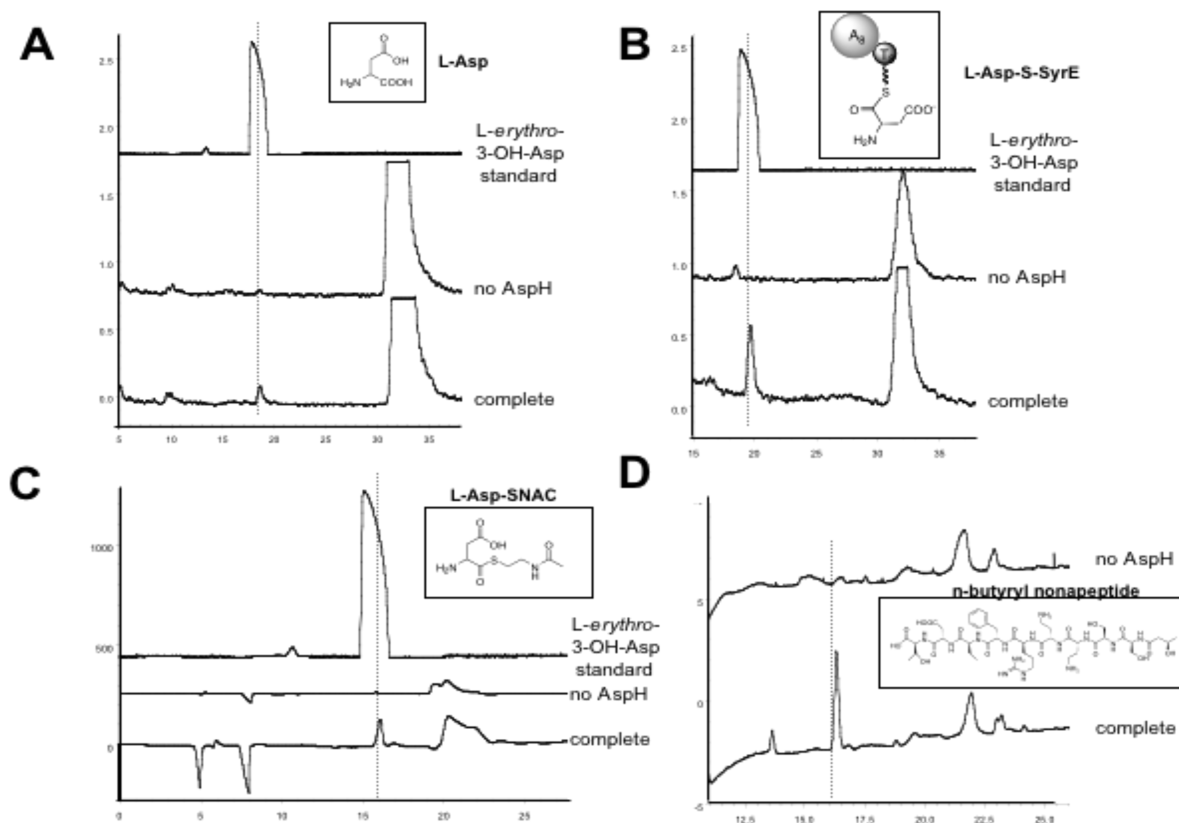


FIGURE 5: HPLC chromatograms of products from AspH incubation with various substrates. “Complete” denotes reactions containing all components, and “no AspH” denotes control reactions lacking the enzyme. The substrates are shown as insets in each trace. (A) Chiral radio-HPLC chromatogram of AspH incubation with L-[14 C]Asp. (B) Chiral radio-HPLC chromatogram of AspH incubation with SyrE-A $_8$ T $_8$ -S-L-[14 C]Asp. (C) Chiral HPLC of AspH incubation with L-Asp-SNAC with detection at 220 nm. (D) C18 HPLC of AspH incubation with the linear *N*-acyl-nonapeptide mimic of the syringomycin lipononapeptide.

Table 1: Kinetic Data Collected for AspH with Aspartate-Containing Substrates

substrate	k_{cat} (min^{-1})	K_{m} (μM)	$k_{\text{cat}}/K_{\text{m}}$
L-Asp	0.15 ± 0.03	52.9 ± 4.7	0.002
L-Asp-SNAC	42.5 ± 5.2	43.6 ± 2.3	0.97
linear nonapeptide	120.0 ± 15.4	26.9 ± 9.9	4.6

substrate is on the assembly line, corresponding with the results of the ATP-PP $_i$ exchange assay.

