Aromatic Prenylation in Phenazine Biosynthesis
DIHYDROPHENAZINE-1-CARBOXYLATE DIMETHYLALLYLTRANSFERASE FROM STREPTOMYCES ANULATUS

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The bacterium Streptomyces anulatus 9663, isolated from the intestine of different arthropods, produces prenylated derivatives of phenazine 1-carboxylic acid. From this organism, we have identified the prenyltransferase gene ppzP. ppzP resides in a gene cluster containing orthologs of all genes known to be involved in phenazine 1-carboxylic acid biosynthesis in Pseudomonas strains as well as genes for the six enzymes required to generate dimethylallyl diphosphate via the mevalonate pathway. This is the first complete gene cluster of a phenazine natural compound from streptomycetes. Heterologous expression of this cluster in Streptomyces coelicolor MS12 resulted in the formation of prenylated derivatives of phenazine 1-carboxylic acid. After inactivation of ppzP, only nonprenylated phenazine 1-carboxylic acid was formed. Cloning, overexpression, and purification of PpzP resulted in a 37-kDa soluble protein, which was identified as a 5,10-dihydrophenazine 1-carboxylate dimethylallyltransferase, forming a C–C bond between C-1 of the isoprenoid substrate and C-9 of the aromatic substrate. In contrast to many other prenyltransferases, the reaction of PpzP is independent of the presence of magnesium or other divalent cations. The \( K_{m} \) value for dimethylallyl diphosphate was determined as 116 \( \mu \text{M} \). For dihydro-PCA, half-maximal velocity was observed at 35 \( \mu \text{M} \). \( K_{\text{cat}} \) was calculated as 0.435 s\(^{-1}\). PpzP shows obvious sequence similarity to a recently discovered family of prenyltransferases with aromatic substrates, the ABBA prenyltransferases. The present finding extends the substrate range of this family, previously limited to phenolic compounds, to include also phenoxy derivatives.

The transfer of isoprenyl moieties to aromatic acceptor molecules gives rise to an astounding diversity of secondary metabolites in bacteria, fungi, and plants, including many compounds that are important in pharmacotherapy. However, surprisingly little biochemical and genetic data are available on the enzymes catalyzing the C-prenylation of aromatic substrates. Recently, a new family of aromatic prenyltransferases was discovered in streptomycetes (1), Gram-positive soil bacteria that are prolific producers of antibiotics and other biologically active compounds (2). The members of this enzyme family show a new type of protein fold with a unique \( \alpha-\beta-\beta-\alpha \) architecture (3) and were therefore termed ABBA prenyltransferases (1). Only 13 members of this family can be identified by sequence similarity searches in the database at present, and only four of them have been investigated biochemically (3–6). Up to now, only phenolic compounds have been identified as aromatic substrates of ABBA prenyltransferases. We now report the discovery of a new member of the ABBA prenyltransferase family, catalyzing the transfer of a dimethylallyl moiety to C-9 of 5,10-dihydrophenazine 1-carboxylate (dihydro-PCA). Streptomyces strains produce many of prenylated phenazines as natural products. For the first time, the present paper reports the identification of a prenyltransferase involved in their biosynthesis.

Streptomyces anulatus 9663, isolated from the intestine of different arthropods, produces several prenylated phenazines, among them endophenazine A and B (Fig. 1A) (7). We wanted to investigate which type of prenyltransferase might catalyze the prenylation reaction in endophenazine biosynthesis. In streptomycetes and other microorganisms, genes involved in the biosynthesis of a secondary metabolite are nearly always clustered in a contiguous DNA region. Therefore, the prenyltransferase of endophenazine biosynthesis was expected to be localized in the vicinity of the genes for the biosynthesis of the phenazone core (i.e., PCA).

In Pseudomonas, an operon of seven genes named \( \text{phzABC-DEFG} \) is responsible for the biosynthesis of PCA (8). The enzyme PhzC catalyzes the condensation of phosphoenolpyruvate and erythrose-4-phosphate (i.e., the first step of the shikimate pathway), and further enzymes of this pathway lead to the intermediate chorismate. PhzD and PhzE catalyze the conversion of chorismate to 2-amino-2-deoxyisochorismate and the subsequent conversion to 2,3-dihydro-3-hydroxanthranilic acid, respectively. These reactions are well established biochemically. Fewer data are available about the following steps.

