Cooperation between a Coenzyme A-Independent Stand-Alone Initiation Module and an Iterative Type I Polyketide Synthase during Synthesis of Mycobacterial Phenolic Glycolipids

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Abstract: Several Mycobacterium tuberculosis strains, Mycobacterium leprae, and other mycobacterial pathogens produce a group of small-molecule virulence factors called phenolic glycolipids (PGLs). PGLs play key roles in pathogenicity and host–pathogen interaction. Thus, elucidation of the PGL biosynthetic pathway will not only expand our understanding of natural product biosynthesis, but may also illuminate routes to novel therapeutics to afford alternative lines of defense against mycobacterial infections. In this study, we report an investigation of the enzymatic requirements for the production of long-chain p-hydroxyphenylalkanoate intermediates of PGL biosynthesis. We demonstrate a functional cooperation between a coenzyme A-independent stand-alone didomain initiation module (FadD22) and a 6-domain reducing iterative type I polyketide synthase (Pks15/1) for production of p-hydroxyphenylalkanoate intermediates in vitro and in vivo FadD22-Pks15/1 reconstituted systems. Our results suggest that Pks15/1 is an iterative type I polyketide synthase with a relaxed control of catalytic cycle iterations, a mechanistic property that explains the origin of a characteristic alkyl chain length variability seen in mycobacterial PGLs. The FadD22-Pks15/1 reconstituted systems lay an initial foundation for future efforts to unveil the mechanism of iterative catalysis control by which the structures of the final products of Pks15/1 are defined, and to scrutinize the functional partnerships of the FadD22-Pks15/1 system with downstream enzymes of the PGL biosynthetic pathway.

Introduction

Mycobacterial infections cause serious diseases. In particular, tuberculosis, produced by Mycobacterium tuberculosis, is responsible for nearly two million deaths per year, and leprosy, produced by Mycobacterium leprae, accounts for over three million people with neuropathy-derived disabilities worldwide. Several M. tuberculosis strains (e.g., strains of the W-Beijing family), M. leprae, and other mycobacterial pathogens produce a group of small-molecule virulence factors called phenolic glycolipids (PGLs) (Figure 1).1 PGLs play key roles in pathogenicity and host–pathogen interaction by virtue of their immunomodulatory properties or, in the case of leprosy, their ability to cause peripheral nerve degeneration.2 Moreover, PGL production has been suggested as a trait predisposing M. tuberculosis strains of the W-Beijing family to their characteristic epidemic spread and increased likelihood of developing drug resistance.3 Thus, elucidation of the PGL biosynthetic pathway will not only expand our understanding of natural product biosynthesis, but may also illuminate routes to novel therapeutics to afford alternative lines of defense against mycobacterial infections.4

We have recently demonstrated that the protein FadD22 is required for PGL biosynthesis and reported a FadD22 inhibitor that blocks PGL production.4b FadD22 is comprised of a coenzyme A-independent stand-alone didomain initiation module (FadD22) and a 6-domain reducing iterative type I polyketide synthase (Pks15/1) for production of p-hydroxyphenylalkanoate intermediates in vitro and in vivo FadD22-Pks15/1 reconstituted systems. Our results suggest that Pks15/1 is an iterative type I polyketide synthase with a relaxed control of catalytic cycle iterations, a mechanistic property that explains the origin of a characteristic alkyl chain length variability seen in mycobacterial PGLs. The FadD22-Pks15/1 reconstituted systems lay an initial foundation for future efforts to unveil the mechanism of iterative catalysis control by which the structures of the final products of Pks15/1 are defined, and to scrutinize the functional partnerships of the FadD22-Pks15/1 system with downstream enzymes of the PGL biosynthetic pathway.

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forms a $p$-hydroxybenzoyl-S-FadD22 thioester intermediate from pHBA and ATP in a coenzyme A (CoA)-independent manner.\textsuperscript{4b} We hypothesize that this intermediate primes the biosynthesis of the long-chain $\beta$-diol-containing moiety of PGLs (phenolphthiocerol [PPOL]) by presenting the pHBA starter unit for PPOL biosynthesis.\textsuperscript{1} Subsequently, PPOL is esterified with the FadD22-Pks15/1 system are likely to be further extended by the FadD22-Pks15/1 system.

### Results and Discussion

#### Loading of Starter and Extender Units onto the FadD22-Pks15/1 System.

We first evaluated whether Pks15/1 could be loaded with pHBA in a FadD22-dependent manner in vitro. To this end, incorporation of covalently bound $[^{14}$C]pHBA-derived label onto Pks15/1 and FadD22 was monitored by standard sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) followed by autoradiography (Figure 2).\textsuperscript{8} Phosphopantetheinylation of the ACP domain was not detected in reactions lacking holo-FadD22 (lane 2), containing FadD22(SS76A) in place of holo-FadD22 (lane 7), or lacking ATP (lane 8).

