

FT-ICR-MS Characterization of Intermediates in the Biosynthesis of the α -Methylbutyrate Side Chain of Lovastatin by the 277 kDa Polyketide Synthase LovF[†]

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ABSTRACT: There are very few fungal polyketide synthases that have been characterized by mass spectrometry. In this paper we describe the *in vitro* reconstitution and FT-ICR-MS verification of the full activity of an intact 277 kDa fungal polyketide synthase LovF of the lovastatin biosynthetic pathway. We report here both the verification of the reconstitution of fully functional *holo*-LovF by using ¹³C-labeled malonyl-CoA to form α -methylbutyrate functionality and also detection of five predicted intermediates covalently bound to the 4'-phosphopantetheine at the acyl carrier protein (ACP) active site utilizing the phosphopantetheine ejection assay and high-resolution mass spectrometry. Under *in vitro* conditions, the diketide acetoacetyl intermediate did not accumulate on the ACP active site of *holo*-LovF following incubation with malonyl-CoA substrate. We found that incubation of *holo*-LovF with acetoacetyl-CoA served as an effective means of loading the diketide intermediate onto the ACP active site of LovF. Our results demonstrate that subsequent α -methylation of the acetoacetyl intermediate stabilizes the intermediate onto the ACP active site and facilitates the formation and mass spectrometric detection of additional intermediates en route to the formation of α -methylbutyrate.

Polyketide natural products have served as a valuable source of useful therapeutics, due in large part to their broad structural variability and resulting functional diversity (1). Numerous compounds of polyketide origin, such as erythromycin (antibiotic) (2), rapamycin (immunosuppressant) (3), doxorubicin (antitumor) (4), and lovastatin (antihypercholesterolemic) (5), are commercially available drugs and widely used. The profound impact of this class of natural products on pharmaceutical development has led to extensive efforts to characterize the enzymes responsible for their biosynthesis. The enzymes, polyketide synthases (PKSs),¹ follow a general biosynthetic scheme in which simple acyl-coenzyme A (CoA) substrates are utilized to generate larger, more complex polyketide products by an array of catalytic domains (6). Polyketide biosynthesis progresses in a stepwise manner in which multiple acyl substrates are selected by the enzyme, and through repeated cycles of Claisen-like condensation reactions, the individual building blocks are linked together

via C–C bonds to form an elongated intermediate. Each round (iteration) of condensation may be followed by various combinations of tailoring reactions; each reaction occurs at the active sites of the various catalytic domains. The variation in structure of each polyketide is ultimately derived from the combination of different catalytic domains that make up the PKS and the acyl-substrate specificity of the acyl transferase (AT) domain(s).

Bacterial type I PKSs are composed of several modules, each of which contains a minimal set of catalytic domains and an array of additional tailoring domains that modify the polyketide intermediate after each iteration of elongation. Each module has, at minimum, a ketosynthase (KS) domain that catalyzes the Claisen-like condensation between the growing intermediate and “extender” acyl units, an AT domain that selects and loads the extender acyl unit from the CoA substrate (the initial module selects/loads the “starter” acyl subunit), and an acyl-carrier protein (ACP) domain that covalently tethers the growing intermediate to the enzyme and transfers it from one module to the next downstream module. The conserved serine at the active site of the ACP domain is posttranslationally modified with a phosphopantetheine (PPant) moiety (7). PPant acts as a flexible/mobile arm, bearing a thiol group to which the growing polyketide intermediate remains covalently bound as a thioester, thereby facilitating the movement of the intermediate to the active sites of the various domains and between each module (8).

In contrast to bacterial type I PKSs, in which each domain is used only once per every molecule of product synthesized, the catalytic domains of fungal type I PKSs are reused multiple times as successive rounds (iterations) of elongation and tailoring occur to yield a polyketide product. Essentially, a fungal type I PKS resembles a single bacterial type I PKS module in terms of

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Abbreviations: ACP, acyl carrier protein; Acac-CoA, acetoacetyl-coenzyme A; CID, collision-induced dissociation; CoA, coenzyme A; DH, dehydratase; DKS, diketide synthase; ER, enoyl reductase; FT-ICR-MS, Fourier transform ion cyclotron resonance mass spectrometry; HMG-CoA, 3-hydroxy-3-methylglutaryl-coenzyme A; IRMPD, infrared multiphoton dissociation; KR, ketoreductase; KS, ketosynthase; MAT, malonyl-CoA:ACP acyltransferase; MS, mass spectrometry; MSⁿ, tandem MS (*n* = number of stages of fragmentation); MT, methyltransferase; NADPH, nicotinamide adenine dinucleotide phosphate; NKS, nonaketide synthase; NRPS, nonribosomal peptide synthetase; Pant, pantetheine; PKS, polyketide synthase; PPant, phosphopantetheine; SAM, S-adenosylmethionine.

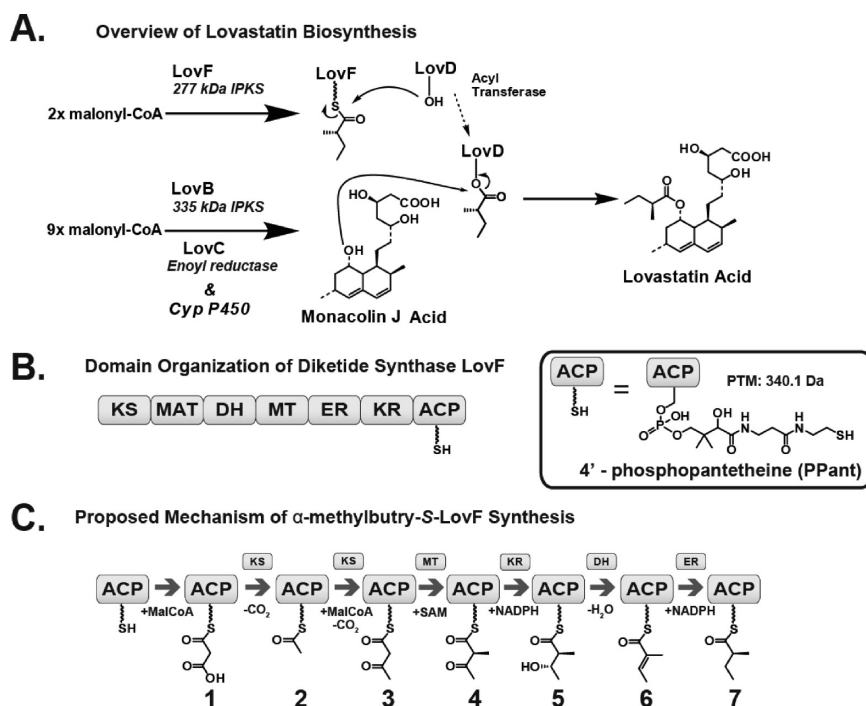


FIGURE 1: Biosynthesis of lovastatin. (A) Overview of the biosynthesis of lovastatin by LovB and LovF. LovD, a broadly acting acyl transferase, mediates the transfer of α -methylbutyrate from LovF to the C8 hydroxyl position of monacolin J acid to form lovastatin acid. (B) LovF consists of multiple catalytic domains: β -ketoacyl synthase (KS), malonyl-CoA:ACP acyltransferase (MAT), dehydratase (DH), methyltransferase (MT), enoylreductase (ER), ketoreductase (KR), and acyl carrier protein (ACP). The 4'-phosphopantetheinylated ACP serves as a covalent tether for the acyl intermediate as it is processed by the other catalytic domains of LovF. (C) Based upon the catalytic domain composition of LovF, the enzyme is predicted to produce multiple covalent intermediates before biosynthesis of α -methylbutyrate.

domain composition. However, fungal PKSs possess a smaller number of catalytic tailoring domains but may use them iteratively in different combinations to generate a polyketide. A subgroup of fungal PKSs of particular interest is highly reducing polyketide synthases (HR-PKSs) from filamentous fungi (9). HR-PKSs are capable of varying combinations of β -keto reduction, dehydration, and/or enoyl reduction after each iteration of elongation, but little is known about how these enzymes are programmed to do so.

The polyketide lovastatin is a potent HMG-CoA inhibitor (statin) (10–12) and was the first compound approved by the FDA to be used specifically for lowering serum cholesterol levels. Lovastatin was widely used and is the progenitor to modern statin analogues, such as the blockbuster drug simvastatin (13). The biosynthesis of lovastatin is accomplished in the filamentous fungus *Aspergillus terreus* by two HR-PKSs: the 335 kDa nonaketide synthase (NKS) LovB and the 277 kDa diketide synthase (DKS) LovF (14–16). LovB, in conjunction with a separate enoyl reductase LovC, synthesizes dihydromonacolin L acid (17, 18), which is subsequently oxidized by CYP450 enzymes to yield monacolin J acid (Figure 1A), the nonaketide core of lovastatin. LovF synthesizes the α -methylbutyrate side chain that is transferred by the dissociated acyltransferase LovD (19) to the C8-hydroxyl of monacolin J acid to yield the complete natural product. Acetyl-CoA, malonyl-CoA, NADPH, and SAM are utilized by LovF to yield α -methylbutyryl-S-LovF, a process involving a single condensation reaction followed by α -methylation, β -ketoreduction, dehydration, and α,β -enoyl reduction reactions at the active sites of the various catalytic domains (Figure 1B). The starting unit acetate and extender unit of malonate, originating from acetyl-CoA and malonyl-CoA, respectively, are received by the malonyl-CoA:ACP acyltransferase (MAT) domain and transferred to the free thiol of the PPant arm at the ACP active

site. Under *in vitro* conditions, malonyl-CoA is the source of both the starter and extender units; therefore, the formation of the malonyl-S-ACP intermediate is followed by decarboxylation to acetyl-S-ACP and the acetyl intermediate is transferred to the KS domain, leaving the ACP unoccupied and available to receive the malonyl extender unit. Subsequent condensation occurs between the acetyl intermediate residing at the KS active site and the decarboxylated malonate extender unit. The resulting acetoacetyl-S-ACP undergoes MT-catalyzed C-methylation of the α -carbon. The methylated diketide undergoes β -ketoreduction, dehydration (forms an α,β -double bond), and enoyl reduction in the presence of NADPH. These reactions are catalyzed by the KR, DH, and ER domains, respectively (Figure 1C).