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\[ \text{PCA} \rightarrow \text{Chorismate} \rightarrow \text{DMAPP} \rightarrow \text{TPS} \rightarrow \text{DMAPP} \rightarrow \text{TPS} \rightarrow \text{Carboxylic acid} \]

2 The abbreviations used are: PCA, phenazine 1-carboxylic acid; LC, liquid chromatography; MS, mass spectrometry; ESI, electrospray ionization; DMAPP, dimethylallyl diphosphate; TAPS, 3-(2-hydroxy-1,1-bis(hydroxymethyl)ethyl)-(amino)-1-propanesulfonic acid; HPLC, high pressure liquid chromatography; MS\(^2\), second level tandem mass spectrometry; MS\(^3\), third level tandem mass spectrometry.
Dihydrophenazine-1-carboxylate prenyltransferase

(i.e. dimerization of 2,3-dihydro-3-hydroxyanthranilic acid, several oxidation reactions, and a decarboxylation, ultimately leading to PCA via several stable intermediates). From *Pseudomonas*, experimental data on the role of PhzF and PhzA/B have been published (8, 9), whereas the role of PhzG is yet unclear. Surprisingly, the only gene cluster for phenazine biosynthesis described so far from streptomycetes (10) was found not to contain a phzF orthologue, raising the question of whether there may be differences in the biosynthesis of phenazines between *Pseudomonas* and *Streptomyces*.

Screening of a genomic library of the endophenazine producer strain *S. anulatus* now allowed the identification of the first complete gene cluster of a prenylated phenazine, including the structural gene of dihydro-PCA dimethylallyltransferase.

**EXPERIMENTAL PROCEDURES**

**Bacterial Strains, Plasmids, and Culture Conditions**—*S. anulatus* 9663 has been isolated previously from the intestine of different arthropods (7, 11). It was grown in liquid YMG medium or on solid MS medium. For production of secondary metabolites, the medium described by Sedmera et al. (12) was used.

*Escherichia coli* XL1 Blue MRF, *E. coli* SURE (Stratagene, Heidelberg, Germany), *E. coli* BW 25113, and *E. coli* ET 12567 (pUB307) were used for cloning and were grown in liquid or on solid (1.5% agar) Luria-Bertani or SOB medium at 37°C. The REDIRECT technology kit for PCR targeting was obtained from Plant Bioscience Limited (Norwich, UK). For inactivation experiments, the *aac(3)IV/oriT* (apramycin resistance) cassette from pIJ773 (13) was used. Carbenicillin (50–100 μg ml<sup>-1</sup>), apramycin (50 μg ml<sup>-1</sup>), kanamycin ([50 μg ml<sup>-1</sup>]), chloramphenicol (25 μg ml<sup>-1</sup>), and nalidixic acid (20 μg ml<sup>-1</sup>) were used for selection of recombinant strains.

**Chemicals**—Kanamycin and carbenicillin were purchased from Genaxxon BioSciences GmbH (Biberach, Germany); apramycin and nalidixic acid were from Sigma; chloramphenicol was from Merck; and phenazine 1-carboxylic acid was from InFormatik. Dimethylallyl diphenolate was synthesized as described in Ref. 14. Endophenazine A was isolated from cultures of *Streptomyces cinnamomeus* DSM 1024 as described in Ref. 7.

**Genetic Procedures**—Standard methods for DNA isolation and manipulation were performed as described by Kieser et al. (15) and Sambrook et al. (16). DNA fragments were isolated from agarose gels by using a PCR purification kit (Amersham Biosciences). Genomic DNA was isolated from *Streptomyces* strains by lysozyme treatment and phenol/chloroform extraction as described by Kieser et al. (15).

**Construction and Screening of the Cosmid Library**—Chromosomal DNA from *S. anulatus* was partially digested with Sau3AI, dephosphorylated and then ligated into the BamHI sites of SuperCos 1 (Stratagene) according to the manufacturer’s instructions. The ligation products were packaged with Gigapack III XL (Stratagene) and transduced into *E. coli* SURE. Colony hybridization was performed on Hybond-N membranes (Amersham Biosciences). *ephzA* from *S. cinnamomeus* (10) was used as hybridization probe for the first screening of the library. The digoxigenin-labeled *ephzA* was generated using the PCR digoxigenin labeling mix (Roche Applied Science) with the primers *ephzA<sub>for</sub>* (5'-ATG AGC ACC CCC CTG ACC ACC-3') and *ephzA<sub>rev</sub>* (5'-TCA GGA GGG GAT CCA GTC CCG-3').