Starter unit loading onto three Pks15/1 site-directed mutants was investigated as well. These mutants had Ala substitutions at the predicted catalytic Cys (Cys-211) of the $\beta$-ketoacyl synthase (KS) domain and/or the phosphopantetheinylation site (Ser-2039) of the acyl carrier protein (ACP) domain. These key amino acid residues were identified by sequence analysis (Figure S1, Supporting Information). Pks15/1(S2039A), the ACP domain mutant defective for carrier domain phosphopantetheinylation (Figure S2, Supporting Information), was radiolabeled like wild-type (cf., lanes 3 and 5). In contrast, holo-Pks15/1(C211A), the KS domain mutant, and Pks15/1(C211A/S2039A), the double mutant, were not competent for label incorporation (lanes 4 and 6, respectively). These observations indicate that a functional KS domain is required for starter unit loading, whereas a functional ACP domain is not. Overall, the results from the starter unit loading analysis are consistent with a model in which the $p$-hydroxybenzoyl unit loaded onto the holo-ArCP domain of FadD22 is transferred to the KS domain active-site Cys-211 residue in Pks15/1. Formation of a $p$-hydroxybenzoyl-Cys-KS domain thioester intermediate would parallel the priming mechanism of other type I PKSs with ordinary acetyl-CoA- or propionyl-CoA-derived starter units.\textsuperscript{9} Pks15/1 was also competent for autoacylation with the predicted extender unit, the malonyl group derived from malonyl-CoA. Covalently bound $[^{14}$C]malonyl-CoA-derived label was readily incorporated onto Pks15/1 (not shown). This labeling is expected to result from AT domain-catalyzed acylation of the ACP domain as seen in other PKSs.\textsuperscript{9}

#### Synthesis of PHPA Products in the in Vitro and in Vivo FadD22-Pks15/1 Reconstituted Systems.

Having demonstrated FadD22-dependent loading of the starter unit onto Pks15/1 and the ability of Pks15/1 for autoacylation with the extender unit, we next evaluated the ability of Pks15/1 to synthesize PHPA products in the in vitro and in vivo FadD22-Pks15/1 reconstituted systems. In these studies, PHPA formation was assessed by liquid chromatography–mass spectrometry (LC–MS) analysis of extracted products released from the enzyme after alkaline hydrolysis of in vitro enzymatic reactions or lysates of engineered E. coli strains expressing the FadD22-Pks15/1 system. Gratifyingly, incubation in vitro of holo-FadD22 and holo-Pks15/1 in the presence of pHBA, malonyl-CoA, ATP, and NADPH led to formation of a characteristic product series. The products in this series had exact masses consistent with predicted PHPAs differing by multiples of the 28-amu unit ($-\text{CH}_2\text{CH}_2-\text{CHO}^{-}$) expected from the presence of different numbers of two-carbon extender units (Figure 3a; Table 1; Figure S3 in Supporting Information).

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(8) See the Supporting Information for additional experimental procedures (plasmid constructions, protein purification, and PHPA standard synthesis), sequence alignment, and Figures S1–S6.
**Scheme 1.** CoA-Independent Loading of the pHBA Starter Unit and Chain Elongation for Biosynthesis of PHPA Intermediates by the FadD22-Pks15/1 System during PGL Assembly

![Scheme Diagram](image)

*Pks15/1* is shown loaded with the malonyl extender unit. The indicated range of iterative cycles and consequent carbon-chain variability in the p-hydroxyphenylalkanoyl-S-Pks15/1 thioester intermediate is that expected during biosynthesis of PGLs from *M. marinum*. Domain abbreviations: A, adenylation; AC, acyl carrier protein; ArCP, aroyl carrier protein; AT, acyltransferase; DH, dehydratase; ER, enoylreductase; KR, keto-reductase; KS, ketosynthase. The thiols of the phosphopantetheinyl group in the carrier domains and the catalytic cysteine in the KS domain are shown.

**Figure 2.** FadD22-dependent loading of pHBA onto Pks15/1. (a) Image of a coomassie blue-stained tri-glycine polyacrylamide gel run under denaturing conditions (7.5%, SDS-PAGE). Lanes 1–9 were loaded with acylation reaction mixtures with the indicated composition of key components. Lane M, molecular weight marker. (b) Autoradiograph image of the gel showing incorporation of [14C]pHBA-derived covalent label onto the proteins.