Properties of the SyrP-Mediated Hydroxylation Reaction. Due to the difficulty of determining the exact concentration of the SyrE-A $_8$ T $_8$ -S-L-[14 C]Asp substrate because of small variations in the loading of A $_8$ T $_8$ with L-[14 C]Asp, the kinetic parameters of SyrP dioxygenase activity with this substrate could not be determined. Despite this obstacle, the catalytic properties of the SyrP-mediated hydroxylation could be characterized by approximately calculating the amount of SyrE-A $_8$ T $_8$ -S-L-[14 C]Asp present using the TCA precipitation assay.

In order to determine whether product formation catalyzed by SyrP is proportional to the amount of substrate present, the hydroxylation reaction was analyzed at increasing concentrations of SyrE-A $_8$ T $_8$ -S-L-[14 C]Asp. This experiment revealed the linear correlation of the amount of SyrE-A $_8$ T $_8$ -S-L-[14 C]Asp to the amount of L-threo-3-OH-Asp product (Figure 8A). To further examine the catalytic role of SyrP, an experiment was carried out in which SyrE-A $_8$ T $_8$ -S-L-[14 C]Asp was held constant at 4 nmol and SyrP was titrated from 25 to 500 pmol. In this case, SyrP was found to catalyze

multiple turnovers at low SyrP:A $_8$ T $_8$ ratios, indicating that it does indeed function catalytically (Figure 8B).

Generation and Characterization of syrP and aspH Knockout Strains of *P. syringae* pv *syringae*. The ΔaspH and ΔsyrP strains of *P. syringae* pv *syringae* B301D were generated by the introduction of gene replacement vectors containing the FRT-flanked, *cat*-disrupted *aspH* or *syrP* genes into the parent strain as described in the Experimental Procedures. The deletions were returned to the *P. syringae* pv *syringae* B301D chromosome through successive selection steps for each of two homologous recombination events. The resulting *aspH::cat* and *syrP::cat* strains were then electroporated with the plasmid pFLP2. Subsequent expression of the Flp recombinase, which acts at the FRT sites flanking the *cat* gene in the *P. syringae* chromosome, resulted in Flp-catalyzed excision of the *cat* elements from the host chromosome and produced the unmarked ΔaspH and ΔsyrP strains. PCR was performed on genomic DNA isolated from the *aspH::cat*, *syrP::cat*, ΔaspH , and ΔsyrP strains to confirm the presence of the desired insertions and deletions.

The *syrP::cat*, ΔsyrP , and wild-type strains were grown in PDB media in still culture at 25 °C for 6 days, and the ΔaspH strain was grown under identical conditions for an additional 2 days due to slow growth. Cultures were extracted in water-saturated 1-butanol and were assayed for syringomycin production by HPLC and LC-MS (Figure 9). This analysis revealed that the ΔaspH strain produced roughly equivalent amounts of syringomycin (1225.6 [(M + H) $^+$] calculated, 1225.6 observed) as the wild-type strain, despite

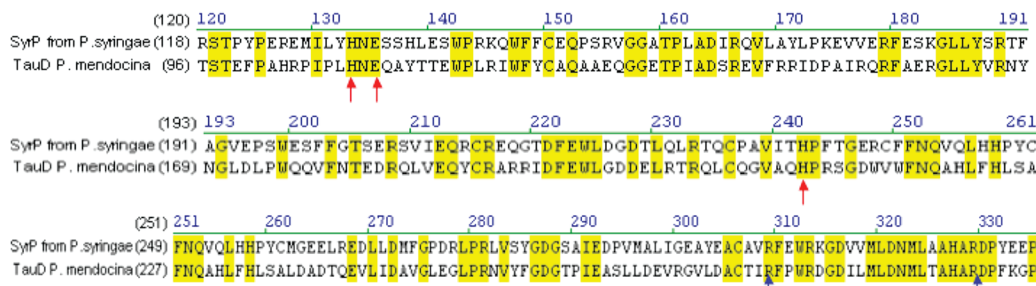


FIGURE 6: Alignment of the protein sequences of SyrP from *P. syringae* pv *syringae* B728a with that of TauD from *P. mendocina*. Red arrows indicate the His₂/Asp iron-binding motif, and blue arrows point to the two conserved Arg residues that bind α -ketoglutarate.