A second screening was performed by PCR for the identification of the following genes: *phzD*, *phzF*, *hmgr* (3-hydroxy-3-methyl-glutaryl-CoA reductase), *hmgs* (3-hydroxy-3-methylglutaryl-CoA synthase), and *mdpd* (mevalonate diposphate decarboxylase). The following primers were used: *PhzD<sub>for</sub>* (5'- CGC GCC GTC TCTG (A/G)TN CA(C/T) GA(C/T) (A/C/T)T-3') and *phzD<sub>rev</sub>* (5'- CGG TGG TGG TCC CGG (G/C)(A/T)(A/G) AA(A/G) TCN (G/C)-3'); *phzF<sub>for</sub>* (5' - CAT CGG GAT CTT GAC CCC NGT NAA (C/T)GA-3') and *phzF<sub>rev</sub>* (5'-GAG GGG GCC CCC ATT(C/T) TCN CAN CC-3'); *HMGR<sub>for</sub>* (5'-GGG CAT CGC CGC GAC CTT CCT GCN GGA GGA GGG-3') and *HMGR<sub>rev</sub>* (5'-GAG ATG ACG GGG AGG AGG CGC CGG TTC TC-3'); *HMGS<sub>for</sub>* (5'-GCC AAG GGG CCN GTN GA(C/T)GT-3') and *HMGS<sub>rev</sub>* (5'-AGG ATG ACG AAG GGG CCN GTN GC(T)/GT-3'); *MDPD<sub>for</sub>* (5'-GAC CCT GGA CGT CTT CCC NAC NAC NAC-3') and *MDPD<sub>rev</sub>* (5'-GCG TTC CCC TGCG GC(T)/G)T AT(C/T) TCN-3'). PCRs were carried out with Taq polymerase.

**Heterologous Expression of Cosmids ppzOS04 and ppzOS02**—The plasmid pIJ787 (17) was first digested with Dral and Bsal, and the 4990-bp fragment containing the integrase cassette was used to replace the *bla* gene in the SuperCos 1 backbone in cosmids 11C7 and 18A9, using λ RED-mediated recombination (18, 19), generating ppzOS02 and ppzOS04, respectively. Both cosmids were first transformed into the nonmethylating host *E. coli* ET12567, and the nonmethylated DNA was introduced into *Streptomyces coelicolor* M512 via triparental conjugation.

**Inactivation of the Gene ppzP**—An apramycin resistance cassette (*aac(3)IV*) was amplified from plasmid pUG019 (17) using the following primers: *ppzP<sub>for</sub>* (5'- CGG TGG TGG TCC CGG (G/C)(A/T)GCN GTN GA(C/T)GA-3') and *ppzP<sub>rev</sub>* (5'-GAC CCT GGA CGT CTT CCC NAC NAC NAC-3'). Underlined are restriction sites for XbaI and SpeI, used for later removal of the cassette. The resulting 1077-bp PCR product was used to replace the *ppzP* gene on cosmid ppzOS04 by λ RED-mediated recombination, resulting in cosmid ppzOS05. Deletion of the *aac(3)IV* cassette from ppzOS02 was carried out by digestion with XbaI and SpeI and religation, resulting in cosmid ppzOS09. The resulting construct was introduced into *S. coelicolor* M512 via triparental conjugation (15).

**Production and Analysis of Secondary Metabolites**—Exconjugants of all mutants as well as wild type *S. anulatus* were precultured for 48 h in liquid YMG medium (50 ml). 50 ml of production medium (20) was then inoculated with 2.5 ml of the precultures. The flasks were agitated on a rotary shaker at 30 °C and 200 rpm for 120 h. For cultivation of mutants, all liquid media contained kanamycin (50 μg ml<sup>-1</sup>). For isolation of endophenazine A, mycelia from 50-ml cultures were centrifuged at 3500 × g for 10 min. The supernatant was discarded, and the cells were extracted with methanol (10 ml) by vortexing. The extract was mixed with sodium acetate buffer (10 ml; 1 M, pH 4.0) and extracted with dichloromethane (5 ml). After separation of the organic phase, the solvent...
was evaporated, and the residue was redissolved in methanol (0.5 ml).

Extracts were analyzed with HPLC (Agilent 1100 series; Waldbronn, Germany) by using an Eclipse XDB-C18 column (4.6 \times 150 \text{ mm}, 5 \mu m; Agilent) at a flow rate of 1 ml min\(^{-1}\) with a linear gradient from 10 to 100\% of solvent B in 20 min (solvent A: water/phosphoric acid (999:1); solvent B, acetonitrile) and detection at 252 and 365 nm. Additionally, a UV spectrum from 200 to 400 nm was logged by a photodiode array detector. The absorbance at 365 nm was used for quantitative analysis, employing authentic reference samples of PCA and endophenazine A as external standards.