Information. The retention time and the observed ion [M – H+] m/z of a synthetic 21-(4-hydroxyphenyl)henicosanoic acid PHPA standard (C_{21}H_{40}O_{3}; calculated exact mass: 418.3447; experimental neutral mass 418.3453, where the experimental neutral mass is the observed ion [M – H+] m/z plus H^+), 1.00728 Da) matched the retention time and the ion [M – H+] m/z of the corresponding PHPA product (C_{21}H_{40}O_{3}) (peak 6 in Figure 3; Table 1; Figure S3 in Supporting Information). This PHPA was one of the most abundant in the PHPA product series. The compounds in the PHPA product series also had the same MS/MS fragmentation pattern of the synthetic PHPA standard. This pattern was consistent with loss of H$_2$O (m/z [M – H$^+$ – 18, M – H$^+$ – H$_2$O$^+$]) or CO$_2$ (m/z [M – H$^+$ – 44, M – H$^+$ – CO$_2$]) by the [M – H$^+$] parent ion (Figure S3, Supporting Information).

Conversely, the PHPA product series was not detected in control reactions lacking a single enzyme or the substrates malonyl-CoA or NADPH (not shown). Traces of the most prominent product peaks in the series were seen in controls lacking pHBA or ATP (not shown). This may result from purification steps involving fractions containing traces of the pHBA-adenylation reaction mixtures with the indicated composition of key components. Lane M, molecular weight marker. (b) Autoradiograph image of the gel showing incorporation of [14C]PheBA-derived covalent label onto the proteins.

The results of the PHPA production studies support the idea that the FadD22-Pks15/1 reconstituted systems produce PHPAs with alkyl chain length variability consistent with a 5-to-14 range of iterative extension cycles with the malonyl-CoA-derived extender unit. This suggests that Pks15/1 is a reducing iterative type I PKS with a relaxed control of catalytic cycle iterations. Importantly, such a mechanistic property would explain the origin of the characteristic alkyl chain length variability seen in the PPGO moiety of PGLs. Notably, one of the most prominent PHPA peaks observed in the product series (peak 7, Figure 3) corresponds to 23-(4-hydroxyphenyl)tricosanoic acid (C_{23}H_{44}O_{3}; calculated exact mass, 446.3760; experimental neutral mass in the in vitro and in vivo reconstituted systems, 446.3760 and 446.3763, respectively; Table 1). This PHPA product is predicted to arise from elongation of the starter unit with 11 extender units, and its generation will require a total of 446.3763. Experimental neutral mass in the in vitro and in vivo reconstituted systems, 446.3760 and 446.3763, respectively; Table 1). This PHPA product is predicted to arise from elongation of the starter unit with 11 extender units, and its generation will require a total of 446.3763. Experimental neutral mass in the in vitro and in vivo reconstituted systems, 446.3760 and 446.3763, respectively; Table 1). This PHPA product is predicted to arise from elongation of the starter unit with 11 extender units, and its generation will require a total of 446.3763.
The 5-to-14 range of extension cycles observed in the reconstituted systems is wider than the 9-to-11 range deduced from reported *M. marinum* PPOL structures. In connection with this observation, it is worth noting that, in nonreducing iterative type I PKS systems, a product template (PT) domain and a thioesterase (TE) domain are proposed to be involved in the control of product chain length, cyclization, and release. In contrast, reducing iterative type I PKSs, such as Pks15/1, lack the PT and TE domains, implying a distinct mechanism for the control of the structure of the final products. In fact, Du and co-workers recently suggested that the final size of highly reduced polyketide products of iterative type I PKSs is not determined by the PKSs alone. It is possible that, in the case of Pks15/1, the complete information for final product size is “programmed” via the interaction of Pks15/1 with other enzymes of the PGL pathway. In this regard, our FadD22-Pks15/1 reconstituted systems lay a first foundation for future efforts to unveil the mechanism of iterative catalysis control by which the structures of the final products of Pks15/1 are defined, and to scrutinize the functional partnerships of Pks15/1 with downstream enzymes of the PGL pathway to achieve PPOL biosynthesis.

