

Comparative Characterization of the Lactimidomycin and iso-Migrastatin Biosynthetic Machineries Revealing Unusual Features for Acyltransferase-less Type I Polyketide Synthases and Providing an Opportunity To Engineer New Analogues

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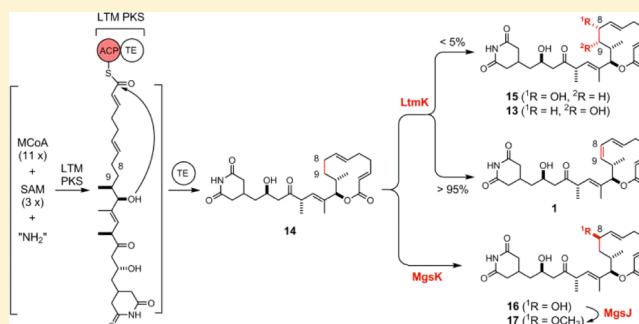
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S Supporting Information

ABSTRACT: Lactimidomycin (LTM, **1**) and iso-migrastatin (iso-MGS, **2**) belong to the glutarimide-containing polyketide family of natural products. We previously cloned and characterized the *mgs* biosynthetic gene cluster from *Streptomyces platensis* NRRL 18993. The iso-MGS biosynthetic machinery featured an acyltransferase (AT)-less type I polyketide synthase (PKS) and three tailoring enzymes (MgsIJK). We now report cloning and characterization of the *ltm* biosynthetic gene cluster from *Streptomyces amphibiosporus* ATCC 53964, which consists of nine genes that encode an AT-less type I PKS (LtmBCDEFGHL) and one tailoring enzyme (LtmK). Inactivation of *ltmE* or *ltmH*

afforded the mutant strain SB15001 or SB15002, respectively, that abolished the production of **1**, as well as the three cometabolites 8,9-dihydro-LTM (**14**), 8,9-dihydro-8S-hydroxy-LTM (**15**), and 8,9-dihydro-9R-hydroxy-LTM (**13**). Inactivation of *ltmK* yielded the mutant strain SB15003 that abolished the production of **1**, **13**, and **15** but led to the accumulation of **14**. Complementation of the Δ *ltmK* mutation in SB15003 by expressing *ltmK* in trans restored the production of **1**, as well as that of **13** and **15**. These results support the model for **1** biosynthesis, featuring an AT-less type I PKS that synthesizes **14** as the nascent polyketide intermediate and a cytochrome P450 desaturase that converts **14** to **1**, with **13** and **15** as minor cometabolites. Comparative analysis of the LTM and iso-MGS AT-less type I PKSs revealed several unusual features that deviate from those of the collinear type I PKS model. Exploitation of the tailoring enzymes for **1** and **2** biosynthesis afforded two analogues, 8,9-dihydro-8R-hydroxy-LTM (**16**) and 8,9-dihydro-8R-methoxy-LTM (**17**), that provided new insights into the structure–activity relationship of **1** and **2**. While 12-membered macrolides, featuring a combination of a hydroxyl group at C-17 and a double bond at C-8 and C-9 as found in **1**, exhibit the most potent activity, analogues with a single hydroxyl or methoxy group at C-8 or C-9 retain most of the activity whereas analogues with double substitutions at C-8 and C-9 lose significant activity.



Lactimidomycin (LTM, **1**) belongs to the glutarimide-containing polyketide family of natural products.^{1,2} Other members of this family include iso-migrastatin (iso-MGS, **2**), migrastatin (MGS, **3**), the dorriginocins (DGNs, **4–6**), NK30424A and -B (**7** and **8**, respectively), the streptimidones (**9** and **10**), and cycloheximide (**11**) (Figure 1). Originally, **1** was isolated from *Streptomyces amphibiosporus* ATCC 53964 for its potent cytotoxicity activity against various tumor cells;³ **3** was isolated in *Streptomyces* sp. MK929-43F1 for its modest tumor cell migration inhibitory activity,^{4,5} and **4–6** were isolated from *Streptomyces platensis* NRRL 18993 as inhibitors of carboxyl methyltransferases involved in the processing of

Ras-related proteins.^{6,7} Fermentation optimization subsequently resulted in the isolation of **2** from *S. platensis* NRRL 18993.⁸ Identical to the cysteine adducts of **2**,¹ **7** and **8** were first isolated from *Streptomyces* sp. NA30424 for their ability to inhibit PLS-induced TNF- α production by suppressing the NF- κ B signaling pathway.⁹ Mostly known for their antifungal activity, **9** and **10** have been isolated from various *Streptomyces* species.^{10,11} Finally, best known as an inhibitor of eukaryotic protein translation, **11** was first isolated from *Streptomyces*