**Analysis by LC-MS**—The extracts were examined with LC-MS and LC-MS\(^2\) analysis using a Nucleosil 100-C18 column (3 \mu m, 100 \times 2 \text{ mm}) coupled to an ESI mass spectrometer (LC/MSD Ultra Trap System XCT 6330; Agilent Technology). Analysis was carried out at a flow rate of 0.4 ml min\(^{-1}\) with a linear gradient from 10 to 100\% of solvent B in 15 min (solvent A: water/formic acid (999:1); solvent B: acetonitrile/formic acid (999.4:0.6)). Detection was carried out at 230, 260, 280, 360, and 435 nm. Electrospray ionization (positive and negative ionization) was performed in positive ionization mode with capillary voltage of 3.5 kV at a heated capillary temperature of 350 °C was used for LC-MS analysis. For LC-MS\(^2\) and LC-MS\(^3\), the analysis was carried out in positive ionization mode with capillary voltage of 3.5 kV at 350 °C. For LC-MS\(^2\) identification of the enzymatic product endophenazine A, the mass 293 ± 0.5 Da was selected for fragmentation. In LC-MS\(^3\), the mass 275 ± 1 Da was selected for fragmentation.

**Overexpression and Purification of PpzP Protein—ppzP was amplified by using the primers ppzP\_pHis\_F (5’-ACC TGT GTA GGA GAA TTC ATG TCA GAA TCC GCT GAT CGT-3’) and ppzP\_pHis\_R (5’-CCG GAC GCG CTC GAC GCT ATC CGG CAT CGG CGG TCA-3’). The underlined letters represent EcoRI and Xhol restriction sites, respectively. The resulting PCR fragment was digested with EcoRI and Xhol and ligated into plasmid pHis8 (21) digested with the same restriction enzymes. The resulting plasmid, pHis8-OS01, was verified by restriction mapping and sequencing.**

**RESULTS**

**Cloning of a Biosynthetic Gene Cluster for Prenylated Phenazines—**A genomic library of the phenazine producer strain *S. anulatus* was constructed in cosmid vector SuperCos 1 (Stratagene). 2000 independent cosmid clones were subjected to a colony blot screening with a labeled probe of the phenazine biosynthesis gene *ephzA* from *S. cinnamonensis* (10). From the genome size of streptomycetes (~8.5 megabases) and the average insert size of SuperCos 1 (~38 kb), it could be estimated that a single genome locus would be represented, on average, in nine different cosmid clones. However, the screening revealed 26 positive clones, indicating that more than one genomic locus hybridized with *ephzA*. In order to identify those cosmids that contained the correct locus and that were likely to contain the entire gene cluster, these 26 cosmids were screened with degenerate primers for two further phenazine biosynthetic genes, *phzD* and *phzF*, and for three genes of the mevalonate pathway for isoprenoid biosynthesis (see “Experimental Procedures”). In another *Streptomyces* strain, it had been shown that the prenyl moiety of endophenazine A is formed via the mevalonate pathway (22), whereas in streptomycetes, the methyl erythritol phosphate pathway is used for the formation of isoprenoids of primary metabolism (23). Two cosmids, 18A9 and 11C7, were identified that gave PCR products with all five primer pairs. Sequencing of the PCR products confirmed that they represented *phzD*, *phzF*, and the three mevalonate pathway genes. Sequencing of cosmids 18A9 revealed a DNA region of 17.5 kb comprising 18 putative coding sequences (Fig. 1B), which together code for all enzymatic functions expected to be involved in endophenazine formation. The results of data base comparisons for these genes are listed in Table 1. Seven of the
putative coding sequences, designated as \textit{ppzA}, \textit{ppzC}, \textit{ppzD}, \textit{ppzE}, \textit{ppzF}, \textit{ppzG}, and \textit{ppzB}, showed obvious similarities to the seven genes that direct the biosynthetic pathway from phosphoenolpyruvate and erythrose-4-phosphate to PCA in \textit{Pseudomonas} strains (8, 9). In addition, the gene \textit{ppzM} was identified, showing striking similarity to \textit{phzM} from \textit{P. aeruginosa}. \textit{phzM} is proposed to be responsible for the \textit{N}-methylation in the biosynthesis of pyocyanine (1-hydroxy-5-\textit{N}-methylphenazine) (24). \textit{ppzM} may therefore be responsible for the \textit{N}-methylation reaction in the biosynthesis of endophenazine \textit{B} (Fig. 1A).

