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Published in:
Applied and Environmental Microbiology

Link to article, DOI:
10.1128/AEM.07955-11

Publication date:
2012

Document Version
Publisher's PDF, also known as Version of record

Citation (APA):
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Published Ahead of Print 27 April 2012.

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Involvement of a Natural Fusion of a Cytochrome P450 and a Hydrolase in Mycophenolic Acid Biosynthesis

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Mycophenolic acid (MPA) is a fungal secondary metabolite and the active component in several immunosuppressive pharmaceuticals. The gene cluster coding for the MPA biosynthetic pathway has recently been discovered in *Penicillium brevicompactum*, demonstrating that the first step is catalyzed by MpaC, a polyketide synthase producing 5-methylorsellinic acid (5-MOA). However, the biochemical role of the enzymes encoded by the remaining genes in the MPA gene cluster is still unknown. Based on bioinformatic analysis of the MPA gene cluster, we hypothesized that the step following 5-MOA production in the pathway is carried out by a natural fusion enzyme MpaDE, consisting of a cytochrome P450 (MpaD) in the N-terminal region and a hydrolase (MpaE) in the C-terminal region. We verified that the fusion gene is indeed expressed in *P. brevicompactum* by obtaining full-length sequence of the *mpaDE* cDNA prepared from the extracted RNA. Heterologous coexpression of *mpaC* and the fusion gene *mpaDE* in the MPA-nonproducer *Aspergillus nidulans* resulted in the production of 5,7-dihydroxy-4-methylphthalide (DHMP), the second intermediate in MPA biosynthesis. Analysis of the strain coexpressing *mpaC* and the *mpaD* part of *mpaDE* shows that the P450 catalyzes hydroxylation of 5-MOA to 4,6-dihydroxy-2-(hydroxymethyl)-3-methylbenzoic acid (DHMB). DHMB is then converted to DHMP, and our results suggest that the hydrolase domain aids this second step by acting as a lactone synthase that catalyzes the ring closure. Overall, the chimeric enzyme MpaDE provides insight into the genetic organization of the MPA biosynthesis pathway.

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Fungi are among the most elaborate chemical producers in nature, producing a range of secondary metabolites, including some that are mycotoxins, food additives, or pharmaceutical drugs. Among the latter is mycophenolic acid (MPA), which is the active component in several immunosuppressants. MPA has also been associated with antiviral, antifungal, antibacterial, and anti-tumor activities (11). Consequently, MPA biosynthesis has received considerable research interest, and a biosynthetic route has been established through chemical labeling and culture feeding studies (3). MPA is a meroterpenoid proposed to be derived from a nonreduced tetraketide moiety through a five-step process involving oxidation, lactonization, and condensation with a farnesyl residue, followed by oxidative cleavage of the terpene part and a final methylation step (Fig. 1A). However, the biosynthesis remained unelucidated at the genetic level until the recent discovery of a putative MPA biosynthetic cluster in *Penicillium brevicompactum* (23). The defined cluster contained eight putative open reading frames (ORFs), including *mpaC*, which encodes a polyketide synthase catalyzing the production of 5-methylorsellinic acid (5-MOA), the first step in MPA biosynthesis (9, 23). Furthermore, it was recently demonstrated that *mpaF* encodes an MPA-insensitive inosine-5'-monophosphate dehydrogenase (IMPDH) conferring self-resistance toward MPA (8, 10). These studies utilized *Aspergillus nidulans* as a fungus of choice for the heterologous expression of *mpaC* and *mpaF*. *A. nidulans* provides a good model system to study the biosynthesis of MPA, since it can produce polyketides and does not produce any of the intermediates in MPA biosynthesis. In addition, the genome has been sequenced, and in the recent years the molecular biology toolbox has been greatly expanded (9, 16, 17, 19).

In the present study, we set out to identify and characterize the enzyme(s) responsible for the conversion of 5-MOA to 5,7-dihydroxy-4-methylphthalide (DHMP), which are the first and second known intermediates in MPA biosynthesis. A bioinformatics study of the MPA biosynthetic cluster, followed by heterologous expression in *A. nidulans*, showed that the conversion of 5-MOA to DHMP is catalyzed by a natural fusion of MpaD, a cytochrome P450, and MpaE, a putative lactone synthase.