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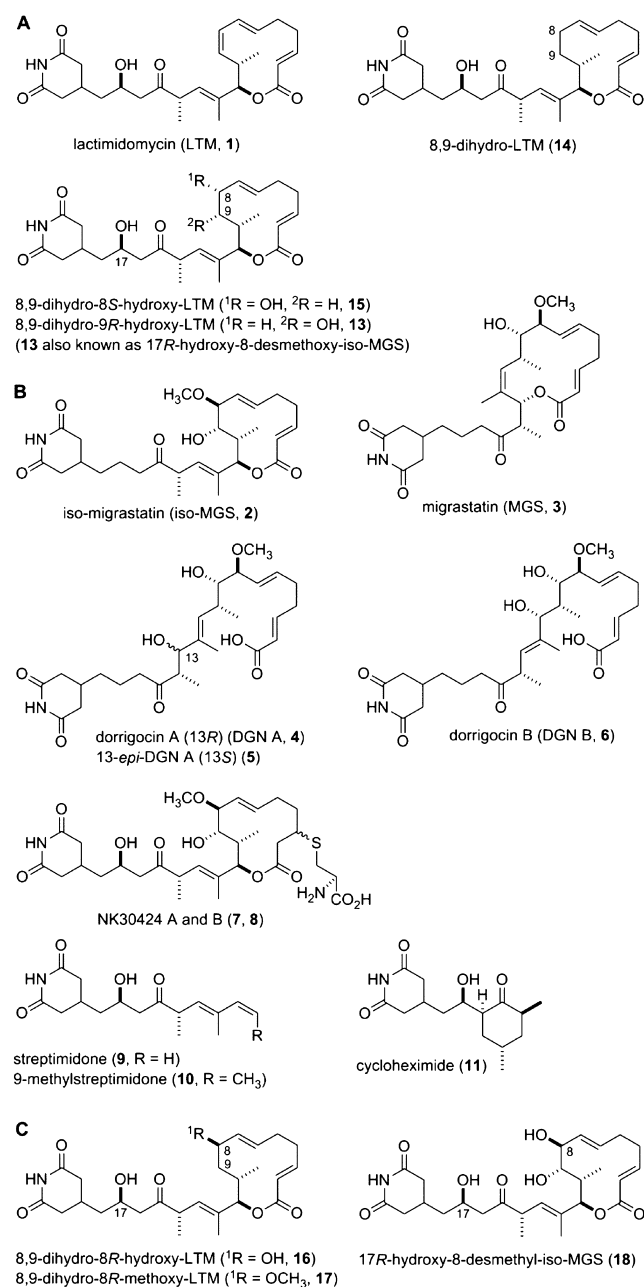


Figure 1. Structures of (A) LTM (1) and congeners (13–15) from *S. amphibiosporus* ATCC 53964, (B) selected members of the glutarimide-containing polyketide family of natural products (2–11), and (C) engineered glutarimide-containing polyketides (16 and 17) bearing structural features of both 1 and 2, and 17R-hydroxy-8-desmethyl-iso-MGS (18), a congener of 2 used as a comparison for cytotoxicity assays in Table 2.

griseus and has since been isolated from numerous *Streptomyces* species.^{12–14}

Among the multitude of activities exhibited by the glutarimide-containing polyketides, the tumor cell migration inhibitory activity of 3 and the protein translation initiation inhibitory activity of 1 have by far most captured the interests of biologists and chemists.^{1,2} Accordingly, synthetic analogues of the 14-membered macrolide 3 have been prepared, some with cell migration inhibitory activity ~3 orders of magnitude greater than that of the natural product, which have been pursued as therapeutic candidates for treating tumor meta-

stasis.^{15–17} A focused library of glutarimide-containing polyketides has been isolated from optimized fermentations of *S. platensis* NRRL 18993 and *S. amphibiosporus* ATCC 53694, and preliminary evaluation of these revealed that 12-membered macrolides, as exemplified by 1 and 2, were also potent inhibitors of tumor cell migration.^{1,2,18–20} While the exact modes of action that dictate and differentiate cell migration inhibition from cytotoxicity for the glutarimide-containing polyketides remain controversial, the actin-bundling protein fascin has been identified as the target for the cell migration inhibitory activity of 3,²¹ and blocking the translocation step in eukaryotic protein translation initiation has been deduced as the mechanism for the cytotoxicity of 1.^{22–24} The latter property of 1, in contrast to 11 that blocks the translocation steps in protein translation elongation, has been exploited in the development of the Global Translation Initiation Sequencing (GTI-seq) technology that allows high-resolution mapping of translation initiation sites across the entire transcriptome.^{23,24} Small molecule inhibitors of protein translation have also shown promise as potential chemotherapeutic agents for treating cancers.²⁵

We have previously reported that 2 is the true natural product of *S. platensis* NRRL 18993, and 3–6 are degradation products of 2 that occur during isolation, which can be readily derived from 2 via a facile, H₂O-mediated ring expansion or ring-opening rearrangement (Figure S1 of the Supporting Information).²⁶ We subsequently cloned and characterized the *mgs* biosynthetic gene cluster from *S. platensis* NRRL 18993 and confirmed 2 as the final product of the iso-MGS biosynthetic machinery, which featured an acyltransferase (AT)-less type I polyketide synthase (PKS) and three tailoring enzymes.²⁷ We have recently revealed that the iso-MGS AT-less type I PKS produces two nascent polyketide intermediates, 16,17-didehydro-8-desmethoxy-iso-MGS (12) and 17R-hydroxy-8-desmethoxy-iso-MGS (13). Further, the biosynthesis of 12 and 13 involves several unusual features that deviate from those of the collinear type I PKS model, and the three tailoring enzymes possess remarkable substrate promiscuity, accounting for the formation of all congeners of 2 known to *S. platensis* NRRL 18993 from 12 and 13 (Figure S2 of the Supporting Information).²⁸