Protein mass spectrometry (MS) has emerged as a valuable tool for the investigation of the biosynthesis done by both PKSs and the related nonribosomal peptide synthetases (NRPSs). The value of mass spectrometry as a tool for dissecting these biosynthetic pathways can be largely attributed to the fact that intermediates are covalently bound to active site residues, thus causing stable mass shifts (20–23). Additionally, the phosphopantetheinyl arm present on carrier domains of these enzymes is prone to be ejected when the protein (or protease-digested peptide) bearing PPant is subjected to thermal activation by collision-induced dissociation (CID) or infrared multiphoton dissociation (IRMPD) (24–26). The resulting ejected PPant arm gives rise to two characteristic 1+ ions at m/z 261.1267 and 359.1036 that will be shifted corresponding to the mass of a covalently bound intermediate. Taking advantage of the aforementioned features of PKSs, and coupled with the use of high-resolution/high-mass-accuracy instruments, an unparalleled level of analysis of these massive enzymatic machines can be achieved.

To date, there are only a handful of studies which utilize mass spectrometry to analyze iterative fungal PKSs, largely due to the

Table 1: HPLC Gradient for Active Site Fractionation - Solvent A is Water (w/ 0.1% TFA) and Solvent B is Acetonitrile (w/ 0.1%TFA)

min	0.0	10.0	15.0	55.0	60.0	60.1	60.2	62.6	63.0	65.0	66.0	71.0
% A	90	90	70	30	10	10	95	95	5	5	90	90
% B	10	10	30	70	90	90	5	5	95	95	10	10

difficulty in obtaining sufficient quantities of these enzymes. Recent efforts to engineer a yeast expression system from which to obtain large quantities of fully functional *holo*-LovF have been successful (27), thereby providing sufficient quantities of the enzyme for thorough MS analysis. Still, many challenges exist which hinder the analysis of relatively small intermediates (< 600 Da) bound to large, multifunctional PKSs, the most obvious of which is their massive size (> 250 kDa). The best studied fungal pathway is PksA, which assembles the precursor to aflatoxin B1 (28). In a recent investigation, PksA was extensively dissected into discrete domains prior to *in vitro* reconstitution in order for the protein to be more amenable to mass spectrometric investigations.

The aim for this study was to use mass spectrometry to detect the covalent intermediates of α -methylbutyrate biosynthesis formed by structurally intact *holo*-LovF. Our goals were to carry out *in vitro* reactions using the intact megasynthase first, followed by limited proteolysis of the PKS to generate smaller, active site-containing peptide fragments that are more amenable to mass spectrometry analysis. We anticipated that this would provide insight into the active site occupancy of the intermediates during the biosynthesis of α -methylbutyrate of the lovastatin pathway. We report here the detection, by FT-ICR-MS, of multiple transient intermediates bound to the ACP active site-containing peptide fragments from the intact LovF. The use of high-mass-accuracy instrumentation allowed us to directly detect the results of incubation of LovF with its substrates and to determine suitable reaction conditions to generate the various intermediates.

EXPERIMENTAL PROCEDURES

LovF ACP Active Site Identification for MS Analysis. The cloning, expression, and purification of C-terminal hexahistidine-tagged LovF, posttranslationally modified by inclusion of a copy of the *npgA* gene from *Aspergillus nidulans* to yield 4'-phosphopantetheinylated *holo*-LovF, were described previously (27). For the purposes of generating and identifying the LovF ACP active site, as well as all subsequent activity assays, purified *holo*-LovF was diluted to a concentration of 14.4 μ M (4 μ g/ μ L) in 100 mM Tris-HCl buffer, pH 7.8, with 1 mM TCEP in 50 μ L final volume (200 μ g total LovF). LovF was prepared as described above and then incubated with 1 mM malonyl-CoA (Sigma-Aldrich) for 10 min. After acyl-CoA incubation, the sample was digested using 1 μ g of proteomic grade trypsin (1:200 w/w) in 50 μ L of 20 mM NH_4HCO_3 reaction buffer, pH 8.2 (Trypsin Singles; Sigma), at 37 °C. The tryptic digestion was quenched following 10 min using an equal volume (100 μ L) of 10% formic acid. Following proteolysis, the resulting mixtures of tryptic peptides from each *holo*-LovF reaction were fractionated using an Agilent 1200 HPLC equipped with a Jupiter reverse-phase C4 5 μ m, 300A, 150 \times 4.6 mm column (Phenomenex) using the water/acetonitrile gradient listed in Table 1. The absorbance of the digestion mixture was monitored at multiple wavelengths (e.g., 205 and 280 nm) by a photodiode array detector. Fractions containing active site peptides were identified by comparing HPLC chromatographs of *holo*-LovF to those of acyl-CoA

substrate-incubated LovF. All fractions analyzed, including those containing active site peptides, were collected and immediately frozen with dry ice and then lyophilized.

HPLC Gradient for Active Site Purification. Solvent A is HPLC grade water (J. T. Baker) with 0.1% TFA (Sigma-Aldrich), and solvent B is HPLC grade acetonitrile (J. T. Baker) with 0.1% TFA. The gradient was run at a flow rate of 1 mL/min.

Generation of LovF-Bound Acyl Intermediates. For all assays, *holo*-LovF was diluted to 4 μ g/ μ L as described above. Samples of LovF were incubated with either 0.25, 0.5, 1.0, 2.0, 4.0, 8.0, 34.7, or 69.4 mol equiv (3.6, 7.2, 14.4, 28.8, 57.6, and 115.2 μ M, 0.5 mM, 1 mM) of malonyl-CoA with and without 1 mM nicotinamide adenine dinucleotide phosphate (NADPH) and 1 mM *S*-adenosylmethionine (SAM) prior to tryptic digestion. [1,2,3- $^{13}\text{C}_3$]Malonyl-CoA (1 mM) (Sigma-Aldrich) was incubated with the *holo*-LovF, 1 mM SAM, and 1 mM NADPH to generate ^{13}C -labeled intermediates. In order to generate acetoacetyl-*S*-LovF by direct acylation, *holo*-LovF was incubated with 1 mM acetoacetyl-CoA (Sigma-Aldrich) at room temperature. For the detection of additional intermediates, *holo*-LovF was coincubated first with 1 mM acetoacetyl-CoA and 1 mM SAM and then subsequently incubated with 1 mM NADPH.

LovF Mass Spectrometric Analysis. Lyophilized HPLC fractions were dissolved in 40–80 μ L of an ESI solution containing 50% LC-MS grade methanol (J. T. Baker), 49% LC-MS grade water (J. T. Baker), and 1% formic acid (Acros Organics), then gently vortexed for 30 s, and centrifuged for 2 min at 14000 rpm. Analysis was performed using a hybrid 6.4T LTQ-FT (Thermo Electron, North America) mass spectrometer. The instrument first tuned to m/z 816.3 using bovine cytochrome *c* (Sigma-Aldrich). Samples were introduced by electrospray ionization (1.30–1.5 kV, 0.3 psi) using a Biversa Nanomate 100 (Advion Biosciences) nanospray device in positive ion mode. Detection using the low-resolution LTQ was performed using 1000 ms maximum ion accumulation time. An 8000 ms maximum ion accumulation time and a resolution of 100000 were used for FT-ICR-MS detection. MS^2 and MS^3 utilized normalized collision energy of 30%, an activation Q of 0.200, and activation times between 100 and 300 ms depending on the intensity of the ions being isolated. Lowering the activation Q was critical for the detection of low m/z fragment ions, in particular PPant ejection ions, that were less than one-third the mass of the parent peptide. Although multiple charge states of the ACP active site peptides were detected, the highly charged peptides with a mass below m/z 800 were found to have very low signal intensity. In the initial rounds of MS analysis, fractions were quickly surveyed, using the ion trap detector, for the presence of phosphopantetheine ejection ion signature by subjecting individual peptide ions to collision-induced dissociation (CID) and looking for the presence of a pantetheine ejection ion at (calculated mass m/z 261.1) and the phosphopantetheine ejection ion at (calculated mass m/z 359.1), as well as the charge-loss postejction parent peptide. This was accomplished using the data-dependent data acquisition method to quickly survey fractions (see below) prior to more thorough analysis of individual ions by FT-ICR-MS.