MATERIALS AND METHODS

Strains and media. The following strains were used in the present study: *P. brevicompactum* strain IBT23078 and *A. nidulans* strains NID210 (argB2 pyrG89 veA1 IS1::PgdA-TrpC::argB, NID211 (argB2 pyrG89 veA1 IS1::PgdA-mpaC-TrpC::argB), NID410 (argB2 pyrG89 veA1 nkuAD IS2::PgdA-mpaDE-TrpC::AFpyrG), NID416 (argB2 pyrG89 veA1 IS1::PgdA-mpaC-TrpC::argB IS2::PgdA-mpaDE-TrpC::AFpyrG), and NID944 (argB2 pyrG89 veA1 IS1::PgdA-mpaC-TrpC::argB IS2::PgdA-mpaDE-TrpC::AFpyrG). Strains NID210 and NID211 were constructed in a previous study (9).

*P. brevicompactum* was grown on Czapek yeast extract agar (CYA) at 25°C. CYA is composed of 5 g of yeast extract (Biokar Diagnostics, Beauvais, France)/liter, 15 g of agar/liter, 35 g of Czapek dox broth/liter, 10 mg of ZnSO₄·7H₂O/liter, and 5 mg of CuSO₄·5H₂O/liter. The pH levels were adjusted to 6.5 with NaOH/HCl. *A. nidulans* strains were grown on minimal medium (MM) containing 1% glucose, 10 mM NaNO₃, 1% salt.
solution (5), and 2% agar for solid media or YES medium containing 20 g of yeast extract (Biokar)/liter, 150 g of sucrose/liter, 0.5 g of MgSO$_4$·H$_2$O and ZnSO$_4$·7H$_2$O/liter, 5 mg of CuSO$_4$·5H$_2$O/liter, and 2% agar (pH 6). The MM was supplemented with 10 mM uridine (Uri), 10 mM uracil (Ura), and 4 mM L-arginine (Arg) when necessary.

RNA purification and cDNA synthesis. Spores from *P. brevicompactum* IBT23078 were harvested and used to inoculate 200 ml of YES medium in 300 ml of shake flasks without baffles. *P. brevicompactum* was grown at 25°C and 150 rpm with shaking. After 48 h, the mycelium was harvested, and the RNA was purified using the fungal RNA purification miniprep kit (EZNA) according to the manufacturer’s instructions. cDNA was synthesized from the RNA using a Finnzymes Phusion RT-PCR kit according to the manufacturer’s instructions. The *mpaDE* transcript was amplified with the primer pair 657 and 660 (657, ATGAAGTC TTTGTCGCTAAC; 660, TTACTTCTGTCCTTCTATGG) and cloned into pJET1.2/blunt using a CloneJET PCR cloning kit (Fermentas) according to the manufacturer’s instructions, resulting in pJet_mpaD_mpaE.

Plasmid construction. Amplification of DNA by PCR to produce DNA fragments suitable for USER cloning was performed in 30 PCR cycles using PfuX7 in 50 µl. USER cloning was performed as previously described (9, 22), with minor modifications. The USER vectors were digested for 6 h with AsISI and Nb.BsmI and AsISI/Nb.BtsI USER cassettes or with PciI for the PciI/Nt.BbvCI USER cassettes A and B, followed by digestion with the appropriate nicking endonuclease for 1 h. Then, 0.1 pmol of purified digested vector was mixed with 1 pmol of purified PCR products amplified with primers that were extended by the appropriate tails for USER cloning into a designated USER cassette.

The PgpdA::USER cassette (AsISI/Nb.BtsI)::TrpC fragment was amplified from pU1111-1 (9) with the primer pair 556/559 (556, CGTGCG AUCTCTCACACAAGGTCAAAAT; 559, CAGCCGAUGCATCTCTGGG TAAAAGCTC) and USER cloned into the AsISI/Nb.BsmI USER cassette in BGHA P147, generating BGHA P123. *mpaDE* was amplified from pJET_mpaDE with the primer pair 570/573 (570, AGAGCGAATGGTCAAAAT; 573, TCTGCGAUGCATTCTGGTTAGG) and USER cloned into the AsISI/Nb.BsmI USER cassette in BGHA P147, generating BGHA P123 and BGHA151.