We have also optimized the fermentation of *S. amphibiosporus* ATCC 53964 and isolated, in addition to 1, three LTM congeners, 8,9-dihydro-LTM (14), 8,9-dihydro-8S-hydroxy-LTM (15), and 8,9-dihydro-9R-hydroxy-LTM (13) (Figure 1A),¹⁸ the latter of which is identical to 17R-hydroxy-8-desmethoxy-iso-MGS (13), one of the two nascent polyketide intermediates synthesized by the iso-MGS AT-less type I PKS (Figure S2 of the Supporting Information).^{27,28} The structural similarity, with distinct variations, between 1 and 2, and the isolation of the same biosynthetic intermediate 13 from both producers of 1 and 2, suggest close relationships in their biosynthetic pathways.

Here we report the cloning and characterization of the *ltm* biosynthetic cluster from *S. amphibiosporus* ATCC 53964. The LTM biosynthetic machinery features an AT-less type I PKS nearly identical to that of 2 but a rare cytochrome P450 desaturase functioning as the sole tailoring enzyme. Comparison of the LTM and iso-MGS biosynthetic machineries allows deduction of a biosynthetic pathway for 1, reveals new insights into the biosynthesis of the glutarimide-containing polyketides, and sets the stage to investigate several unusual features common to AT-less type I PKSs. Exploitation of the tailoring

enzymes for **1** and **2** biosynthesis affords two analogues, 8,9-dihydro-8*R*-hydroxy-LTM (**16**) and 8,9-dihydro-8*R*-methoxy-LTM (**17**), which feature the 17*R*-hydroxyl group present in **1** and the 8*R*-hydroxyl or 8*R*-methoxyl groups present in **2** (Figure 1C) and provide new insights into the structure–activity relationship (SAR) of **1** and **2**.

MATERIALS AND METHODS

Bacterial Strains, Plasmids, and Culture Conditions.

Bacterial strains and plasmids used are summarized in Table S1 of the Supporting Information. *S. amphibiosporus* wild-type and recombinant strains were cultivated on ISP4 or tryptic soy broth (TSB) medium at 28 °C.²⁹ *Escherichia coli* strains were grown on lysogeny broth (LB) medium at 37 °C.³⁰ Strains carrying antibiotic resistance markers were selected in medium supplemented with ampicillin (50 µg/mL), apramycin (50 µg/mL), chloramphenicol (50 µg/mL), kanamycin (50 µg/mL), nalidixic acid (20 µg/mL), thiostrepton (5 µg/mL), or trimethoprim (50 µg/mL), where appropriate.

DNA Isolation, Manipulation, and Sequencing. Standard procedures were used to manipulate purified DNA.³⁰ Genomic DNAs were purified from *S. amphibiosporus* strains using the salting out procedure.²⁹ Plasmids were isolated using available kits (Qiagen). Cosmid DNAs were purified by the alkaline lysis method.¹⁷ PCRs were performed using the high-fidelity Expand PCR system according to the manufacturer's protocol (Roche). Oligonucleotide primers that were used are listed in Table S2 of the Supporting Information. Southern analysis of *S. amphibiosporus* genomic DNA was conducted using DIG-High Prime DNA Labeling and Detection Starter Kit I following the manufacturer's protocol (Roche).

Genome Scanning and Sequence Analysis. Isolation of the *ltm* biosynthetic gene cluster was performed by the genome scanning method as described previously.^{27,31} A SuperCos1-based library of *S. amphibiosporus* ATCC 53964 was constructed according to the manufacturer's protocol (Agilent Technologies). Three cosmids, pBS15001, pBS15002, and pBS15003, covering the entire *ltm* biosynthetic gene cluster, were identified by probing with PKS genes identified from a genome sampling library and were sequenced (Figure 2A). The DNA sequences obtained were analyzed by the Genetic Computer Group (GCG) program, BLAST-X, N, and P available at National Center for Biotechnology Information (NCBI).

Gene Inactivation in *S. amphibiosporus* by λ-RED-Mediated PCR Targeting Mutagenesis. The *ltmE*, *ltmH*, and *ltmK* genes, located on pBS15001 and pBS15003 (Figure 2A), were inactivated in *S. amphibiosporus* ATCC 53964 by λ-RED-mediated PCR targeting mutagenesis.³² Specifically designed primers (Table S2 of the Supporting Information) were used to amplify the disruption cassette from the template plasmid pJ773.³² The PCR-amplified cassette is comprised of the *aac(3)IV* gene, conferring apramycin resistance, and an RK2 origin of transfer (*oriT*), and is flanked by DNA sequence homologous to regions immediately upstream and downstream of the gene to be deleted. This cassette is used to replace the target gene on the cosmid by homologous recombination, yielding pBS15004 (Δ*ltmE*), pBS15005 (Δ*ltmK*), and pBS15008 (Δ*ltmH*) (Figure S3 of the Supporting Information). The resulting mutant cosmids were introduced into the *S. amphibiosporus* wild-type strain through intergeneric conjugation between *S. amphibiosporus* and *E. coli* S17-1.²⁹ Exconjugants with desired double-crossover recombination