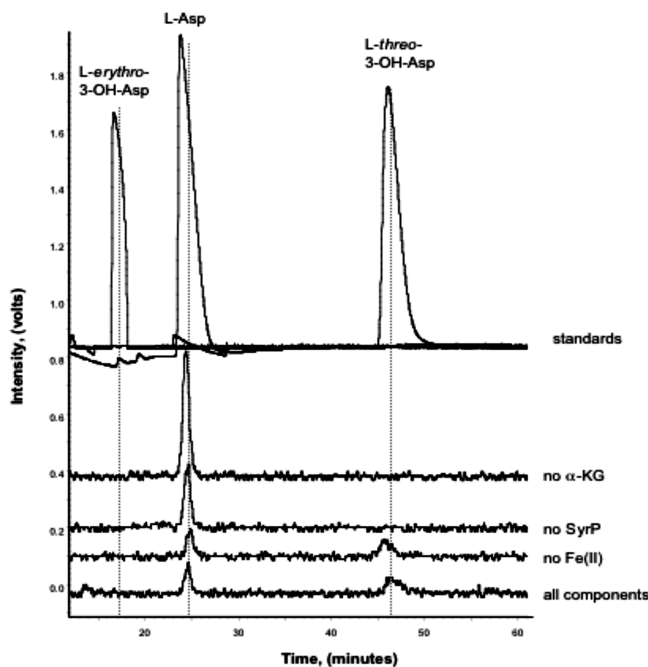


FIGURE 7: Chiral radio-HPLC traces of products from the reaction of SyrP with the SyrE-A₈T₈-S-L-[¹⁴C]Asp substrate and products of the control reactions. In the reaction containing all components, a peak at \sim 49 min is evident in the B-RAM chromatogram, corresponding to the retention time of the *L*-threo-3-OH-Asp authentic standard. A peak with similar retention time is evident in the control reaction lacking Fe(II), suggesting that SyrP may have been purified with Fe(II) already bound. The mass of the *L*-threo-3-OH-Asp product was verified by LC-MS.

its reduced growth rate, supporting the hypothesis that AspH is not involved in syringomycin biosynthesis. However, analysis of the *syrP::cat* strain indicated that syringomycin production was entirely abolished. This led us to suspect that the insertion of the *cat* gene in the *P. syringae* chromosome was causing polar effects on downstream genes in the syringomycin gene cluster. Subsequent HPLC analysis of the unmarked Δ *syrP* strain suggested that it, too, did not produce any syringomycin but instead produced a new compound with a retention time that is 0.3 min later than that of syringomycin. The LC-MS analysis of material eluting in this peak showed it to have *m/z* of 1216.6 rather than the *m/z* 1209.6 expected for the *des*-hydroxy species.

The *syrP* knockout strain was grown in the presence of [¹³C]Asp and [¹⁵N]NH₄Cl in order to obtain ¹³C- and ¹⁵N-labeled syringomycin derivatives. NMR studies of the compound produced by the Δ *syrP* species indicated that it is, in fact, an immature form of syringomycin lacking the β -OH form of Asp₈. However, due to difficulties in extracting

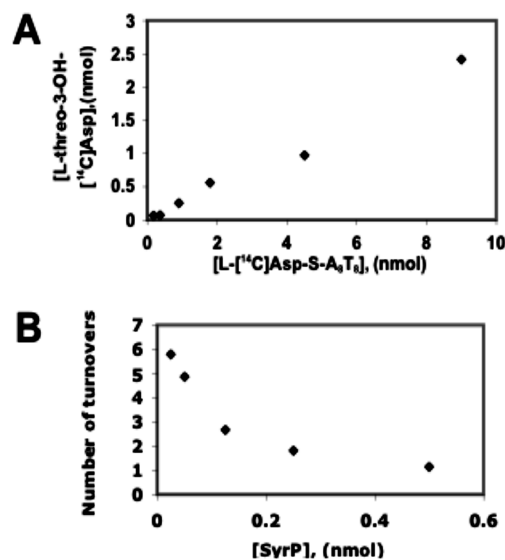


FIGURE 8: (A) In a SyrP-mediated reaction, formation of the *L*-threo-3-OH-[¹⁴C]Asp product is proportional to the amount of SyrE-A₈T₈-S-L-[¹⁴C]-Asp substrate. (B) When SyrE-A₈T₈-S-L-[¹⁴C]-Asp is held constant at 4 nmol and SyrP is titrated, multiple turnovers are observed at low SyrP:A₈T₈ ratios, indicating that SyrP functions catalytically.

sufficient amounts of the labeled compound, these NMR studies were unable to fully elucidate the structure of this species.