Genetic transformation and cross. Protoplasting and gene-targeting procedures were performed as described previously (19). A total of 5 µg of BGHA P146, BGHA P151, and BGHA P127 was digested with SwaI to liberate the gene-targeting substrates for bipartite transformation (19). Fragments from BGHA P146 and BGHA P127 or from BGHA P151 and BGHA P146 were used as bipartite transformation substrates for the transformation of *A. nidulans* NID1 (BGHA P146 and BGHA P127), resulting in NID410 or NID211 (BGHA P151 and BGHA P146) resulting in NID944 (see Fig. S1 in the supplemental material). Correct integration was confirmed by using the primer pairs 157/183 (157, TACTCCCCAGCGACTAC; 183, CATTCCGAGATCTCTGAGAC) and 61/156 (61, GGCTACGCTAGCGGATCCACTTAACGTTACTGAA; 156, GTCTCT GACTTCCGGATC). Strain NID211 and NID410 were crossed according to the protocol previously published (28), resulting in strain NID416.

UHPLC-HRMS analysis. Three plugs were taken from each strain grown for 8 days at 37°C in three point inoculations on YES media (with supplements, if necessary) and transferred to a 2-ml vial. After the addi-
tion of 1 ml of methanol-dichloromethane-ethyl acetate (1:2:3 [vol/ vol]), the vials were capped and subjected to ultrasonication for 60 min. The supernatants were transferred to clean vials, and the organic phase was evaporated under an N₂ flow at 30°C. The residues were redissolved in 200 μl of acetonitrile-water (1:1 [vol/vol]) (with subjection to ultrasonication) and filtered through a 0.22-μm-pore-size PTFE syringe filter.

UHPLC-HRMS was performed on a maxis G3 quadrupole time-of-flight mass spectrometer (Bruker Daltonics, Bremen, Germany) equipped with an electrospray ionization (ESI) ion source. The mass spectrometer was connected to an Ultimate 3000 UHPLC system (Dionex, Sunnyvale, CA) equipped with a diode-array detector scanning 200 to 700 nm. Separation of 1- to 3-μl samples were performed at 40°C on a Kinetex C18 column (100 by 2.1 mm [inner diameter], 2.6 μm; Phenomenex, Torrance, CA) using a linear water-acetonitrile gradient (both buffer with 20 mM formic acid) at a flow of 0.4 ml min⁻¹ starting from 10% acetonitrile and increasing to 100% in 10 min, remaining at 100% acetonitrile for 3 min. Mass spectrometry (MS) analyses were performed in both ESI⁺ and ESI⁻ (separate runs) with a data acquisition range of m/z 100 to 1,000, and the MS was calibrated using sodium formate automatically infused prior to each analytical run, providing a mass accuracy of <1.5 ppm.

Reference standards of nuclear magnetic resonance (NMR)-validated DHMP and 5-MAO (see below) and 3-methylorsellinic acid (Ambinter, Paris, France), as well as orsellenic acid (Apin Chemicals, Oxon, United Kingdom), were coanalyzed. 4,6-Dihydroxy-2-(hydroxymethyl)-3-methylbenzoic acid (DHMB) was tentatively identified in ESI⁻ with a matching accurate mass (<1 ppm accuracy, no other candidate compositions) and isotope pattern (SigmaFit better than 10), loss of CO₂ (diagnostic of a carboxylic acid), and UV absorptions at 210 nm (100%), 261 nm (30%), and 304 nm (25%). The retention time (1.61 min) compared to DHMP (2.78 min) fit well with the calculated LogD values of 0.60 and 2.39, respectively (at a pH of 3.2 as in the solvent) (18). Extracted ion chromatograms of the [M+H]+ ions (± m/z 0.001) for all target peaks were constructed for all extracts to exclude that the compounds were produced in various quantities or present in trace amounts in the medium or the wild-type strains. DHMB was detected as the [M−H]⁻ ions. The target compounds were identified by the whole UV/VIS spectrum, the retention time (±0.02 min), the accurate masses (±1.5 ppm), and the relative intensities of the [M+H]+, [M+Na]+, [M+H-H₂O]+, [M−H]⁻, and [M−H−CO₂]⁻ ions, as well as by their respective isotope patterns.

Structure verification of 5-MAO and DHMP. 5-MAO was isolated from a large-scale ethyl acetate extract prepared from 100 petri dishes with MM/H₁₁₀₀₁ and increased to 100% in 10 min, remaining at 100% acetonitrile for 3 min. Mass spectrometry (MS) analyses were performed in both ESI⁺ and ESI⁻ (separate runs) with a data acquisition range of m/z 100 to 1,000, and the MS was calibrated using sodium formate automatically infused prior to each analytical run, providing a mass accuracy of <1.5 ppm.