For the in-depth FT-ICR-MS analysis of the ACP occupancy, the 17+ charge state of the larger LovF ACP active

site-containing peptide (Y₂₄₂₀–H₂₅₃₈ at m/z 803.9991) was isolated and fragmented in order to detect substrates and/or intermediate occupancy of the ACP active site. Specifically, ions in the range of m/z 800–820 were isolated and fragmented. This mass range contained mostly peptide ions that resulted from the 17+ charge ACP active site peptide or mass-shifted forms, shifts that were due to covalently bound intermediates, oxidation, or noncovalent adducts such as sodium and/or potassium. There were few contaminating ions. All ions between m/z 800 and 820 were collectively trapped and simultaneously fragmented by CID to efficiently generate ejection ions bearing a range of intermediates depending on the reaction conditions used. Following the *in vitro* incubation reactions between *holo*-LovF and the various substrates, mass-shifted ACP peptides bearing covalently bound intermediates still fit well within the m/z 800–820 range. The identity of intermediates of α -methylbutyrate covalently bound to the pantetheine ejection ion was confirmed by both the exact mass as determined by FT-MS² analysis and IT-MS³ detection of diagnostic MS³ fragments. Pantetheine ejection ions subjected to MS³ were fragmented using a normalized collision energy of 30%, an activation Q of 0.200, and activation time of 100 ms at each stage of MS^{*n*}. Typically, a minimum of 250–500 individual FT-ICR-MS spectra were acquired for each at each stage of MS¹, MS², or MS³. The spectra were averaged for post acquisition data analysis using Qualbrowser (Thermo Electron Corp.).

Data-Dependent PPant Ejection Screening Method. In order to rapidly screen mixtures of tryptic peptides in each HPLC fraction for ACP active site-containing peptides, a data-dependent tandem MS method was employed. As each lyophilized HPLC fraction was initially infused into the mass spectrometer using the Nanomate nanospray source, tandem MS data were gathered using a preprogrammed data acquisition method that was generated with the instrument's controlling software: Xcalibur 1.0 (Thermo Electron Corp.). This data acquisition method involved repeatedly acquiring a single low-resolution MS¹ scan followed by data-dependent CID (MS²) of the top five most abundant ions from the MS¹ scan. Each ion was trapped and fragmented twice before being added to an exclusion list. The exclusion list size was set to 500 ions (maximum allowable by the software), with an exclusion duration that lasted the full 5 min duration of the acquisition method. The m/z isolation width was set to ± 1.5 , activation Q at 0.200, the activation time set to 1000 ms, and the activation energy set to 35%. The use of the low-resolution, high-sensitivity LTQ for the data-dependent data acquisition permitted the rapid detection of low-intensity PPant ejection ions that may have been below the detection limits of the FT-ICR-MS (even when employing in-source fragmentation) without incurring significantly longer data acquisition times. As the data-dependent method continued to collect spectra from each fraction, the previously acquired MS² spectra were continuously refreshed and visually screened for the presence of PPant ejection ions. After the conclusion of the data-dependent method, any candidate parent ions that appeared to yield pantetheine ejection ions were subsequently trapped and fragmented. All parent ions identified as being PPant modified were subjected to more thorough FT-ICR-MS analysis.

RESULTS

HPLC-Directed LovF Active Site Detection. To analyze the biosynthetic intermediates covalently bound to LovF by FT-ICR-MS, the entire 277 kDa *holo*-megasynthase was treated to

brief proteolytic digestion following *in vitro* reactions. This approach was employed to yield a complex mixture of peptide (typically 5–35 kDa) fragments, many of which contained multiple missed trypsin cleavage sites. Digested samples of LovF were first subjected to *off-line* fractionation by HPLC prior to MS analysis. In order to rapidly identify the peptide fragments containing active sites, two samples of LovF were prepared: one incubated with malonyl-CoA prior to proteolysis and the other not. Both reaction mixtures were incubated with trypsin and then separated by HPLC using identical gradients. Great care was taken to prepare the two samples identically (incubation time, pH, duration of proteolysis) with the sole difference being the presence of the CoA substrate. It was anticipated that after incubation with malonyl-CoA the active site-containing peptide fragments of LovF would have differences in HPLC retention times on account of the covalently linked substrate, resulting in identifiable peak shifts on the chromatograms when compared to a substrate-free sample. The resulting HPLC chromatograms (205 nm absorbance) of the two digested samples were aligned to reveal separate peaks that appeared shifted in the malonyl-CoA treated sample of LovF (Figure 2). Fractions were collected, flash frozen, and lyophilized, then redissolved in an ESI solution containing 50% MeOH and 1% formic acid, and directly introduced into the mass spectrometer using a Biversa Nanomate nanospray ESI source, a sample introduction system that allows extensive time for multiple methods of data acquisition. Each fraction was subjected to an automated, data-dependent tandem MS method that fragmented the majority of peptide ions, and MS² spectra were quickly inspected, using just the linear ion trap detector, for the presence a characteristic ejection ions m/z 261.1 or m/z 359.1 which would indicate the presence of the ACP active site.

During the data-dependent spectra acquisition, several different parent peptides produced fragment ions with masses similar to PPant ejection ions. Therefore, to confirm the identity of these fragment ions as being PPant-related, the candidate parent ions were individually retrapped in the ion trap portion of the instrument and subjected to an additional round of fragmentation (MS³) (29). Fragmentation of the PPant ejection ions resulted in a series of diagnostic MS³ fragments. Parent peptide ions that were determined to yield PPant ejection ions were reanalyzed by FT-ICR-MS in order to unambiguously determine the exact mass of the peptide as well as confirming the exact mass of the PPant ejection ion. Using this overall data acquisition strategy, it was possible to map these parent peptides to the primary sequence of *holo*-LovF.

LovF ACP Active Site Detection by FT-ICR-MS. When comparing the chromatograms of the trypsin-digested LovF and that of LovF incubated with malonyl-CoA prior to digestion, two distinct differences were noted (Figure 2, regions A and B) between 51.25% and 52.00% acetonitrile (36.25 to 37.00 min during the gradient) and 52.75% to 53.25% acetonitrile (37.75 to 38.25 min). FT-ICR-MS analysis of the fraction collected at point A revealed the presence of multiple charge states (11+ through 20+ observed) of a single peptide with a monoisotopic mass of 13650.86 Da (Figure 3A), within 0.9 ppm of the calculated exact mass (13650.85 Da) of the *holo*-ACP active site-containing tryptic peptide Y₂₄₂₀–H₂₅₃₈ of LovF (Figure 3B). The HPLC fraction collected at point B was also found to contain multiple charge states (7+ through 15+) of a peptide with a monoisotopic mass of 10440.26 Da (Figure 3C), which was within 1.4 ppm of the theoretical mass (10440.25 Da) of the *holo*-ACP active site-containing tryptic peptide A₂₄₄₈–H₂₅₃₈ of LovF

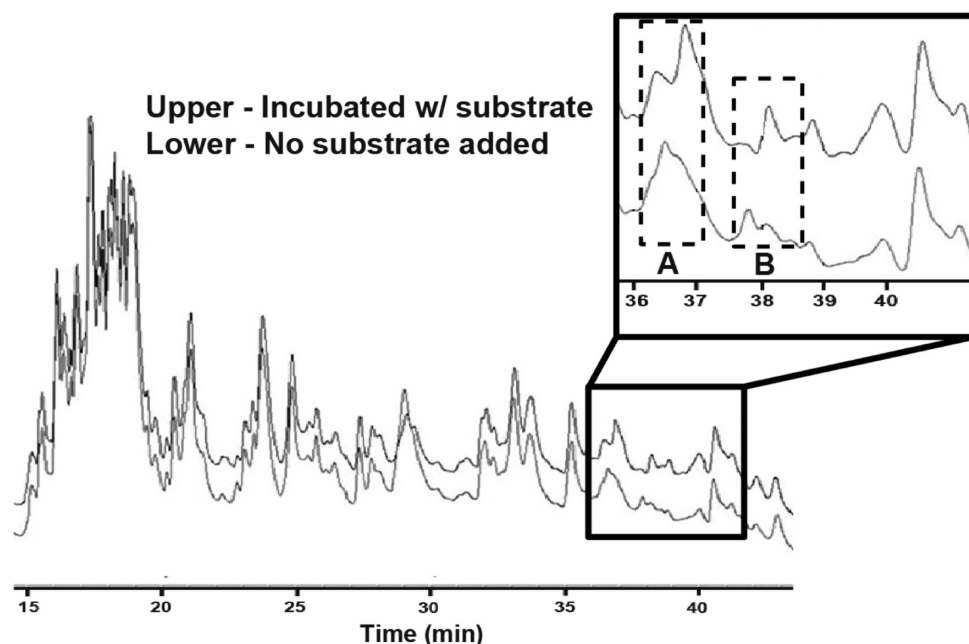


FIGURE 2: HPLC chromatogram comparison of LovF digests indicates active sites. Samples of *holo*-LovF and *holo*-LovF incubated with 1 mM malonyl-CoA were each subjected to limited proteolysis by trypsin and separated by HPLC. Comparing the chromatograms (absorbance 205 nm) of these separate samples of LovF revealed two regions (A and B) where peak shifts occurred due to the presence of substrate covalently bound to active site-containing tryptic peptides. Points A and B correlate to ACP active site-containing peptides of two different sizes.