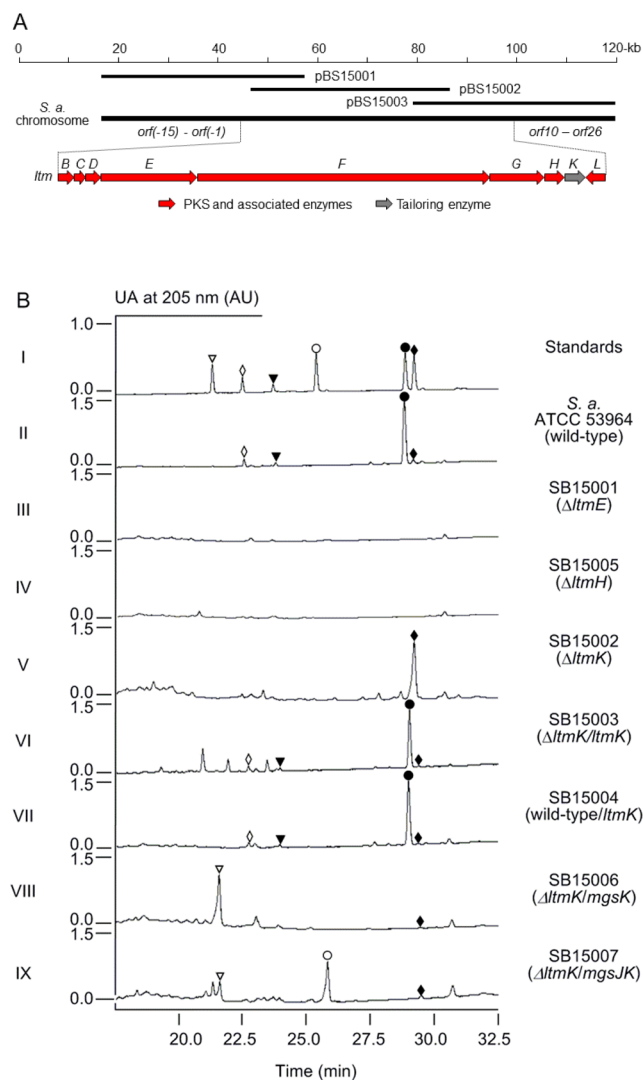


Figure 2. Cloning and sequencing of the *ltm* gene cluster from *S. amphibiosporus* ATCC 53964 and confirmation of its role in LTM (**1**) biosynthesis. (A) Sequenced 100 kb DNA region covered by three overlapping cosmids and genetic organization of the *ltm* biosynthetic gene cluster. Proposed functions for individual open reading frames are summarized in Table 1. (B) Inactivation of selected genes within the *ltm* cluster supporting the proposed pathway for **1** biosynthesis and exploitation of the tailoring enzymes for **1** and **2** biosynthesis affording two novel analogues. HPLC chromatograms show authentic standards of **1** (●), **14** (◆), **13** (▼), **15** (◇), **16** (▽), and **17** (○).

were selected on the basis of apramycin resistance and kanamycin sensitivity. The genotype of each mutant was confirmed by Southern analysis, and the mutants were subsequently named SB15001 (Δ*ltmE*), SB15002 (Δ*ltmK*), and SB15005 (Δ*ltmH*) (Figure S3 of the Supporting Information).

Generation of the Δ*ltmK* Complementation Strain SB15003. A 2.5 kb SacI–MscI DNA fragment carrying *ltmK* was first cloned from pBS15003 into pWHM79³³ as a SacI–SmaI fragment, creating pBS15006. From there, a 3.0 kb EcoRI–HindIII fragment was purified, Klenow treated (Roche), and ligated into the EcoRV site of pBS11016,²⁷ in which the *aac(3)IV* gene is replaced by the *neo* gene conferring kanamycin resistance, affording pBS15007. Expression of *ltmK* in pBS15007 is under the control of the *Erme** promoter.²⁹

Table 1. Deduced Functions of ORFs in the *ltm* Gene Cluster in Comparison with Those in the *mgs* Gene Cluster

| gene | deduced product/ amino acid ^a | protein homologues ^{a,b} | % identity/ % similarity | | proposed function ^c |
|------------------------------|---|--------------------------------------|-----------------------------|--------------------------------------|---|
| <i>orf(-15)- orf(-1)</i> | | | | | beyond the upstream boundary |
| – | | MgsA/1217 (ACY01386) | – | | transcriptional activator ^d |
| <i>ltmB</i> | LtmB/539 (ACY01397) | MgsB/539 (ACY01387) | 63/71 | | type II thioesterase |
| <i>ltmC</i> | LtmC/88 (ACY01398) | MgsC/83 (ACY01388) | 68/85 | loading module | ACP |
| <i>ltmD</i> | LtmD/656 (ACY01399) | MgsD/656 (ACY01389) | 82/87 | loading module | amidotransferase |
| <i>ltmE</i> | LtmE/3437 (ACY01400) | MgsE/3192 (ACY01390) | 68/74 | module-2, -3, -4 | KS-DH-KR-ACP-KS-B-ACP-KS |
| <i>ltmF</i> | LtmF/8360 (ACY01401) | MgsF/8021 (ACY01391) | 67/73 | module-4, -5, -6, -7, -8, -9, -10 | KR-ACP-KS-ACP-KS-DH-KR-MT-ACP-KS-KR -MT-ACP- KS-KS-DH-KR-ACP-ACP-KS-DH |
| <i>ltmG</i> | LtmG/2098 (ACY01402) | MgsG/1953 (ACY01392) | 58/68 | module-10, -11 | ACP-KS-DH-KR-ACP-TE |
| <i>ltmH</i> | LtmH/768 (ACY01403) | MgsH/751 (ACY01393) | 65/76 | | acyltransferase |
| – | | MgsI/338 (ACY01394) | – | | oxidoreductase ^d |
| – | | MgsJ/281 (ACY01395) | – | | O-methyltransferase ^d |
| <i>ltmK</i> | LtmK/413 (ACY01404) | MgsK/403 (ACY01396) | 32/47 | | cytochrome P450 desaturase |
| <i>ltmL</i> | LtmL/247 (ACY0140%) | Ppta/246 (AAG43513) | 57/65 | | phosphopantetheinyltransferase |
| <i>orf10-orf26</i> | | | | | beyond the downstream boundary |