Sequence analysis further revealed a group of six coding sequences with striking similarity to genes of the mevalonate pathway (23), leading from acetyl-CoA and acetoacetyl-CoA to isopentyl diphosphate and DMAPP (see Table 1). Orthologs of these six genes, arranged in the exact same order, are found in the biosynthetic gene cluster of furaquinocin (25), naphterpin (26), terepenticin (27), and BE 40644 (28). \textit{S. anulatus}, there-
fore, is one of the rare Streptomyces species that possesses the genes of the mevalonate pathway, a feature limited to ~1% of the strains of this genus (29).

Directly adjacent to the genes of the mevalonate pathway, a gene designated ppzP was identified (Fig. 1B). Its predicted gene product showed 44% identity to Fnq26, a prenyltransferase of the ABBA family that is involved in the biosynthesis of the meroterpenoid furanonaphthoquinone I (5).

Heterologous Expression of the Endophenazine Cluster in S. coelicolor M512—A RED-mediated recombination in E. coli was used to replace the β-lactamase gene within the SuperCos 1 backbone of cosmid 18A9 with the cassette pIJ787 containing a tetracycline resistance gene and the integration function of the phage ΦC31 (13). The resulting cosmid ppzOS04 was introduced into the genome of S. coelicolor M512 using triparental conjugation (15).

As control, the same pIJ787 cassette was introduced in an empty SuperCos 1 vector, and the resulting construct, ppzOS30, was also integrated into the genome of S. coelicolor M512. The secondary metabolite production of the resulting integration mutants was analyzed by HPLC-UV and LC-MS, using the wild type strain S. anulatus as comparison. In accordance with previous results (7), S. anulatus was found to produce PCA and endophenazine A (Fig. 2A). The polar compound PCA was predominantly found in the culture medium, but some amount was also present in the mycelial extract. The more lipophilic, prenylated compound endophenazine A is predominantly found in cell extracts, not in the culture medium.

LC-MS analysis showed that both S. anulatus and S. coelicolor (ppzOS04) also produced a small amount of endophenazine B as well as phenazine 1-carboxylic acid methyl ester (data not shown). Both compounds were identified by LC-MS2 in comparison with authentic reference samples.

As mentioned above, the screening of the cosmid library had identified a second cosmid, 11C7. Sequencing of the termini of its insert and comparison with the sequence obtained from 18A9 showed that 11C7 lacked the gene ppzP (the putative prenyltransferase) and 744 nucleotides of the mevalonate pathway. The heterologous expression strain S. coelicolor (ppzOS04) produced both phenazine 1-carboxylic acid (most of which was found in the medium) and endophenazine A (most of which was associated with the cells) (Fig. 2B). The presence of both compounds was also shown by LC-MS at m/z = 225 [M + H]+ and m/z = 293 [M + H]+ for PCA and endophenazine A, respectively. LC-MS2 showed that both compounds had the same fragmentation pattern as the authentic reference samples (data not shown). In contrast, the control strain S. coelicolor M512 (ppzOS30), which had integrated the empty vector, did not produce any phenazines (data not shown).
kinase gene but contained all other genes of the putative endophenazine cluster (see Fig. 1B). Cosmid 11C7 was expressed heterologously in S. coelicolor M512 using the same method described above. The resulting strain S. coelicolor(ppzOS02) produced phenazine 1-carboxylic acid and its methyl ester but no prenylated phenazines (Fig. 2C).

The results of the heterologous expression experiment with 18A9 prove that this cosmid contains all of the genes required for endophenazine biosynthesis. The experiment with 11C7 indicates that ppzP may be responsible for the prenylation reaction in this pathway. However, 11C7 lacked not only ppzP but also the promoter of the putative operon of mevalonate biosynthesis genes (see Fig. 1B) as well as part of the mevalonate kinase gene. Therefore, the heterologous expression of 18A9 and 11C7 did not provide conclusive evidence for the function of ppzP.