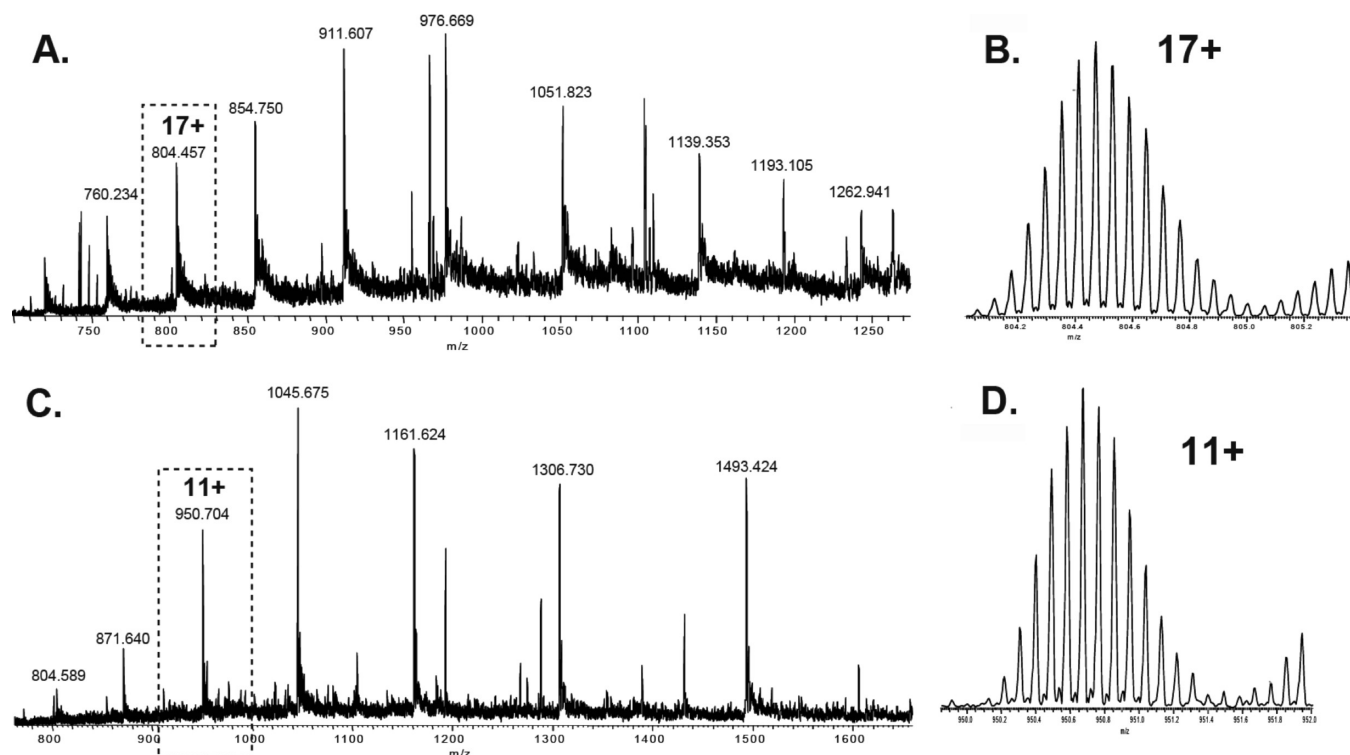


FIGURE 3: FT-ICR-MS of *holo*-LovF ACP active site-containing peptides. Analysis of individual HPLC fractions by FT-ICR-MS revealed the presence of two different ACP active site-containing peptides in two separate HPLC fractions. Multiple charge states of each tryptic peptide were detected. (A) FT-ICR-MS of the HPLC fraction collected between 36 and 37 min resulted in detection of multiple charges of the ACP peptide $Y_{2420}-H_{2538}$. (B) Detail of the 17+ charge state of the ACP active site peptide $Y_{2420}-H_{2538}$. (C) FT-ICR-MS of the HPLC fraction collected between 37 and 38 min revealed multiple charges of the ACP peptide $A_{2448}-H_{2538}$. (D) Detail of the 11+ charge state of the ACP active site peptide $A_{2448}-H_{2538}$.

(Figure 3D). Subjecting these ACP active site peptides to CID activation yielded phosphopantetheine ejection ions at m/z 261.1267 and m/z 359.1036 (Figure 4), serving as additional confirmation that these two tryptic peptides contained the phosphopantetheinylated active site residue (S_{2480}). These two ACP peptides

were consistently generated, in approximately equal quantities, following subsequent limited tryptic digestions of *holo*-LovF. All HPLC fractions were also inspected for peptides containing the KS and MAT active sites, but these specific peptides were not identified following limited proteolysis.

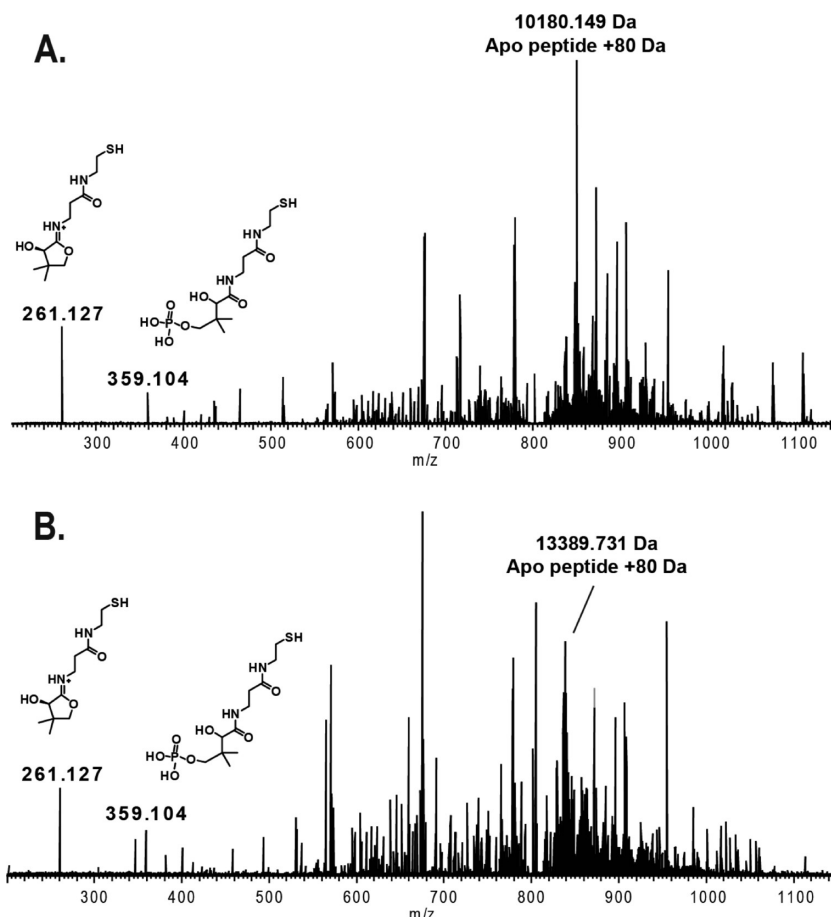


FIGURE 4: Phosphopantetheine elimination from *holo*-LovF ACP active site peptides. FT-ICR-MS² of the two different sizes of ACP active site-containing tryptic peptides from a sample of *holo*-LovF not incubated with malonyl-CoA revealed the presence of both a pantetheine ejection ion (Pant) at m/z 261.127 and a phosphopantetheinyl (PPant) ejection ion at m/z 359.104. The resulting charge loss parent phosphopeptide was clearly detectable among the fragment ions. (A) MS² spectrum of the ACP active A₂₄₄₈–H₂₅₃₈. (B) MS² spectrum of the ACP active Y₂₄₂₀–H₂₅₃₈.

LovF and Malonyl-CoA Incubation. Having the ability to reproducibly generate ACP active site peptides from the intact *holo*-LovF PKS, we turned our attention toward monitoring this active site's occupancy following incubation of the megasynthase with substrate. We anticipated that incubation with malonyl-CoA would result in the formation of acetoacetyl-S-ACP intermediate. To this end, *holo*-LovF was initially incubated with only malonyl-CoA, digested, and then subjected to MS analysis. When analyzing the fraction that contained the ACP peptides, a wide range (e.g., m/z 800–820) of ions were trapped, such that only a single charge state (e.g., 17+) of the ACP active site peptide was isolated and fragmented. The use of this wide isolation window permitted the detection of mass additions to both the intact peptide and to the resulting PPant ejection ions. This allowed the determination of the relative quantities of different acyl intermediates bound to the ACP active site at the time of proteolysis. Following incubation with malonyl-CoA, both of the *holo*-LovF ACP active site-containing peptides had mass additions of 86.00 Da, corresponding to the formation of malonyl-S-LovF. Upon CID, these peptides yielded a pantetheine ejection ion with a mass of m/z 347.128 (calculated 347.1271 Da, 1.8 ppm) (Figure 5). The identities of the pantetheine ejection ions were further verified by MS³ and analysis of the resulting fragmentation pattern of the pantetheine ejection ion (Supporting Information Figure 1). In addition, acetyl-S-ACP was detected in the same spectra as malonyl-S-ACP following incubation of *holo*-LovF with malonyl-CoA (Figure 5). The acetyl intermediate at

the ACP active site was the result of decarboxylation of the malonyl-S-ACP.