^aNumbers are in amino acids, and given in parentheses are NCBI accession numbers. ^bHomologues from *mgs* clusters are given, and other homologues are included if *mgs* homologues are absent. ^cDomain abbreviations: ACP, acyl carrier protein; B, branching; DH, dehydratase; KR, ketoreductase; KS, ketosynthase; MT, methyltransferase; TE, thioesterase. ^dFunction proposed for the genes from the *mgs* cluster that are missing from the *ltm* cluster.

pBS15007 was transformed into *E. coli* S17-1 and conjugated into SB15002. Exconjugants were selected on the basis of resistance to apramycin and kanamycin, yielding SB15003 as the Δ *ltmK* complemented strain.

Generation of the *ltmK* Overexpression Strain SB15004. pBS15007 was transformed into *E. coli* S17-1 and conjugated into the *S. amphibiosporus* wild-type strain. Exconjugants were selected on the basis of resistance to kanamycin, yielding SB15004.

Construction of Recombinant Strains SB15006 and SB15007 for Production of Novel Analogues. The three genes, *mgsI*, *mgsJ*, and *mgsK*, encoding the three tailoring enzymes for **2** biosynthesis were amplified by PCR from pBS11006²⁷ individually. The *ErmE** promoter was amplified by PCR from pWHM79.³³ All primers for *mgsI*, *mgsJ*, *mgsK*, and *ErmE** contained different restriction enzyme sites (Table S2 of the Supporting Information). The resultant PCR products were digested accordingly and ligated sequentially into the EcoRV and XbaI sites of pBS11016²⁷ to yield pBS15009, in which the expression of each of the *mgsIJK* genes was under the control of *ErmE** (Figure S4 of the Supporting Information). pBS15009 was then digested with either Acc65I and PmeI or Acc65I and AflII, and the resultant fragments were Klenow treated and self-ligated to afford the expression construct pBS15010 (for *mgsK*) or pBS15011 (for *mgsJK*), respectively. pBS15010 or pBS15011 was finally introduced into the Δ *ltmK* mutant strain of SB15002 by conjugation, and exconjugates, selected on the basis of apramycin and kanamycin resistance, afforded the recombinant strain SB15006 (i.e., Δ *ltmK/mgsK*) or SB15007 (i.e., Δ *ltmK/mgsJK*), respectively (Figure S4 of the Supporting Information).

***S. amphibiosporus* Wild-Type and Recombinant Strain Fermentation and Isolation and HPLC Analysis of **1** and Related Metabolites **13–15**.** *S. amphibiosporus* ATCC 53964 wild-type and recombinant strains were fermented as previously described.¹⁸ Large-scale fermentation to isolate **14** from SB15002 for structural characterization was similarly conducted after incubation in production medium for 7 days. All metabolites were isolated as previously described, and their identities were confirmed by either HPLC analysis in comparison with authentic standards (for **13–15**) or ¹H and ¹³C NMR analyses (for **14**).^{18,28}

Isolation and Structural Elucidation of Analogues **16 and **17**.** SB15006 and SB15007 strains were grown on ISP4 medium for sporulation, and spores were harvested and preserved as a 20% glycerol suspension at –80 °C. Large-scale fermentations and metabolite isolation were similarly performed as previously described.¹⁸ From a 2 L fermentation of SB15006, **15** (5.0 mg) was purified, and from a 2 L fermentation of SB15007, **15** (2.0 mg) and **16** (5.2 mg) were purified.

8,9-Dihydro-8R-hydroxy-LTM (16**).** Colorless oil: $[\alpha]_D^{27} + 95.4$ (*c* 0.24, DMSO); ¹H NMR (700 MHz) and ¹³C NMR (175 MHz) data in Table S3 of the Supporting Information; HRESIMS for the $[M + Na]^+$ ion at *m/z* 498.2461 (calcd $[M + Na]^+$ ion for C₂₆H₃₇NO₇ at *m/z* 498.2466).

8, 9-Dihydro-8R-methoxy-LTM (17**).** Colorless oil: $[\alpha]_D^{27} + 123.3$ (*c* 0.24, DMSO); ¹H NMR (700 MHz) and ¹³C NMR (175 MHz) data in Table S3 of the Supporting Information; HRESIMS for the $[M + Na]^+$ ion at *m/z* 512.2617 (calcd $[M + Na]^+$ ion for C₂₇H₃₉NO₇ at *m/z* 512.2623).