Inactivation of the Phenyltransferase Gene ppzP and Heterologous Expression of the Resulting Construct in S. coelicolor M512—A RED-mediated recombination was used to create an in-frame deletion of the gene ppzP within the integrative cosmid ppzOS04. For this purpose, the cassette plg773, containing the apramycin-resistant gene acc(5)/IV and flanked with XbaI and SpeI restriction sites, was used to replace the coding sequence of the ppzP gene. Subsequently, the cassette was excised by XbaI and SpeI digestion and religation of the compatible overhangs, leaving only 6 bp as a “scar” sequence between the start and the stop codon of ppzP. The resulting integrative cosmid ppzOS09 was introduced into the genome of S. coelicolor M512 as described above. HPLC and LC-MS analysis of the cultures (Fig. 2D) showed phenazine 1-carboxylic acid and its methyl ester but no endophenazine A or B. This strongly supported the hypothesis that ppzP is responsible for the prenylation reaction in endophenazine biosynthesis in the heterologous expression experiment. However, a polar effect of the deletion on the expression of the genes downstream of ppzP cannot be excluded with certainty.

Overexpression and Purification of PpzP—For the biochemical investigation of the putative prenyltransferase, the gene ppzP was cloned into a plasmid for expression as an N-terminally His-tagged protein (see “Experimental Procedures”). The resulting construct was introduced into E. coli BL21 (DE3)pLysS. Induction with isopropyl 1-thio-β-D-galactopyranoside resulted in expression of a protein of ~37 kDa, as determined by SDS-PAGE (Fig. S1), coinciding with the calculated molecular mass of 37.138 kDa. Ni²⁺ affinity chromatography resulted in a protein of ~90% purity. 30 mg of purified soluble PpzP were obtained from 2 liters of culture. This protein was used for the biochemical investigations described below.

Identification of PpzP as 5,10-Dihydrophenazine 1-Carboxylate Prenyltransferase—Phenazine 1-carboxylic acid has been suggested as the product of the core pathway of phenazine biosynthesis from which the differently substituted or dimerized phenazines are derived by various tailoring reactions (8). In different Pseudomonas strains, PCA is converted by the monooxygenase PhzO to 2-hydroxy-PCA, by the amide synthetase PhzH to phenazine 1-carboxamide, and by the consecutive action of PhzM (methyltransferase) and PhzS (oxidoreductase) to pyocyanine (1-hydroxy-5-N-methyl-phenazine). However, when we incubated PCA with PpzP and DMAPP under various conditions, we did not observe product formation in HPLC analysis. From chemical reasoning, the reduced form of PCA (i.e. dihydro-PCA) (Fig. 3A), appeared to be an attractive candidate for the prenylation substrate, since it is much more reactive than PCA for an electrophilic attack at C-9. Dihydro-PCA is presumed to be the direct biosynthetic precursor of PCA (8). We therefore generated dihydro-PCA by reduction of PCA with sodium dithionite (Fig. 3A). When dihydro-PCA was incubated with PpzP and DMAPP, the time-dependent formation of an enzymatic product was readily observed by HPLC (Fig. 3B). Due to the very rapid oxidation of dihydro-PCA and its derivatives, they could not be quantified in the reduced form. After incubation, the reaction mixture was therefore oxidized with sodium persulfate (Na₂S₂O₈), and substrate and product were analyzed in the oxidized form. The enzymatic product was thereafter identified as endophenazine A by LC-MS analysis showing the same retention time, UV spectrum, molecular ion (m/z = 293 in positive ionization), and fragmentation pattern in ESI-MS² and ESI-MS³ as an authentic reference sample of endophenazine A.

Biochemical Properties of PpzP—In the enzymatic assay described under “Experimental Procedures,” the formation of endophenazine A showed a linear dependence on the amount of PpzP (up to 1 μg) and on the reaction time (up to 45 min). The reaction was strictly dependent on the presence of active PpzP, DMAPP, and dihydro-PCA. Maximal product formation was observed at pH 7.5, with half-maximal values at pH 9.5 and pH 5.0. In sharp contrast to the trans-prenyldiphosphate synthases like FPP synthase (30) and to the aromatic prenyltransferase NphB of naphterpin biosynthesis (3), the catalytic activity of PpzP was independent of the presence of Mg²⁺ or any other divalent metal ions. The addition of EDTA (10 mM) even increased reaction velocity 1.5-fold. Similarly, the addition of 500 mM NaCl increased product formation 3-fold, whereas the addition of 50 mM MgCl₂ and 100 mM CaCl₂ increased product formation ~1.5-fold. Therefore, 500 mM NaCl was routinely included in all assays (see “Experimental Procedures”). Although 10 mM Zn²⁺ had no effect on the reaction, the addition of 10 mM FeSO₄ completely abolished the formation of endophenazine A.