Remote homology modeling: mpAD encodes a putative hydroxylase. Initial annotation of the eight proposed ORFs that constitutes the putative MPA biosynthetic cluster in P. brevicompactum suggested the following candidates for the established biosynthetic reactions: (i) polyketide synthase (PKS), mpAC; (ii) cytochrome P450, mpAD; (iii) prenyl transferase, mpAα; (iv) oxidative cleaving, mpAH; (v) and finally O-methyltransferase, mpAG (23). However, the lactonization occurring at the second step in MPA biosynthesis is not a standard cytochrome P450 reaction, which prompted us to investigate this step in detail. In order to pinpoint what type of reaction MpaD catalyzes, we conducted a BLASTP search to identify homologs with known functions. The search identified putative homologs with >50% identity, indicating that orthologs are present in other filamentous fungi (data not shown).

RESULTS AND DISCUSSION

Remote homology modeling: mpAD encodes a putative hydroxylase. Initial annotation of the eight proposed ORFs that constitutes the putative MPA biosynthetic cluster in P. brevicompactum suggested the following candidates for the established biosynthetic reactions: (i) polyketide synthase (PKS), mpAC; (ii) cytochrome P450, mpAD; (iii) prenyl transferase, mpAα; (iv) oxidative...
assigned as CYP631B4, and SNOG_06679 was assigned as CYP631C2 by D. R. Nelson of the P450 Nomenclature Committee (Department of Molecular Sciences, University of Tennessee). The output from HHpred indicated that MpaD is very likely to catalyze a hydroxylation reaction, and we hypothesized that the target is the methyl group in ortho position to the carboxylic acid group on 5-MOA (Fig. 1B). Lactonization then yields DHMP. We considered it unlikely that this reaction is also catalyzed by MpaD and turned our attention toward the remaining unassigned two putative ORFs in the gene cluster, mpB and mpE, annotated as encoding a protein of unknown function and a zinc-dependent hydrolase, respectively. Since the lactone formation from hydroxylated 5-MOA to DHMP requires what resemble a reverse hydrolysis, we turned our attention to mpE.

Remote homology modeling: mpE encodes a putative lactone synthase. MpaE was subjected to bioinformatic analysis as described for MpaD with similar results. The highest-scoring hits from HHpred for MpaE were acyl homoserine lactone (AHL) lactonases from Stenotrophomonas maltophilia (29), Agrobacterium tumefaciens (15), and Bacillus thuringiensis (14). This homology indicates that MpaE has the same structure as enzymes cleaving a lactone bond. Furthermore, multiple sequence alignments confirmed that the signature sequence HXHXDH, which is completely conserved and essential for activity in all Zn-dependent hydrolases, is present in MpaE (Fig. 2). We hypothesized that MpaE catalyzes the reverse reaction, i.e., the formation of a lactone through dehydration and is thereby a lactone synthase. The predicted activity of MpaD and MpaE would in combination result in the conversion of 5-MOA to the second intermediate DHMP in MPA biosynthesis in fungi (3).

**mpaD and mpE is a single gene that encodes a fusion protein.** We decided to undertake a heterologous expression approach to investigate our hypothesis. We have previously used such an approach successfully for expressing the MPA PKS in the MPA-nonproducer fungus *A. nidulans* (9). The putative ORFs of *mpaD* and *mpaE* are located in tandem within the defined MPA cluster and were annotated as separate ORFs (23) using GenScan software (4). GenScan, however, is based on invertebrate sequences and, to be more certain that we would clone the correct and full sequences of the genes, we decided to confirm the proposed annotations with two additional algorithms, FgeneSH (25) and Augustus (27). Both of these programs have been trained on fungal sequences. Surprisingly, predictions from both of these two programs suggested that *mpaD* and *mpaE* are a single gene that encodes a single polypeptide (data not shown), hereafter named *mpadE*. Curiously, the MpaD homologs P450 BM3 and CYP505A1 (12) are natural fusion proteins as well, unlike almost every other known cytochrome P450s. In P450 BM3 and CYP505A1, the CYP domain is fused with an electron-donating domain. To establish whether *mpadE* and *mpaE* are transcribed as two separate or one fused ORF, RNA was extracted from *A. nidulans* strain NID211, NID410, and NID416. (A) Extracted ion chromatogram, m/z 183.06519 ± 0.001 corresponding to the [M+H]+ ion of 5-methylorsellinic acid. The MS spectrum at 4.0 min is inserted in relevant chromatograms. (B) Extracted ion chromatogram (m/z 181.04954 ± 0.001) corresponding to the [M+H]+ ion of DHMP. The MS spectrum at 2.9 min is inserted in relevant chromatograms. 3-MOA, 3-methylorsellinic acid; 5-MOA, 5-methylorsellinic acid; DHMP, 5,7-dihydroxy-4-methylphthalide.