Most surprising was our inability to detect the acetoacetyl intermediate, which was expected to form in the presence of *holo*-LovF and malonyl-CoA, a result of decarboxylation and condensation between a malonyl extender unit and the acetate that resulted from decarboxylation of the initial malonyl unit loaded to the ACP (30). Initially, *holo*-LovF was incubated with 1 mM malonyl-CoA for 10 min prior to proteolysis. Different incubation durations (e.g., 1, 2, 4, 8, 15, 30 min) were tested but did not yield different results. Inclusion of SAM and NADPH during incubation also failed to generate the acetoacetyl-S-ACP. Therefore, we hypothesized that the 1 mM malonyl-CoA concentration (~70 mol equiv relative to *holo*-LovF used) was excessively high and was overwhelming the ability of *holo*-LovF to shuttle the diketide from the KS active site back to the ACP active site by *congesting* the ACP with malonyl. To test this hypothesis, *holo*-LovF was incubated with lower concentrations of malonyl-CoA such that 0.5, 1, 2, 4, and 8 mol equiv of substrate were used, both with and without 1 mM SAM and 1 mM NADPH. Given that 2 mol of malonyl-CoA and 1 mol *holo*-LovF are needed to generate 1 mol of acetoacetyl-S-LovF, we anticipated that the lowered malonyl-CoA concentration in the incubations would yield the sought-after intermediate(s). However, the lower substrate concentrations also failed to yield detectable quantities of the acetoacetyl-S-ACP peptide (Figure 6), producing mostly malonyl-S-ACP and, to a lesser extent, acetyl-S-ACP. At this point

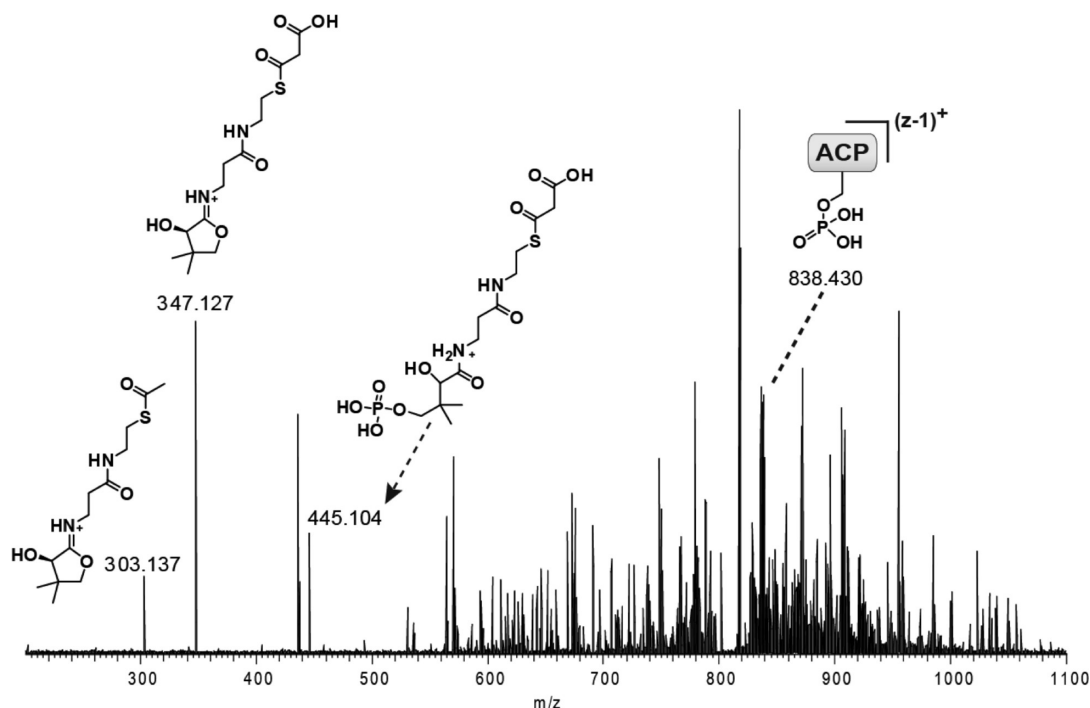


FIGURE 5: Detection of malonyl-S-LovF by PPant ejection. The FT-ICR-MS² of ACP active site-containing peptide following incubation of *holo*-LovF with 1 mM malonyl-CoA. Both pantetheine and phosphopantetheine ejection ion with a mass of 347.127 Da corresponding to the addition of malonyl was detected. The postejection charge loss parent phosphopeptide (MS¹ 17+ to MS² 16+) was detected at m/z 838.430 (13398.90 Da). FT-ICR-MS² of the ACP active site-containing peptide following incubation of *holo*-LovF with low concentrations (<0.1 mM) of malonyl-CoA resulted in the detection of a pantetheine ejection ion at m/z 303.137 corresponding the presence of an acetyl intermediate.

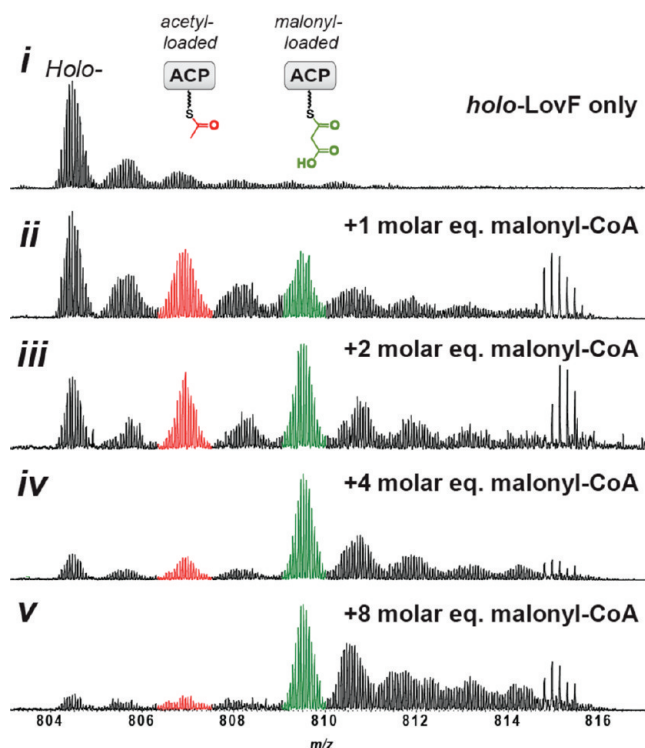


FIGURE 6: Concentration-dependent malonyl-S-LovF accumulation. *Holo*-LovF was incubated with different molar equivalent concentrations of malonyl-CoA. *In vitro* reactions contained 14.4 μ M *holo*-LovF incubated with SAM, NADPH, and 0, 14.4, 28.8, 57.6, or 115.2 μ M malonyl-CoA. The ACP active site peptide was predominately occupied with malonyl after incubation with 115.2 μ M malonyl-CoA. The relative intensity of acetyl-S-LovF was substantially higher at lower concentrations of malonyl-CoA.

in time the occupancy of the LovF KS active site cannot be determined.

During these sets of experiments, scaling up the quantity of enzyme being used led us to detect, by FT-ICR-MS, that a small quantity (<5%) of α -methylbutyryl-S-LovF was detectable after incubation with 0.5 equiv of malonyl-CoA, as previously reported (27). PPant ejection and MS³ were used to verify this assignment. However, the α -methylbutyryl product was found to be present even in the absence of malonyl-CoA incubation (Supporting Information Figure 2). Therefore, we were faced with two unanticipated obstacles in our attempt to detect multiple LovF intermediates. First was the lack of detectable acet-oacetyl-S-ACP, and second was the presence of a small amount of α -methylbutyryl-S-ACP regardless of whether or not substrate was included. In light of these two observations, it was unclear whether LovF was capable of production of α -methylbutyrate in the absence of LovD under these particular *in vitro* reaction conditions. We address both of the aforementioned obstacles in the next section.