DISCUSSION

Syringomycin and related nonribosomal peptides from pseudomonads are pore-forming *N*-acyl-peptidolactones. The long-chain acyl group is added at the N-terminal residue at the start while the lactone/depsipeptide linkage is formed in the terminal chain release/cyclization step. The full-length nonapeptidyl chain, tethered as a pantetheinyl-S-peptide on the ninth and last module of the megasynthetase SyrE enzyme, is released by intramolecular capture of the thioester carbonyl of Cl-Thr₉ by the OH side chain of Ser₁. In the pseudomonal lipodepsipeptide family the residues at positions 7–9 are conserved as dehydrobutyrine₇-3-OH-Asp₈-4-Cl-Thr₉, arising respectively from the proteinogenic monomers, Thr, Asp, and Thr, in a set of enzyme-mediated tailoring steps.

In principle, the tailoring of those three residues could happen (a) prior to NRPS assembly line action, (b) during assembly line action, or (c) after release of the nonapeptidolactones from the assembly lines (35). Our prior studies on 4-Cl-Thr₉ have established that it is chlorinated on the assembly line, as a Thr-S-pantetheinyl-SyrB1 covalent

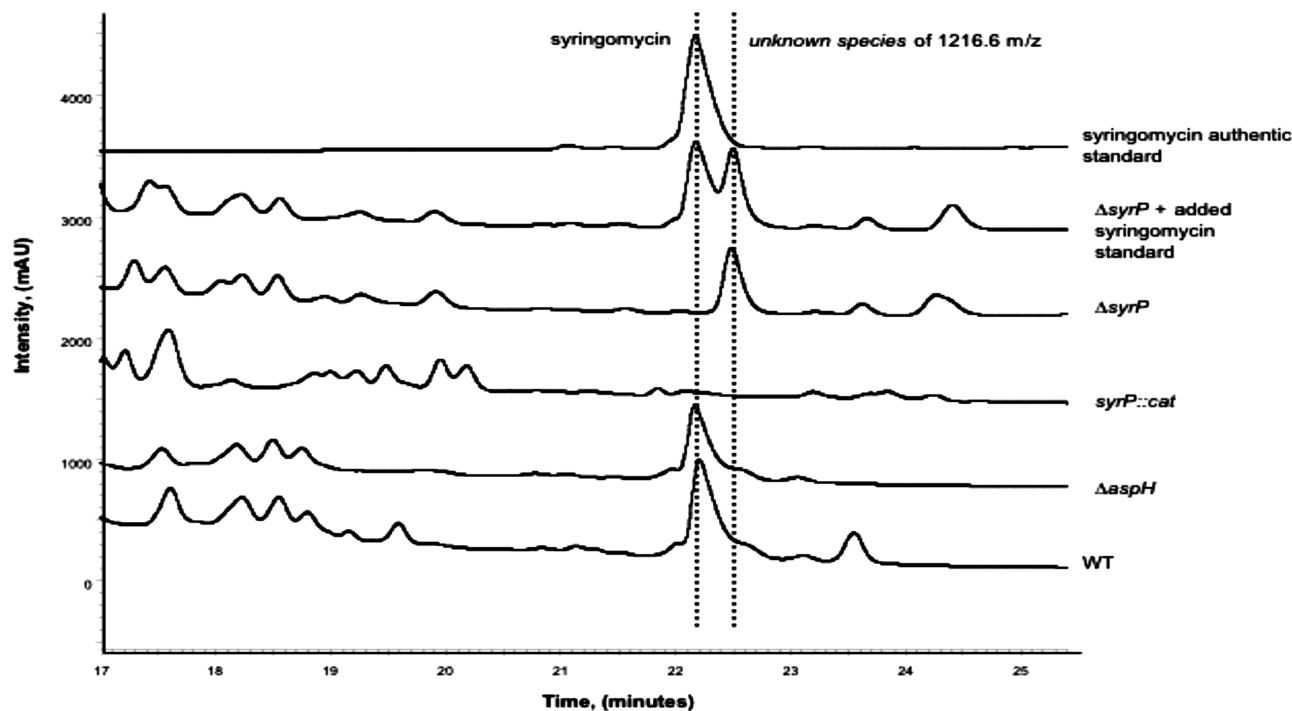


FIGURE 9: HPLC analysis of 1-butanol extracts of wild type, $\Delta aspH$, $syrP::cat$, $\Delta syrP$, and $\Delta syrP$ spiked with syringomycin authentic standard and syringomycin authentic standard. The traces show a new product formed in the $\Delta syrP$ culture with a retention time that is 0.3 min later than syringomycin. LC-MS analysis of this product shows it to have a mass of m/z 1216.6, and NMR studies indicate that it is an immature form of syringomycin, missing the β -OH group on Asp₈. Insufficient amounts of the ^{13}C - and ^{15}N -labeled compound were obtained for determination of the other structural variations in this compound.

intermediate and then transferred by the agency of the shuttle enzyme SyrC to the T₉ thiolation domain on SyrE. Nothing is known on the timing of dehydration of Thr₇ to dehydrobutyryne although it most likely occurs after Thr₇ has been incorporated into the growing chain to avoid enolization and adventitious hydrolysis to 2-ketobutyrate. In this study we have investigated the timing of Asp₈ enzymatic tailoring.

The adenylation–thiolation didomain pair, A₈T₈, expected to activate and load Asp₈ onto the syringomycin synthetase assembly line is embedded as part of the eighth module in the megadalton protein SyrE. We have been able to express A₈T₈ as a free-standing fragment and posttranslationally prime T₈ with phosphopantetheine via the enzyme Sfp (36) to yield the holo form of A₈T₈ as a soluble, well-folded 70.1 kDa protein. The A domain shows a preference for activating L-Asp by 60-fold over the 3-OH *erythro* diastereomer and 30-fold over the 3-OH *threo* diastereomer, consistent with the view that hydroxylation occurs after selection, activation, and installation of Asp as the *S*-pantetheinyl thioester on T₈. By use of [^{14}C]-L-Asp, we could generate A₈T₈-S-L-[^{14}C]Asp as a potential substrate for the two candidate hydroxylases, AspH and SyrP, examined in this work.