**MpaDE catalyzes the conversion of 5-MOA to DHMP.** *mpaDE* was introduced into an *A. nidulans* strain (NID211) expressing MPA PKS (*mpaC*) and therefore capable of producing 5-MOA. In contrast to the reference strain NID211, this new strain (NID416) produced a compound eluting as a prominent peak at 2.9 min with the mass expected for DHMP (Fig. 3). The compound was purified and identified as DHMP by NMR (see Fig. S3 and Table S2 in the supplemental material) and by comparison with the published spectra. In addition, we note that, in contrast to NID211, the DHMP producing NID416 contained very little 5-MOA, which also points to that 5-MOA is converted by MpaDE (Fig. 3). We next addressed the question of whether the unique compound produced by NID416 is due to the *mpaDE* gene product converting 5-MOA to DHMP or whether it results from the conversion of an endogenous *A. nidulans* metabolite. We thus expressed *mpaDE* in an *A. nidulans* strain not producing 5-MOA. The resulting strain (NID410) did not produce detectable levels of DHMP. Finally, we constructed a strain NID944 where the *mpaD* part of *mpaDE* is expressed in a strain expressing *mpaC*. We found that strain NID944 contains a high amount of DHMB, whereas this metabolite was not detected in the other strains. This result conclusively shows that MpaD catalyzes the conversion of 5-MOA to DHMB. Interestingly, NID944 does produce DHMP, which shows that the conversion from DHMB can happen nonenzymatically or that *A. nidulans* has an endogenous enzyme that can
catalyze the lactonization. AN0028 and AN2639 do both have high structural similarity to MpaE, and in HHpred the top hits are the same as for MpaE (data not shown). Therefore, they are good candidates for catalyzing the lactone formation when a high concentration of DHMB is available. Taken together, these results show that the natural fusion protein MpaDE catalyzes the formation of DHMP in MPA biosynthesis, and the buildup of DHMB in NID944 supports that MpaD catalyzes the hydroxylation (Fig. 1B and see Fig. S5 in the supplemental material). The results also suggest that the lactonization step is potentially aided by MpaE.

MpaD and MpaE orthologs involved in lactonization in other fungi? To investigate whether the fusion of MpaDE is widespread in nature, we performed a BLASTP search. This search did not identify any orthologs in any of the organisms within the NCBI database. However, as previously mentioned, BLASTP analysis identified several orthologs of MpaD and MpaE as separate enzymes in a number of fungi. Interestingly, we noticed that in both *Talaromyces stipitatus* and *Phaeosphaeria nodorum*, orthologs of MpaD (TSTA 060710 and SNOG 06679) and MpaE (TSTA 060680 and SNOG 06681) are located very close to each other (Fig. 4). In addition, they are placed in the vicinity of a PKS. The two PKSs have ~40 and ~50% sequence identity, respectively, and the same domain architecture as MpaC (9) and AusA from *A. nidulans* that catalyzes the production of 3,5-dimethylo lactamase-type thioesterase (20). For the remaining members of the *P. brevicompactum* MPA gene cluster, BLAST hits were found for MpaB and MpaG (see Table S1 in the supplemental material), although not in the vicinity of the corresponding PKSs. Based on the homology and genomic proximity, it can be hypothesized that the PKSs from *T. stipitatus* and *P. nodorum* are catalyzing the production of methylsuccinil acid and that the MpaD orthologs are involved in converting this methylorsellinic acid into a lactone. Furthermore, the role of MpaE identified here in the lactonization step of the MPA pathway adds to the list of diverse biochemical functionalities of the polyketide biosynthesis enzymes that belong to metallo-β-lactamase family (2, 13).

In conclusion, our results provide new insights into the biochemical and genetic organization of the MPA biosynthesis pathway. The fusion protein MpaDE may provide an advantage by increasing the rate of reaction due to the close proximity of the two catalytic domains and also by possibly decreasing the side-products from the pathway. As more fungal genomes and polyketide gene clusters get sequenced, it will be of interest to find the extent to which such fusion proteins play a role in shaping the molecular diversity and species-level specificity of fungal polyketides.

ACKNOWLEDGMENTS

We are grateful to David Nelson (http://drnelson.utmem.edu/CytochromeP450.html) for naming MpaD and its orthologs.

The study was supported by grants 09-064967 and 09-064240 from the Danish Council for Independent Research, Technology, and Production Sciences.

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