PpzP was found to be specific for both DMAPP and dihydro-PCA. When geranyldiphosphate was used instead of DMAPP, no product formation was observed. Likewise, no prenylated products could be observed when PpzP was incubated with other phenazine substrates, such as phenazine, phenazine 1-carboxylic acid methyl ester, N-methyl-phenazine (as methyl sulfate salt), or pyocyanine (1-hydroxy-5-N-methyl-phenazine). These phenazine substrates were reduced with dithionite to their dihydro analogs in the same way as described for PCA. Since N-methyl-phenazine and pyocyanine are compounds with quaternary nitrogen, these reaction mixtures were analyzed directly without prior extraction with ethyl acetate.

We also tested the aromatic substrates of previously examined ABBA prenyltransferases (i.e. 4-hydroxyphenylpyruvate, flavilon (2,5,7-trihydroxynaphthoquinone), 1,3-dihydroxy-naphthalene, and 1,6-dihydroxynaphthalene). Of these, only flavilon was prenylated by PpzP in the presence of DMAPP. LC-MS confirmed that the product was a monoprenylated fla-
violin derivative. However, the reaction velocity was only 0.5% of that obtained with dihydro-PCA. A reaction mechanism of the C-prenylation of flavilin under catalysis of Fnq26 was suggested by Haagen et al. (5). A similar mechanism may be expected for the prenylation of flavilin by PpzP. However, the amount of the prenylated product was too low for a structural identification by NMR spectroscopy.

Using a constant concentration of dihydro-PCA (0.8 mM) and varying concentrations of DMAPP, a typical hyperbolic curve of product formation over substrate concentration was obtained (Fig. 3C), indicating that the reaction followed Michaelis-Menten kinetics. Nonlinear regression analysis resulted in a $K_m$ value of $116 \pm 9 \mu M$ and a $K_{cat}$ of $0.435 \pm 0.006$ s$^{-1}$. Using different concentrations of dihydro-PCA in the presence of 0.4 mM DMAPP, a sigmoidal dependence of product formation on substrate concentration was observed (Fig. 3D). The half-maximal reaction velocity was obtained at $\approx 35 \mu M$ dihydro-PCA. Using this value (35 $\mu M$) as an estimate for the $K_m$ value for dihydro-PCA, the catalytic efficiency ($K_{cat}/K_m$) of PpzP was calculated as $12,400 \text{ M}^{-1} \text{s}^{-1}$. This is significantly higher than the value of $7.7 \text{ M}^{-1} \text{s}^{-1}$ reported for the conversion of the (artificial) substrate 1,3-dihydroxynaphthalene by NphB (6) but is comparable with the value of $5280 \text{ M}^{-1} \text{s}^{-1}$ calculated for the prenylation of the (genuine) substrate 4-HPP by CloQ (4). We therefore suggested that dihydro-PCA is the genuine substrate of PpzP.

**Investigation of Fnq28 from S. cinnamonensis DSM 1042 for Dihydro-PCA Dimethylallyltransferase Activity**—Our group recently reported a gene cluster of prenylated naphthoquinone and prenylated phenazine biosynthesis from *S. cinnamonensis* DSM 1042 (10). This strain produces, besides the prenylated naphthoquinone derivative furanonaphthoquinone I, the same prenylated phenazines as *S. anulatus* (10, 12, 20). The gene cluster from *S. cinnamonensis* contains genes assigned to furanonaphthoquinone I biosynthesis as well as a contiguous group of six genes (i.e. ephzBCDEGA) assigned to phenazine biosynthesis. By gene inactivation (10) and biochemical investigation (5), the prenyltransferase Fnq26 had been identified as the aromatic prenyltransferase of furanonaphthoquinone I biosynthesis. A second gene with similarity to ABBA prenyltransferases, designated as *fnq28*, was found immediately adjacent to the genes of the phenazine biosynthesis. *Fnq28* shows 36% sequence identity with PpzP on the amino acid level. We tested whether *Fnq28* also prenylates dihydro-PCA. For this purpose, *Fnq28* was expressed as an N-terminally His-tagged protein, using the same method as employed for PpzP. Ni$^{2+}$ affinity chromatography readily yielded a soluble protein of apparent homogeneity (data not shown). However, no prenylation of
Dihydrophenazine-1-carboxylate prenyltransferase

dihydro-PCA was observed with this protein, in clear contrast to the results with PpzP.