LovF Incubation with ¹³C-Labeled Malonyl-CoA Substrate. In order to determine whether a portion of the α -methylbutyryl-S-LovF observed, following incubation of *holo*-LovF with substrate, was due to *de novo* synthesis under *in vitro* reaction conditions, it was necessary to further evaluate our assay using ¹³C-labeled substrate. Samples of *holo*-LovF were incubated with 1 mM each [1,2,3-¹³C₃]malonyl-CoA, SAM, and NADPH for 10 min prior to trypsin digestion. Upon CID fragmentation of the ACP active site peptide, [1,2,3,4-¹³C₄]- α -methylbutyryl-loaded PPant ejection ion (m/z 349.198) was clearly detectable in the resulting tandem mass spectrum, along with the unlabeled α -methylbutyrate product that copurified with *holo*-LovF (Figure 7). These results demonstrated that under our *in vitro* reaction

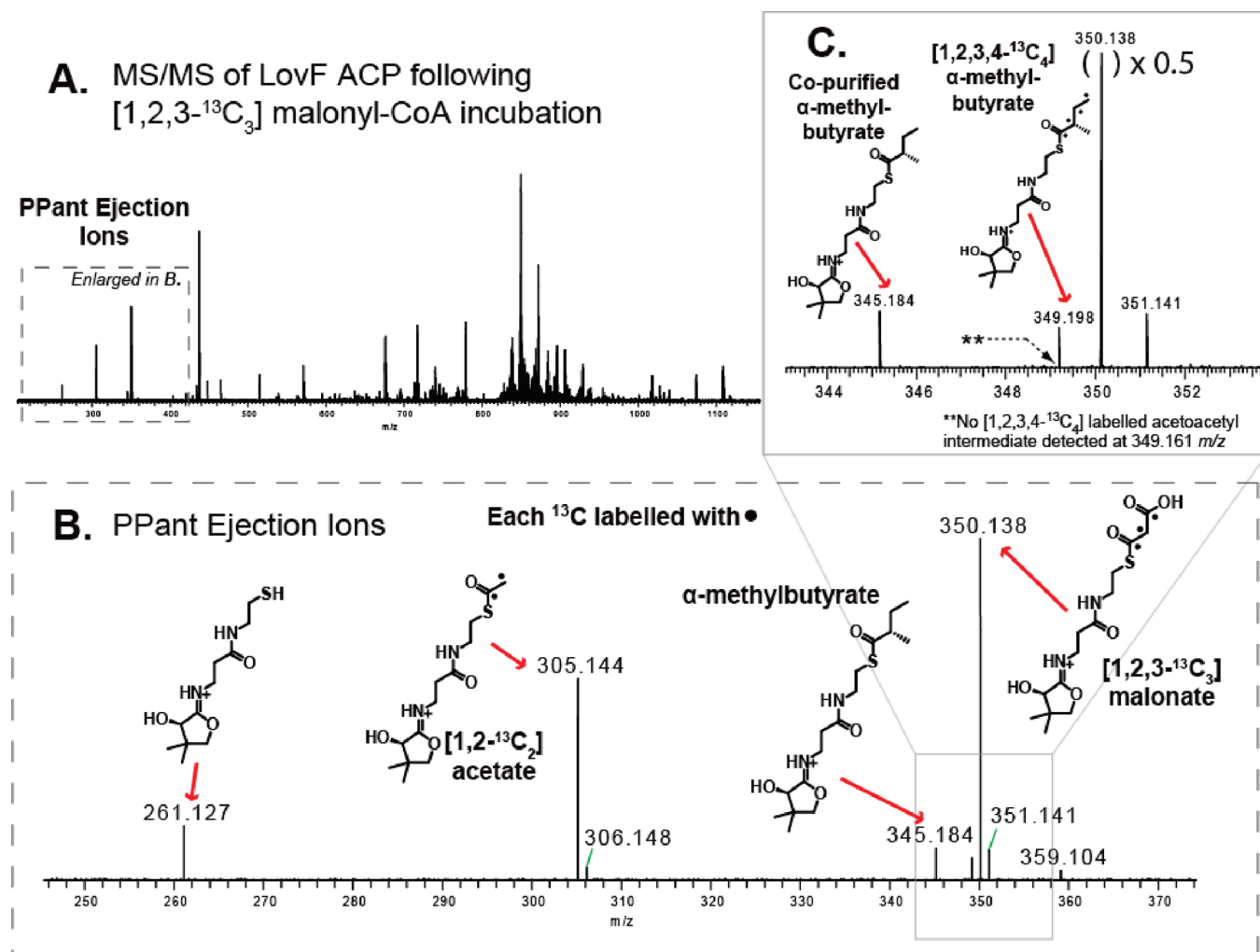


FIGURE 7: [1,2,3,4-¹³C₄]-α-Methylbutyrate product detection. To confirm the *de novo* biosynthesis of α-methylbutyrate, *holo*-LovF was incubated with [1,2,3-¹³C₃]-malonyl-CoA and analyzed by FT-ICR-MS. (A) FT-ICR-MS² of all parent ions between *m/z* 800 and 820 resulted in detection of multiple PPant ejection ions. (B) A more detailed view reveals the presence of [1,2-¹³C₂]acetyl-loaded and [1,2,3-¹³C₃]malonyl-loaded PPant ejection ions. (C) The unlabeled α-methylbutyryl-loaded PPant ejection ion was present. However, the [1,2,3,4-¹³C₄]-α-methylbutyryl-loaded PPant ejection ion was detected in very low quantity, validating that the *holo*-LovF used in these assays was fully competent and capable of generating the product from malonyl-CoA added to the *in vitro* reactions.

conditions *holo*-LovF was a fully competent enzyme capable of *de novo* biosynthesis of [1,2,3,4-¹³C₄]-α-methylbutyrate using [1,2,3-¹³C₃]-malonyl-CoA. Just as when *holo*-LovF was incubated with unlabeled malonyl-CoA and there was no detectable formation of acetoacetate-S-LovF, we did not detect the presence of a [1,2,3,4-¹³C₄]acetoacetyl-loaded PPant ejection ion (calculated: *m/z* 349.161) which would have been distinguishable by FT-ICR-MS from the [1,2,3,4-¹³C₄]-α-methylbutyryl-loaded PPant ejection ion.

Direct Acylation with Acetoacetyl-CoA. Because we did not detect the acetoacetyl intermediate on the ACP active site, we sought alternate solutions for evaluating the function of the individual catalytic domains by detecting the formation of the additional biosynthetic intermediates by LovF en route to α-methylbutyrate. In an attempt to load acetoacetate onto the ACP active site, *apo*-LovF was incubated with acetoacetyl-coenzyme A and Sfp (31, 32). At pHs between 7.0 and 8.0, Sfp-mediated phosphopantetheinylation of *apo*-LovF to acetoacetyl-S-ACP failed. The Sfp reaction was repeated in MES buffer at lower pHs (6.0–6.5), but *apo*-LovF began precipitating as the pH was lowered below 7.0. Sfp reactions were typically incubated for 3 h, so an overnight Sfp reaction at pH 7.5 was attempted.

This reaction successfully yielded acetoacetyl-S-LovF, but only an extremely low conversion occurred. Incubation of *apo*-LovF with Sfp and CoA failed to produce *holo*-LovF, whereas a truncated protein comprised of just the *apo*-ACP domain, derived from LovF, was efficiently phosphopantetheinylated by Sfp to yield either *holo*-ACP or acetoacetyl-S-LovF.

An alternate approach was employed which relied upon a direct acylation of acetoacetate from acetoacetyl-CoA to the free thiol of the phosphopantetheine at the ACP active site (33) of *holo*-LovF. Using this direct approach, *holo*-LovF was incubated with 1 mM acetoacetyl-CoA for 15, 60, 90, and 120 min prior to tryptic digestion and MS analysis (Figure 8A). It was determined that maximal formation of acetoacetyl-S-LovF was achieved after 15 min (Figure 9B). Following direct acylation with acetoacetyl-CoA, a PPant ejection ion bearing an acetyl intermediate was detected, suggesting a retro-aldol cleavage of acetoacetyl-S-LovF to yield acetyl-S-LovF (Figure 8B). This same trend was not observed after using the truncated *apo*-ACP protein, acetoacetyl-CoA, and Sfp to form acetoacetyl-S-ACP.

Direct Observation of Additional LovF Intermediates. Direct acylation with acetoacetyl-CoA yielded sufficient quantities of acetoacetyl-S-LovF for probing the formation of additional