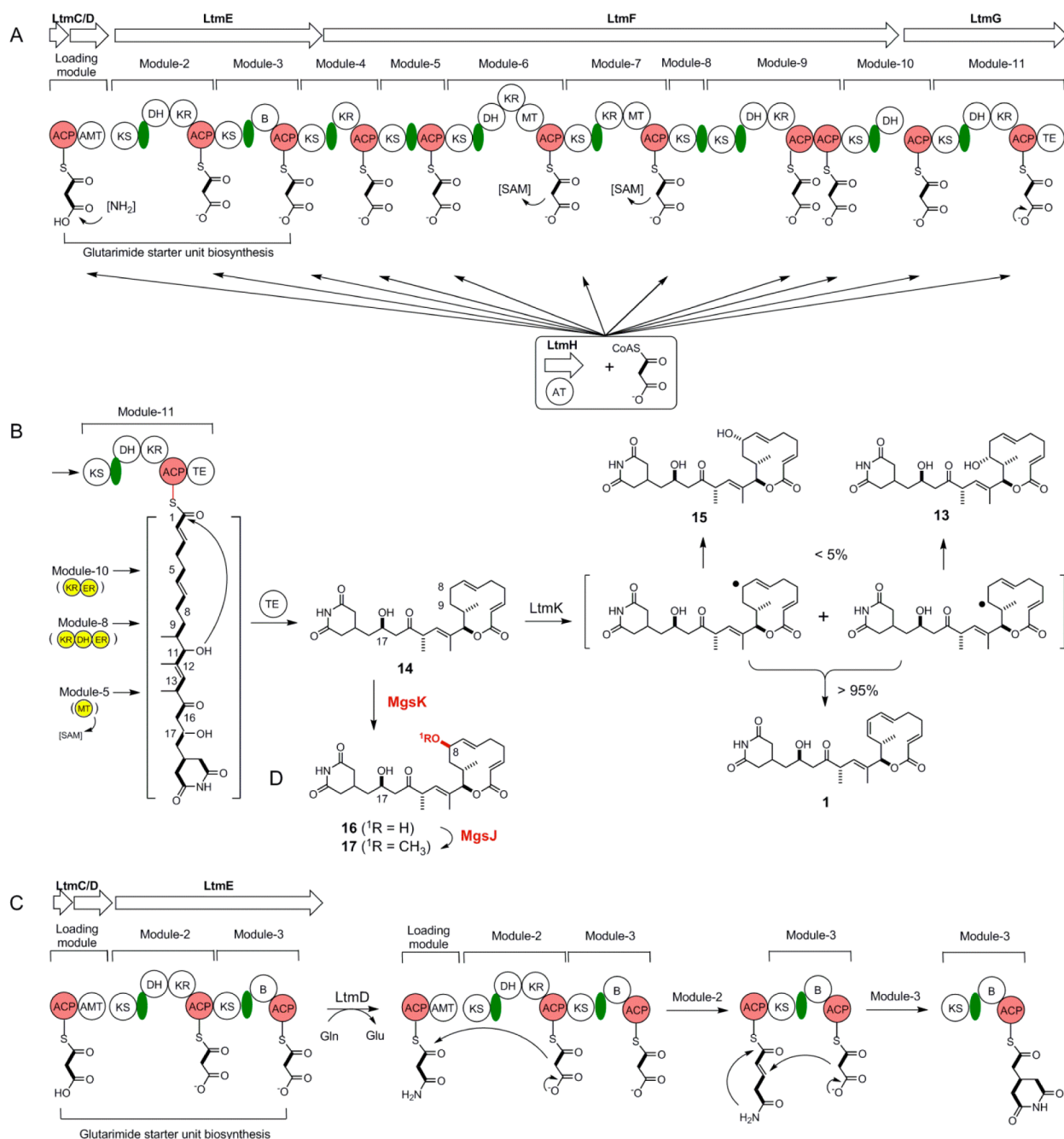


Figure 3. Proposed biosynthetic pathway for LTM (**1**) featuring the LTM AT-less type I PKS that affords **14** as the nascent product and the LtmK P450 desaturase that catalyzes the conversion of **14** to **1**, and production of novel analogues by exploiting the tailoring enzymes in **1** and **2** biosynthesis. (A) Schematic depiction of the LTM AT-less type I PKS consisting of LtmCDEFKG with the discrete LtmH AT that acts iteratively in trans, loading malonate onto each of the 11 modules. (B) LtmK P450 desaturase converts **14** to **1** via transient radical intermediates accompanied by **13** and **15** as minor hydroxylated products. (C) Revised proposal for glutarimide starter unit biosynthesis featuring the LtmD amidotransferase that furnishes an ACP-tethered malonamoyl thioester intermediate and the KS-B-ACP tridomain PKS module-3 of LtmE that affords the ACP-tethered glutarimide starter unit via a Michael addition. (D) Production of novel analogues **16** and **17**, featuring the 17R-hydroxyl group as in **1** and 8R-hydroxyl or 8R-methoxyl group as in **2**, by expressing *mgsK* or *mgsJ* in the Δ *ltmK* mutant. Abbreviations: ACP, acyl carrier protein; AMT, amidotransferase; B, branching; DH, dehydratase; ER, enoylreductase; KR, ketoreductase; KS, ketosynthase; MT, methyltransferase; TE, thioesterase. [NH₂] represents the amino donor for the AMT domain. [SAM] denotes S-adenosylmethionine as the methyl donor for the MT domain. Green ovals represent AT-docking domains, and the yellow domains highlight the missing domains for the LTM AT-less type I PKS predicted according to the colinearity model for **14** biosynthesis.