Fnq26 has been shown previously to transfer a geranyl moiety to flavioliol (2,5,7-trihydroxynaphthoquinone) or to the artificial substrate 1,3-dihydroxynaphthalene. We now tested whether Fnq26 could also use dihydro-PCA and either DMAPP or geranyldiphosphate as substrates. However, again no prenylation products were detected. These observations are consistent with the results from inactivation experiments, which had shown that endophenazine A is still formed in *S. cinnamonensis* after inactivation of both *fnq26* and *fnq28* (10). Therefore, neither Fnq26 nor Fnq28 catalyzes the prenylation reaction of endophenazine biosynthesis in *S. cinnamonensis*. In this organism, the enzyme that functionally corresponds to PpzP of *S. anulatus* is yet to be identified.

**DISCUSSION**

In this study, we have identified, for the first time, a prenyltransferase involved in the biosynthesis of prenylated phenazines. Sequence similarities and biochemical properties suggest that PpzP belongs to the recently discovered family of ABBA prenyltransferases (1). The present functional characterization of PpzP as 5,10-dihydrophenazine-1-carboxylate 9-dimethylallyltransferase now extends the substrate range of this family, previously limited to phenolic compounds, to include also phenazine derivatives.

At present, 13 genes with sequence similarity to ABBA prenyltransferases can be identified in the data base. A phylogenetic analysis of these genes (1) separates them into two clades. One of them comprises the 4-hydroxyphenylpyruvate 3-dimethylallyltransferases CloQ and NovQ from *Streptomyces* strains as well as four genes of unknown function from fungal genomes. The other clade comprises genes involved in the biosynthesis of prenylated naphthoquinones in different streptomycetes. A ClustalX analysis (data not shown) places PpzP into this second clade. Its closest ortholog is Fnq26, the prenyltransferase of furanonaphthoquinone I biosynthesis (5).

The x-ray structural analysis of the ABBA prenyltransferase NphB (formerly designated as Orf2) had revealed a novel protein fold, characterized by a β/α-barrel with 10 antiparallel β-strands. A structural model of PpzP (Fig. S2), generated using NphB as template, suggests a close similarity of the three-dimensional structure of the two proteins. The active center of NphB is localized within the spacious central cavity of the barrel, which has been suggested to explain the promiscuity of this enzyme for different aromatic substrates (3).

NphB requires the presence of Mg$^{2+}$ for catalytic activity. X-ray crystallography has shown that the Mg$^{2+}$ ion is coordinated by a single aspartate residue, by four water molecules, and by one of the oxygen atoms of the α-phosphate group of the isopenoid diphasphate (3). In contrast, PpzP does not require Mg$^{2+}$ ions for its activity, and the same has been reported for the ABBA prenyltransferases CloQ (4), SCC01790 (3) and Fnq26 (5). Modeling studies suggested that only charged residues (Lys$^{54}$ in CloQ, Arg$^{51}$ in SCC01790, and Arg$^{50}$ in Fnq26) may functionally substitute for Mg$^{2+}$ in the binding of the α-phosphate group of the isopenin diphosphate (1, 3). An alignment of PpzP with the enzymes named above (Fig. S3) shows that also PpzP contains an arginine residue (Arg$^{49}$) in the respective position of the first β strand. This may explain the Mg$^{2+}$ independence of the PpzP reaction. Like the other ABBA prenyltransferases, PpzP does not contain the DDXXD motif typical for trans-prenyldiphosphate synthases (1).

In the biosynthesis of phenazines in different microorganisms, a common pathway leads to PCA, and this compound was believed to represent the branching point from which the pathways to differently substituted phenazines diverge (8). Our study now suggests that, at least for the prenylation of phenazines, the branching point is dihydro-PCA rather than PCA. This compound has been suggested (8) to be the product of the last enzyme-catalyzed step of the biosynthesis of the phenazine core, whereas its oxidation to PCA may occur nonenzymatically. Therefore, dihydro-PCA is likely to be available as substrate for PpzP *in vivo*.

Our study reports the first complete gene cluster of phenazine biosynthesis from streptomycetes, after a previously identified cluster turned out to be incomplete (10). Natural phenazines have important biological activities, including antibacterial, antitumor, antioxidant, and testosterone 5α-reductase-inhibiting activity (31). The present discovery of the endophenazine gene cluster from *S. anulatus* and the functional identification of PpzP as dihydro-PCA prenyltransferase may pave the way to the generation of new prenylated phenazines with improved biological activities by metabolic engineering and chemo-enzymatic synthesis.

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