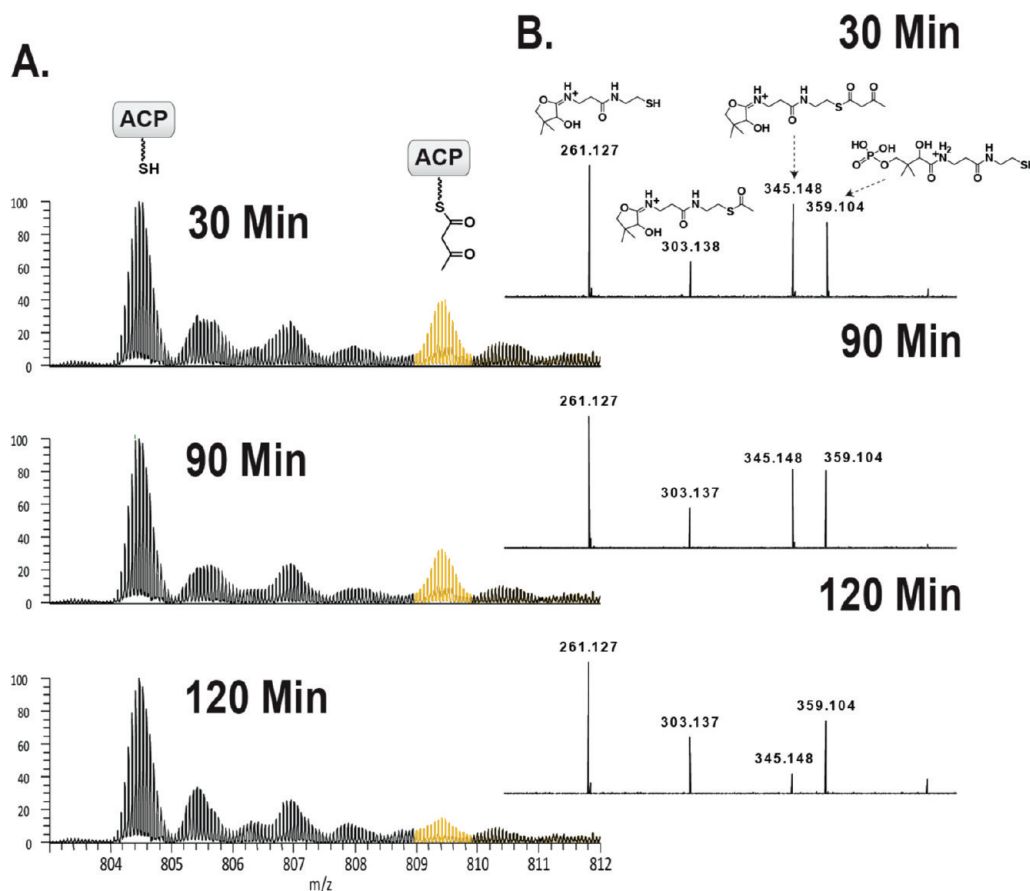


FIGURE 8: Acetoacetyl-CoA transthioesterification reaction. Samples of *holo*-LovF were incubated with 1 mM acetoacetyl-CoA (acac-CoA) in order to directly transfer acetoacetate to the free thiol of the 4'-phosphopantetheine of the ACP active site. Using this strategy it was possible to circumvent the absence of this intermediate when the *holo*-LovF was incubated with malonyl-CoA. *Holo*-LovF was incubated with 1 mM acetoacetyl-CoA for various lengths of time. (A) A mass shift of 84.0211 Da was observed on the ACP active site peptide corresponding to the formation of acetoacetyl-S-LovF. Longer incubation times were found to result in lower quantities of detectable acetoacetyl-S-LovF. The optimal incubation time was ultimately found to be 15 min. (B) FT-ICR-MS² detection of the relative abundance of the acetoacetyl-loaded PPant ejection ion. In addition, acetyl-loaded PPant ejection ion was detected.

intermediates in the presence of SAM and NADPH. In the biosynthetic pathway of α -methylbutyrate, methylation of the α -carbon is proposed to occur following formation of the diketide intermediate, which is catalyzed by the MT domain of LovF. Therefore, the logical next step following *in vitro* direct acylation was addition of SAM. *Holo*-LovF and acetoacetyl-CoA were coincubated with SAM for 15 min prior to limited trypsin digestion, and the resulting mixture was separated by HPLC as done previously. The product of C-methylation at the α -carbon of the acetoacetyl-S-LovF was readily detected on the ACP active site peptide. During our assays, >95% of the *holo*-ACP peptides were found to be loaded with the α -methylated diketide following addition of SAM; the majority of the ACP active site peptide was detected as α -methylacetoacetyl-S-ACP (Figure 9C). In the absence of SAM, a different trend was observed. After a 15 min incubation of *holo*-LovF with acetoacetyl-CoA alone, at most 60% of the ACP active site peptide was occupied (Figure 9B) and detected as acetoacetyl-S-ACP.

Following methylation of the acetoacetyl intermediate, LovF is predicted to catalyze β -ketoreduction, dehydration of the β -hydroxyl group, and α,β -enoyl reduction of the intermediate to yield α -methylbutyrate (Figure 1C). *Holo*-LovF was coincubated with acetoacetyl-CoA and SAM, and then NADPH was added followed by an additional 15, 30, or 60 min of coincubation (Figure 10A, iv–vi). This reaction yielded detectable, albeit small, amounts of the reduced β -hydroxy- α -methylbutyryl intermediate.

The formation of this β -keto reduced intermediate could only be detected via PPant ejection assay, yet it was still possible to observe a time-dependent buildup of this intermediate (Figure 10B). Under the *in vitro* incubation conditions attempted, there was no detection of the α -methylbut- α,β -enoyl intermediate, suggesting that further reduction of this intermediate to α -methylbutyrate occurs very rapidly when it is present on the ACP active site. Also, coincubation of *holo*-LovF with acetoacetyl-CoA and NADPH, while excluding SAM, did not yield a detectable amount of *shunt* intermediates or butyryl-S-ACP. The utility of the high-mass accuracy capabilities of the FT-ICR-MS for detecting the formation of the different intermediates became exceptionally clear from this series of experiments, as several of the PPant ejection ions shared the same nominal mass, yet their exact masses differed by a mere <0.04 amu.

There were two instances in which FT-ICR-MS capabilities were crucial for accurately identifying pantetheine ejection ions bearing covalent intermediates. First, we were able to determine the identity of the α -methylbutyryl-loaded ejection ion, which copurified following heterologous expression of *holo*-LovF, by FT-ICR-MS from its exact mass (m/z 345.184). Following acylation by acetoacetyl-CoA, the pantetheine ejection assay yielded an ion with a nominal mass of m/z 345 which was found to have an exact mass of m/z 345.148 (Figure 10A, ii). When attempting to observe the low intensity of the MS³ fragmentation patterns of the α -methylbutyryl-loaded and acetoacetyl-loaded

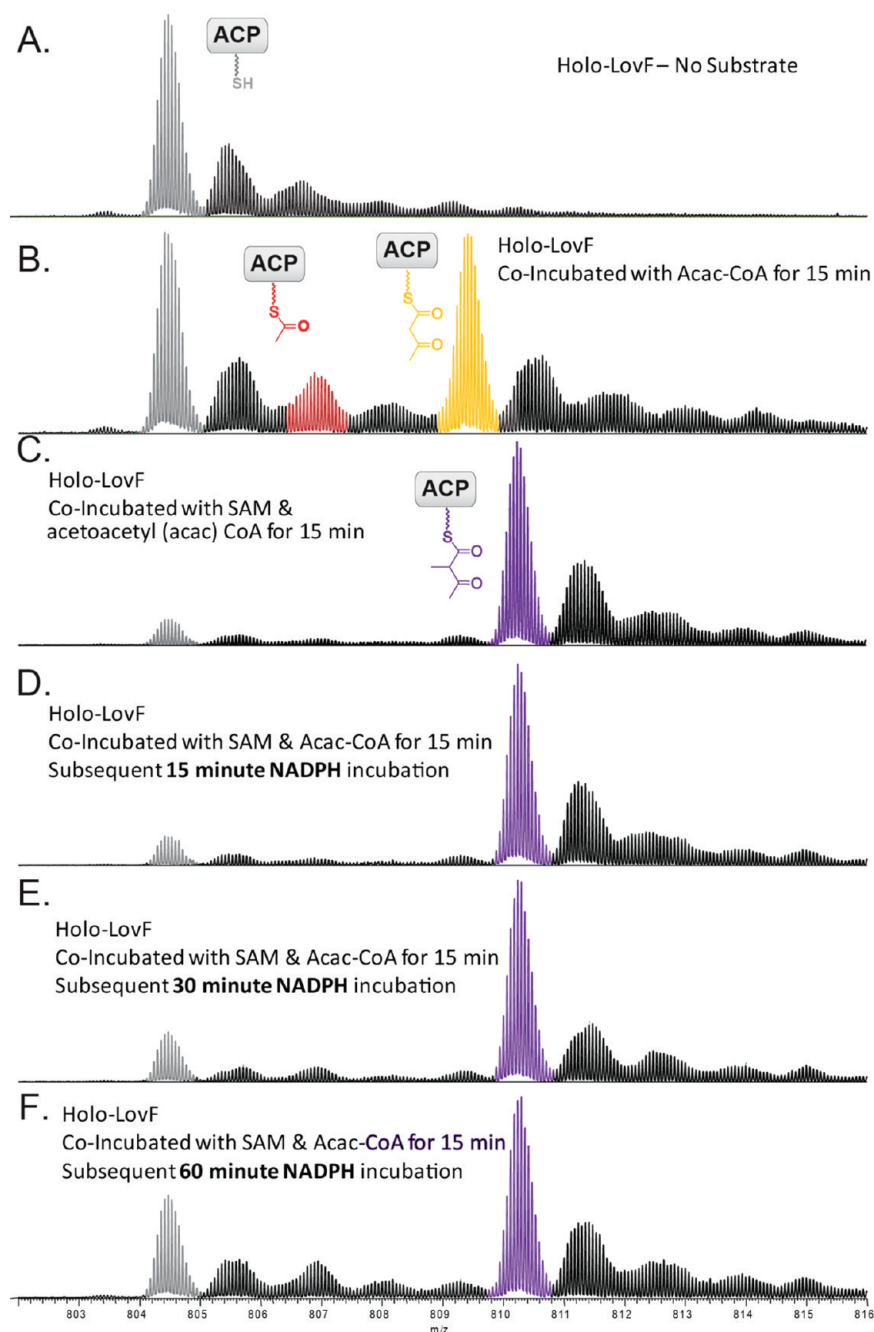


FIGURE 9: Detection of additional intermediates. Following coinubation of *holo*-LovF with acetoacetyl-CoA and SAM, NADPH was added to the solution, and the reaction was allowed to incubate for 15, 30, and 60 min prior to tryptic digestion. (A) *Holo*-LovF ACP active site for comparison. (B) *Holo*-LovF with acetoacetyl-CoA and (C) *holo*-LovF with acetoacetyl-CoA and SAM. Incubation of *holo*-LovF plus SAM, followed by subsequent incubation of NADPH for (D) 15 min, (E) 30 min, and (F) 60 min. From these three reactions, the formation of α -methylacetoacetyl-S-LovF was pronounced.