### Conclusion

The studies reported herein demonstrate the functional partnership between FadD22, a pHBA-specific CoA-independent stand-alone didomain initiation module, and Pks15/1, a 6-domain reducing iterative type I PKS, for production of PHPA intermediates from the PGL biosynthetic pathway. To the best of our knowledge, this is the first demonstration of aroyl-5-ArCP-dependent priming of an iterative PKS. Our results are consistent with a model in which (1) pHBA is transferred from the ArCP domain in *p*-hydroxybenzoyl-FadD22 to the KS domain active-site Cys residue of Pks15/1 and (2) the formed *p*-hydroxybenzoyl-KS transient intermediate serves as a substrate for Pks15/1-dependent acyl chain elongation using the malonyl-CoA-derived extender unit for biosynthesis of PHPA intermediates during PGL production (Scheme 1). The presented data suggest that Pks15/1 has a relaxed control of catalytic cycle iterations, a mechanistic property that explains the origin of a characteristic alkyl chain length variability seen in mycobacterial PGLs. Overall, this study provides novel insight into the biosynthesis of PGLs, an important group of mycobacterial virulence factors.

### Experimental Section

**Reagents.** Unless otherwise stated, all commercially available materials were purchased from Aldrich, Acros Organics, Fisher Scientific, TCI, or Alfa Aesar and were used without any further purification. When necessary, solvents and reagents were dried prior to use, using standard protocols. Radiolabeled [carboxy-14C]p-hydroxybenzoic acid and [2-14C]malonyl-CoA were purchased from American Radiolabeled Chemicals, Inc. 12-N-Biotinyl-(aminododecanoyl)-Coenzyme A (biotinyl-CoA) was purchased from Avanti Polar Lipids, Inc. Monoclonal antibiotin-alkaline phosphatase conjugated antibody was obtained from Sigma-Aldrich. All other molecular biology reagents were from Sigma, Invitrogen, New England Biolabs, Novagen, QIAGEN, or Stratagene. Oligonucleotide primers were purchased from Integrated DNA Technologies, Inc. The synthetic PHPA standard [21-(4-hydroxyphenyl)heneicosanoic acid; C27H46O3] was synthesized as described in the Supporting Information and outlined in Figure S4.
Incorporation of Covalently Bound $^{14}$C$p$HBA-Derived Label onto Pks15/1 and FadD22 in Vitro. For these and all other experiments with FadD22 and Pks15/1 described in this study, we used the proteins from the opportunistic human pathogen *Mycobacterium marinum*. The proteins utilized in the experiments were recombinantly produced and purified as described in the Supporting Information (Figure S5). The starter unit loading experiments were conducted using reported approaches for monitoring $^{14}$C/$^{3}$H acyl-enzyme thioester intermediate formation. The standard reaction mixture (20 µL) contained 12 µM $^{14}$C$p$-hydroxybenzoic acid (pHBA), 100 mM sodium phosphate buffer (pH 7.2), 10% glycerol, 1 mM tris(2-carboxyethyl)phosphine hydrochloride (TCEP), 0.5 mM MgCl$_2$, 1 mM ATP, 4 µM FadD22 (wild-type or mutant), and 8 µM Pks15/1 (wild-type or mutant). In control reactions, selected components were omitted as indicated in Figure 2. After incubation (30 °C, 2 h), the reactions were quenched by the addition of SDS-PAGE loading buffer (100 µL, 2 × concentrated, lacking reducing reagent) and analyzed by standard SDS-PAGE (7.5%). For detection of $^{14}$C-labeled proteins, the polyacrylamide gels were treated with Amplify Fluorographic Reagent (GE Healthcare) as recommended in the manufacturer’s manual. The treated gels were analyzed by standard SDS-PAGE (7.5%).

Table 1. Mass Spectrometry Data for $p$-Hydroxyphenylalkanoate Products

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Synthetic PHPA Standard

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$^a$ Peak numbers correspond to those shown in Figure 3. $^b$ The calculated exact masses for the molecular formulas shown were determined using Agilent’s MassHunter Qualitative Analysis Software package version B.02.00 (Agilent Technologies, Inc.). $^c$ The experimental exact masses were determined by processing the data using Agilent’s MassHunter Qualitative Analysis Software’s “Find Compounds by Formula” feature. The software searches the data for ions based on a given molecular formula and, in this case, extracts the ions for the [M – H$^-$]$^+$ ion for the molecular formulas of interest. The experimental exact mass is calculated by the addition of a proton (H$^+$; 1.00728 Da) to the measured m/z value. The experimental exact mass is used to calculate the ppm difference from the calculated exact mass for a given molecular formula. $^d$ The ppm error was determined by Agilent’s MassHunter Qualitative Analysis Software.
polyacrylamide gels were exposed to storage phosphor screens (GE Healthcare), which were subsequently scanned using a Typhoon Trio Imager (Amersham Biosciences).