Cytotoxicity Assay. The cytotoxicity assay was conducted using the MTT method as previously described.³⁴ Generally, cells were seeded in 96-well plates (HeLa cervical carcinoma cells, 3000 per well; MDA-MB-231 breast cancer cells, 10000 per well; and Jurkat T cell lymphoma cells, 40000 per well) and

cultured in the presence of serial dilutions of the test compound, with vehicle as a negative control, in a 5% CO₂ incubator for 4 days. MTT reagent (Millipore CT0-A) was added (10 μ L/well), and incubation continued for 5 h. Then, 100 μ L of 2-propanol with 0.04 N HCl was added to each well,

and UV absorption for each well was determined using a Synergy II plate reader at 570 and 630 nm. Experiments were conducted in triplicate and repeated twice. Representative dose–response curves are shown in Figure S5 of the Supporting Information, from which EC_{50} values were determined using GraphPad Prism (Table 2).

RESULTS

Cloning of the *ltm* Biosynthetic Gene Cluster from *S. amphibiosporus* ATCC 53964. The *ltm* biosynthetic gene cluster was cloned from *S. amphibiosporus* ATCC 53964 by genome scanning (Materials and Methods).³¹ Genome scanning and sequence analysis of a 100 kb DNA region from *S. amphibiosporus* ATCC 53964 revealed a gene cluster that is highly conserved with the *mgs* gene cluster of *S. platensis* NRRL 18993 (Figure 2A and Figure S2 of the Supporting Information). The putative *ltm* cluster was covered by three overlapping cosmids, pBS15001, pBS15002, and pBS15003. Construction of the SB15001 ($\Delta ltmE$), SB15005 ($\Delta ltmH$), and SB15002 ($\Delta ltmK$) mutant strains and subsequent metabolite analysis by HPLC showed these mutants all abolished LTM production, confirming the involvement of this gene cluster in LTM biosynthesis (Figure 2B). Because the boundary of the *mgs* cluster in *S. platensis* had been previously determined by systematic gene inactivation,²⁷ we assigned the boundary of the *ltm* cluster by direct comparison between the *mgs* and *ltm* clusters (Figure 2A and Figure S2A of the Supporting Information). The *ltm* cluster consists of nine genes, and as would be expected from the structural similarity between iso-MGS and LTM (Figure 1), eight of the nine genes within the *ltm* cluster shared significant homology with genes involved in iso-MGS production in *S. platensis* NRRL 18993 (Figure 2A, Table 1, and Figure S2 of the Supporting Information).

Assignment and Annotation of the *ltm* Cluster. The overall organization of the *ltm* cluster is highly conserved with respect to the *mgs* cluster from *S. platensis* NRRL 18993 (Figure S2A of the Supporting Information), and the deduced gene products from both clusters show remarkable sequence homologies (Table 1 and Figure S2B of the Supporting Information). ORFs immediately upstream [*orf*(–15)–*orf*(–1)] and downstream [*orf*10–*orf*26] of the *ltm* cluster encode proteins with no apparent function in I biosynthesis. These findings greatly facilitated the assignment of the *ltm* cluster to nine genes (LtmBCDEFGHKL) that span approximately 55 kb (Figure 2A and Table 1).

Eight of the nine genes within the *ltm* cluster have homologues in the *mgs* cluster (Table 1 and Figure S2A,B of the Supporting Information).²⁷ As a result, *ltmC* encodes an acyl carrier protein (ACP), *ltmD* encodes an amidotransferase, *ltmEFG* encode three AT-less type I PKSs, *ltmH* encodes a discrete AT, and *ltmB* encodes a type II thioesterase (TE). The three LtmEFG AT-less type I PKSs together comprise 10 PKS modules. Bioinformatics analyses of each of the 10 modules revealed that all ketosynthase (KS) and ACP domains are functional as they contain the conserved Cys-His-His catalytic triad for KS domains and the Ser residue for phosphopantetheinylation in ACP domains. However, module-8 and module-9 of LtmF are unusual, with the former lacking an ACP domain and the latter containing two ACP domains. Other domains within the 10 PKS modules are also predicted to be functional. Thus, the six ketoreductase (KR) domains (in module-2, -4, -6, -7, -9, and -11) all have the conserved GxGxxG motif for NAD(P)H binding. The five dehydratase (DH) domains (in

module-2, -6, -9, -10, and -11) all contain the conserved HxxxGxxxxP motif. Both methyltransferase (MT) domains (in module-6 and -7) are characterized with the ExxxGxG motif highly conserved among various S-adenosylmethionine (SAM)-dependent MTs. Module-11 of LtmG features a TE domain at its C-terminus, which catalyzes macrolactonization and release of the completed polyketide product. Taken together, LtmCDEFG constitute the functional LTM AT-less type I PKS, with LtmH providing the AT activity in trans, for the biosynthesis of the nascent glutarimide-containing polyketide intermediate (Figure 3A,B). The glutarimide starter unit is assembled by the loading module of LtmCD and PKS module-2 and -3 of LtmE. LtmE module-3 features the newly discovered branching (B) domain, a hallmark of PKS modules that generate a β -branch by catalyzing a Michael addition of a C2 unit to the growing polyketide chain (Figure 3C).^{11,14,35,36} LtmB, a type II TE, plays an editing role to ensure polyketide biosynthesis by removing mischarged substrates or aberrant intermediates that might otherwise block the LTM AT-less type I PKS.^{37,38}