ejection ions, it was necessary to use the high-sensitivity linear ion trap detector. However, the MS³ fragmentation patterns of these two ejection ions were identical. Following incubation of *holo*-LovF with acetoacetyl-CoA and SAM, the formation of the α -methylacetoacetyl intermediate was evident on the intact ACP active site tryptic peptide. The resulting pantetheine ejection ion bearing this intermediate (m/z 359.164) could be distinguished by FT-ICR-MS from the unloaded phosphopantetheine ejection ion (m/z 359.104). When initially optimizing the direct acylation reaction conditions, the ability to detect that a small amount of the α -methylated acetoacetyl intermediate was present and to distinguish it from the phosphopantetheine ejection ion (both of which were present only in low quantities) was critical for determining

the efficacy of the various reaction conditions attempted. This feat would not have been possible on a low-resolution instrument alone. Following our discovery that direct acylation of *holo*-LovF by acetoacetyl-CoA in the presence of SAM resulted in a highly abundant, highly stable intermediate, the addition of NADPH did not immediately seem to reveal the presence of β -hydroxy- α -methylbutyryl present of the ACP active site. However, closer inspection of the pantetheine ejection ions following these reactions revealed that this intermediate was in low abundance and the monoisotopic peak of the β -hydroxy- α -methylbutyryl-loaded pantetheine ejection ion nearly overlapped with an isotope of the highly abundant α -methylacetoacetyl-loaded pantetheine ejection ion (Figure 10B, iii–vi). The two ions differed in mass by a mere

biosynthetic steps performed by an intact megasynthase. The fact that the biosynthesis of α -methylbutyrate, a diketide, involves only a handful of catalytic steps made this fungal type I polyketide synthase an attractive system to study by FT-ICR-MS. Many of the experimental hurdles that were overcome during the analysis of LovF are being used to guide ongoing mass spectrometric investigations of LovB and other iterative type I PKSs.

Deconstructed/reconstituted type I PKS systems are widely utilized to study the catalytic processes responsible for the creation of a diverse array of compounds, and for good reason: the expression of separate domains as individual proteins allows for the fine-tuning of molar ratios of different pieces of a PKS pathway in order to facilitate examination of the functional role or substrate tolerance of a given component, much the same way type II PKSs are studied (30, 38, 39). In addition, it is possible in some cases to determine the kinetics of individual catalytic steps (40). However, keys to understanding the basis for controlling many of the catalytic steps, during successive iterations, may only be gained by studying fully intact megasynthases.

After we determined the trypsin digestion conditions that consistently generated abundant quantities of the ACP active site peptide, we anticipated that incubation of *holo*-LovF with malonyl-CoA would yield readily detectable quantities of several acyl intermediates. Incubation with 1 mM malonyl-CoA alone resulted primarily in a high quantity of malonyl-S-ACP and to a lesser extent acetyl-S-ACP. Much to our surprise (and annoyance), there was no detectable acetoacetyl-S-ACP under these reaction conditions. Inclusion of SAM and NADPH did not change this observed trend, nor did shortening or increasing malonyl-CoA incubation times. Our initial hypothesis was that 1 mM concentration of malonyl-CoA was simply too high relative to the quantity of *holo*-LovF in the *in vitro* reaction, resulting in congestion of malonyl at the ACP, wherein the ACP was immediately reoccupied by malonyl after decarboxylative condensation between the starter and extender unit and transfer of the resulting diketide to the KS active site. However, lowering the concentration of malonyl-CoA during incubation with LovF failed to produce detectable levels of acetoacetyl-S-LovF or α -methylbutyryl-S-LovF. Therefore, the diketide intermediate may remain at the KS active site following condensation of the starter and extender units, and upon exiting the KS active site the intermediate undergoes α -methylation, β -ketoreduction, dehydration, and enoyl reduction very rapidly to form α -methylbutyrate. This hypothesis is supported by our previous observation that after *in vitro* reaction of LovF, LovD, monacolin J acid, and the complement of various substrates and cofactors, only α -methylbutyrate (not acetoacetate, nor α -methylacetoacetate) was transferred to monacolin J acid by LovD.

Further clues regarding the location of the acetoacetyl intermediate could be gleaned from the direct acylation experiments. We found that Sfp was unable to efficiently transfer phosphopantetheine to *apo*-LovF, while it readily converted a LovF *apo*-ACP construct to *holo*-ACP. The failure of promiscuous phosphopantetheinyl transferase (PPTase) Sfp, from *Bacillus subtilis*, to effectively transfer acetoacetyl-S-phosphopantetheine to *apo*-LovF led us to seek an alternative. As others have recently noted (33), direct acylation serves as a reliable alternative for generating acyl-S-ACPs. Direct acylation of the acetoacetate from acetoacetyl-CoA proved to be a very efficient means for generating the acetoacetyl-S-LovF, although the ACP active site was at most only 60% occupied in the presence of an $\sim 80\times$ molar excess of acetoacetyl-CoA relative to *holo*-LovF.

While it was possible to generate acetoacetyl-S-LovF by direct acylation, this intermediate did not stably remain on the ACP active site. With time, the ACP active site quickly returned to its unoccupied *holo* form even in the presence of a substantial molar excess of acetoacetyl-CoA. Whether return of the ACP active site to the unoccupied *holo* state is due to transfer of the acetoacetyl intermediate to the KS active site or due to hydrolysis of the intermediate off of the ACP active site remains unclear at this point. Following direct acylation to form acetoacetyl-S-LovF, we observed the reverse reaction of the acetoacetyl intermediate to acetyl-S-LovF. This provides further insight into why we did not detect acetoacetyl-S-LovF following incubation of LovF with malonyl-CoA, as it appears that the diketide may be susceptible to a retro-aldol cleavage. Ultimately, it was clear from these experiments that under *in vitro* conditions the diketide acetoacetate does not stably reside at the *holo*-LovF ACP active site at steady-state.

Our ability to monitor the occupancy of the active site of LovF provided us with an unprecedented snapshot of several of the catalytic steps carried out by this megasynthase. First, a small amount of α -methylbutyrate that purified with the overproduced protein in yeast was detected by FT-ICR-MS. However, ^{13}C -labeling verified that α -methylbutyrate synthesized by LovF originated from the malonyl-CoA alone under *in vitro* reaction conditions. Also, under these same conditions, the majority of the *holo*-LovF ACP active site peptides were occupied by the malonyl-S-thioester starter unit and to a lesser degree the acetyl intermediate. The small amount of α -methylbutyryl observed relative to these two ACP-bound species, in addition to lack of all other LovF intermediates, suggested that the acetoacetyl intermediate quickly progressed through all subsequent enzymatic tailoring reactions (α -methylation, β -ketoreduction, dehydration, and enoyl reduction) when it was present on the ACP active site. We observed that incubation of *holo*-LovF with acetoacetyl-CoA and SAM resulted in the abundant formation of stable α -methylacetoacetyl-S-LovF. The large quantity of α -methylacetoacetyl-S-LovF observed, after incubation of LovF with acetoacetyl-CoA and SAM, relative to all other intermediates, strongly suggested that α -methylation of the diketide, catalyzed by the MT domain, prevented the retrograde movement of the intermediate to the KS active site or protected it from hydrolysis. In the case of LovF, methylation of the diketide may act as one measure of quality control in which the enzyme prevents the additional chain elongation by limiting the accessibility of this intermediate to the KS active site. We had previously shown that under *in vitro* conditions LovD was capable of transferring both nonmethylated intermediates from LovF to the C8 position of monacolin J (via LovD), in addition to the α -methylacetoacetyl intermediate (27). Of the various lovastatin (monacolin K) analogues formed, the C8- α -methylacetoacetyl-monacolin J (monacolin X) was the predominant shunt product. Monacolin X is also produced by a mutant strain of the lovastatin producer *Monascus ruber* (41). Our observation presented in this report, that α -methylation of the diketide essentially locked in the methylated intermediate at the ACP active site ($>95\%$ as α -methylacetoacetyl-S-Lov), provides additional insight into why the monacolin X tends to be the predominant shunt product of the lovastatin biosynthetic pathway.

SUPPORTING INFORMATION AVAILABLE

Figure S1, confirmation of pantetheine-loaded intermediates by linear ion trap MS³ of loaded phosphopantetheine ejection

ions; Figure S2, detection of a minor quantity of the α -methylbutyryl-loaded PPant ejection ion active site of LovF. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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