**Incorporation of Covalently Bound [14C]Malonyl-CoA-Derived Label onto Pks15/1.** The extender unit loading experiments were conducted using reported approaches for monitoring [14C]labeled acyl-enzyme thioester intermediate formation.46,15 The proteolytic enzymes utilized were recombinantly produced and purified as described in the Supporting Information. The standard reaction mixture (20 µL) contained 50 mM [14C]malonyl-CoA, 100 mM sodium phosphate buffer (pH 7.2), 10% glycerol, 1 mM TCEP, 0.5 mM MgCl2, 1 mM ATP, and 8 µM Pks15/1 (wild-type or mutant). After incubation (30 °C, 3 min), the reactions were quenched by the addition of SDS-PAGE loading buffer14 (7 µL, 3x concentrated, lacking reducing reagent) and analyzed by standard SDS-PAGE (7.5%).16 For detection of [14C]-labeled proteins, the polyacrylamide gels were treated and exposed to storage phosphor screens as described in the preceding paragraph. The screens were then scanned using a Typhoon Trio Imager.

**Protein Phosphopantetheinylation Analysis.** Phosphopantetheinylation was assessed as incorporation of the biotinylated phosphopantetheinyl group derived from the CoA analog biotinyl-CoA (Avanti Polar Lipids) onto the proteins. The promiscuous Bacillus subtilis phosphopantetheinyl transferase Sfp11 was utilized to phosphopantetheinylate Pks15/1 proteins in vitro. Sfp was recombinantly produced and purified as previously reported.11 The Pks15/1 proteins were produced recombinantly and purified as described in the Supporting Information. The standard reaction mixture (20 µL) contained 4 µM Pks15/1 (wild-type or mutant), 100 mM sodium phosphate buffer (pH 7.2), 10% glycerol, 2 mM TCEP, 0.5 mM MgCl2, 50 nM Sfp, and 50 µM biotinyl-CoA. After incubation (30 °C, 15 min), the reactions were quenched by addition of SDS-PAGE loading buffer14 (7 µL, 3x concentrated, lacking reducing reagent). The samples were analyzed by standard SDS-PAGE (7.5%).16 For detection of [14C]labeled proteins, the polyacrylamide gels were treated and exposed to storage phosphor screens as described in the preceding paragraph. The screens were then scanned using a Typhoon Trio Imager.

**LC–MS Instrumentation and Analysis.** Mass spectral data were collected at the City University of New York (CUNY) Mass Spectrometry Facility at Hunter College (New York, NY) on an Agilent Technologies G6520A high-resolution Q-TOF mass spectrometer attached to an Agilent Technologies 1200 Capillary HPLC system. Samples were infused by electrospray ionization in negative mode. Chromatography was performed on a Zorbax 3.1 × 30 mm SB-C18 3.5 µm column (part no. 873700-902) using water containing 0.1% formic acid and 50 µM ammonium formate (solvent A) and 95% acetonitrile containing 0.1% formic acid and 50 µM ammonium formate (solvent B) at a flow rate of 350 µL/min. The gradient program was as follows: 2% B (0–3 min), 2–100% B (3–15 min), 100% B (15–30 min). Total analysis time was 30 min. The HPLC flow was diverted to waste for the first 2.5 min. The temperature of the column was held at 45 °C. The entire instrument parameters were as follows: fragmentor = 140 V; drying gas temperature = 300 °C; drying gas flow = 12 L/min; nebulizer pressure = 40 psi; and capillary voltage = 3500 V. Data were collected with the instrument set to low mass range (100–1700 m/z) under high-resolution conditions at 4 GHz, and data were stored as both centroid and profile mode. The mass spectra were collected over a range of 115–1600 m/z at 1 spectra/s, and MS/MS spectra were collected over a range of 60–800 m/z at 1 spectra/s with an isolation width of ∼4 m/z. The collision energy was ramped using a slope of 6.5 V/100 Da and an offset of 5 V using nitrogen as the collision gas. The reference masses used were purine with (M + H)+ at 119.03632 m/z and HP-922 with (M + formate)− ion at 966.00072 m/z. They were infused into the spray chamber using Agilent’s calibrant delivery system. The instrument was

controlled with Agilent MassHunter Workstation Acquisition Software, and data were analyzed using Agilent MassHunter Workstation Qualitative Analysis Software (Agilent Technologies, Santa Clara, CA). Funding for this instrument system was provided by NIH Shared Instrumentation Grant 1S10RR022649-01 and the CUNY Instrumentation Fund.

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Note Added after ASAP Publication. Two names were added to the list of authors of this publication on October 30, 2009.

Supporting Information Available: Experimental procedures for plasmid constructions, protein purification, and PHPA standard synthesis; sequence alignment; and Figures S1–S6. This material is available free of charge via the Internet at http://pubs.acs.org.

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