Based on precedent, we expected the conversion of the L-Asp residue into the L-3-OH-Asp residue in syringomycin would be carried out by an iron-based hydroxylase. Such an oxygenase could possibly be a heme protein but more likely would be a non-heme mononuclear iron with a two-His, one-carboxylate facial triad as the active site ligands (22, 33). Initial inspection of the syringomycin biosynthetic gene cluster annotations (15) did not reveal either type of iron hydroxylase within the cluster. Reasoning that such a hydroxylase might be encoded elsewhere in the genome or even be borrowed from another pathway, we examined all

the orfs in the producing *P. syringae* genome. The most likely candidate was an ORF at the primary locus Psyr_1584. Residues 149–229 of AspH align well with Asp non-heme iron hydroxylases from *Alcanivorax* and from *Ralstonia* as well as LpxO from *S. typhimurium*. In particular, the putative iron ligands, two histidines, and a conserved aspartate in an HXD array, as well as the two arginine side chains observed to coordinate α -ketoglutarate in this superfamily (22, 33), are all conserved, strongly suggesting AspH would be an Asp-selective α -ketoglutarate-requiring non-heme iron hydroxylase.

Indeed, AspH is such a catalyst. Our evidence shows that free aspartate is a detectable but poor substrate ($k_{cat} = 0.15 \text{ min}^{-1}$) for hydroxylation in an O₂- and Fe^{II}-dependent manner. Asp-SNAC, a soluble mimic of the Asp-S-pantetheinyl tether for covalent intermediates on NRPS assembly lines, has a 280-fold higher k_{cat} at 42 min^{-1} . While this might point to oxygenative tailoring on module 8 of SyrE, we also examined the ability of AspH to hydroxylate a linear *N*-acyl-nonapeptide surrogate of syringomycin and found an even higher k_{cat} of 120 min^{-1} . This latter finding, that an Asp side chain in the peptide is hydroxylated 800-fold faster than the free Asp amino acid, would argue that the function of AspH may be to hydroxylate Asp side chains in a peptide scaffold, but that has yet to be resolved.

However, in each of the above three contexts, free Asp, Asp-SNAC, and the linear nonapeptide, the AspH enzyme generated the L-*erythro* 3-OH diastereomer, with no detectable formation of the L-*threo* diastereomer. Since it is the L-*threo* form that is found in syringomycin, we were forced to the following conclusion. AspH, while indeed a mononuclear iron aspartyl β -hydroxylase with robust activity toward the aspartyl side chain when the carboxyl was

derivatized, is not likely to be involved in a syringomycin tailoring step.

At this juncture we returned to a bioinformatics approach by reexamining the Syr gene cluster and made an unanticipated observation about SyrP. The publication in 1997 (15) assigning a regulatory function to the 40 kDa SyrP protein noted similarity to phospho-transfer and phospho-receiver domains found in the *E. coli* chemotaxis protein CheA although SyrP lacked the domains associated with autophosphorylation activity. Despite this predicted lack of autophosphorylation capability, Zhang et al. suggested, but did not directly prove, that SyrP might have a regulatory phospho-transfer role in control of Syr gene cluster expression (5).

However, a contemporary BLAST analysis shows a high degree of homology between SyrP and the *P. mendocina* TauD protein rather than to chemotaxis homologues. TauD is the non-heme iron enzyme taurine hydroxylase, a paradigm for mechanistic and structural knowledge about the mononuclear Fe^{II} hydroxylase family (31–33). Most notably, SyrP contains the conserved HXE motif that comprises two of the three facial triad ligands to the Fe^{II} in the TauD active site. In addition, SyrP also has the third conserved His and the two Arg residues that function in this enzyme family to position the cosubstrate α -ketoglutarate.

To evaluate whether SyrP could indeed be the sought after syringomycin Asp hydroxylase, we procured a synthetic gene, expressed it in *E. coli*, and obtained a soluble, active hydroxylase. It was fortunate that we had prepared the variety of substrates used to profile AspH activity because SyrP showed no hydroxylase activity toward free Asp or the linear nonapeptide analogue of syringomycin. However, SyrP was active when presented with Asp-S-pantetheinyl T₈ in the context of the didomain A₈T₈ fragment excised from the nonamodular SyrE. Most reassuringly, chirality analysis after hydrolytic release of product from thioester linkage showed the product to be the desired *L*-threo-3-OH-aspartate. Kinetic characterization of SyrP with the incompletely loaded Asp-S-A₈T₈ 70 kDa protein substrate was not feasible, but the hydroxylation reaction is linearly dependent on SyrP concentrations and multiple turnovers are detected.

Several conclusions can be drawn. SyrP is indeed a mononuclear iron-containing Asp hydroxylase, but not one that acts on the free amino acid. It works on *L*-Asp tethered in pantetheinyl thioester tethered to the T₈ carrier protein domain of SyrE. This accords with our observation that A₈ activates Asp about 30 times faster than *L*-threo-3-OH-Asp, consistent with oxygenative tailoring after the A₈ adenylation domain has activated loaded Asp on the T₈ domain.

The generation of a Δ syrP strain of *P. syringae* pv *syringae* B301D indicated that mature syringomycin is no longer produced, but instead an immature form of the species, which lacks the β -OH moiety on Asp₈, is generated. NMR and MS data indicate that this immature form of syringomycin also lacks other structural features of the mature species, but insufficient amounts of this compound were obtained for complete structural characterization.

At this juncture SyrP joins the SyrB1/B2 couple as a third enzyme in the Syr gene cluster acting in tandem to tailor Asp₈ and Thr₉ on the NRPS assembly line. SyrC has been assigned by us (13) as the 4-Cl-Thr aminoacyl shuttle enzyme. SyrD is annotated as an efflux pump. As there are

no other genes in the cluster, the enzymatic dehydration of Thr₇ to dehydrobutyrine₇ during maturation of syringomycin has yet to be determined.

Finally, this work reveals that the *P. syringae* genome encodes two aspartyl hydroxylases, both non-heme mononuclear iron catalysts decarboxylating cosubstrate α -ketoglutarate during oxygen transfer. AspH and SyrP, respectively, are stereospecific for generating the *erythro*- or the *threo*-3-OH-aspartyl hydroxylated products. While SyrP acts on residue 8 in the assembly of the nonapeptidolactone syringomycin, the substrate and product from AspH action are not known. Unlike SyrP, AspH is not in an operon context that is revealing or predictive of function. Given that 3-OH-Asp residues are formed posttranslationally in proteins (16, 17), it may be that a comparative proteomics search of 3-OH-Asp-containing proteins in wild-type and AspH knock-out strains would produce mass spectrometric evidence of proteins generated by AspH action.

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