Distinct differences also exist between the *mgs* and *ltm* clusters (Table 1 and Figure S2 of the Supporting Information). Within the *ltm* cluster, pathway-specific regulatory genes are absent, whereas the *mgs* cluster has a regulatory *mgsA* gene. While there are three genes within the *mgs* cluster that encode the tailoring enzymes MgsIJK, *ltmK* is the only gene within the *ltm* cluster that encodes a tailoring enzyme, indicative of variations in post-PKS modifications between 1 and 2 biosynthesis. Sequence analysis revealed that LtmK is a cytochrome P450 protein with the conserved oxygen-activating (GxxT) and heme binding (GxxxCxG) motifs, both critical for the activity of P450 enzymes. LtmK shares homology with MgsK (32% identical and 47% similar)¹⁷ and other cytochrome P450 monooxygenases, such as PldB (39% identical and 54% similar) for pladienolide B biosynthesis from *S. platensis* Mer-11107³⁹ and Ema1 (34% identical and 50% similar) for avermectin biosynthesis from *Streptomyces tubercidicus* R-922 (Figure S1D of the Supporting Information).⁴⁰ Finally, the *ltm* cluster contains an additional *ltmL* gene that is absent in the *mgs* cluster. LtmL shares homology with phosphopantetheinyl transferases, including PptA (63% identical and 72% similar) for bleomycin biosynthesis from *Streptomyces verticillus*,⁴¹ SePptII (57% identical and 65% similar) for erythromycin biosynthesis from *Saccharopolyspora erythraea*,⁴² and NysF (57% identical and 65% similar) for nystatin biosynthesis from *Streptomyces noursei*.⁴³ LtmL ensures that the LTM AT-less type I PKS is in its functional holo form by phosphopantetheinylating the serine residues of each of the 11 ACPs.⁴¹

Development of a Genetic System for *S. amphibiosporus* ATCC 53694. An efficient genetic system for *S. amphibiosporus* ATCC 53694 was established by *E. coli*–*Streptomyces* intergeneric conjugation (Materials and Methods). *S. amphibiosporus* grows well on ISP4 medium, and sporulation typically requires incubation for 10–14 days at 28–30 °C. The *S. amphibiosporus* wild-type strain shows extreme sensitivity to apramycin, kanamycin, and thiostrepton, but its growth is not affected by trimethoprim (50 μ g/mL) or nalidixic acid (20 μ g/mL). Although protoplast-mediated transformation in *S. amphibiosporus* was not successful under all conditions examined, *E. coli*–*S. amphibiosporus* intergeneric conjugation can proceed with high efficiency.²⁹ With *E. coli* S17-1 (DNA methylation proficient) or *E. coli* ET12567/pUZ8002 (DNA methylation deficient) as donors and *S. amphibiosporus* spores

as recipients, integrative plasmids such as pSET152 can be introduced into *S. amphibiosporus* with conjugation frequencies of 10^{-6} or 10^{-7} , respectively (Table S1 of the Supporting Information). Selected *E. coli*–*Streptomyces* shuttle vectors, commonly used in *Streptomyces* genetics, including pKC1218, pKC1139, pBS3031, and pHZ1358 (Table S1 of the Supporting Information), with copy numbers varying from one to 300 copies per cell,²⁹ were all successfully introduced by conjugation and were stably maintained in *S. amphibiosporus* ATCC 53694.

Inactivation of *ltmE* and *ltmH* Abolishing the Production of **1, as Well as **13**–**15**, in SB15001 and SB15005.** The identity of the cloned gene cluster that encodes the production of **1** was confirmed by inactivation of *ltmE* and *ltmH* in *S. amphibiosporus* ATCC 53694 (Materials and Methods). The *ltmE* gene encodes module-2 and -3 of the LTM AT-less type I PKS that is responsible for the biosynthesis of the glutarimide starter unit, while *ltmH* encodes the sole AT that loads malonyl-CoA onto each of the PKS modules for LTM biosynthesis (Table 1 and Figure 3A,C). Both *ltmE* and *ltmH* were inactivated by replacement of an internal fragment, encoding the KS domain in PKS module-2 of *LtmE*, and a 2.3 kb fragment, encoding essentially the entire *LtmH*, respectively, with the *aac(3)IV-oriT* cassette through λ -RED-mediated PCR targeting mutagenesis.³² The genotypes of the resulting mutant strains of SB15001 (i.e., $\Delta ltmE$) and SB15005 (i.e., $\Delta ltmH$) were confirmed by Southern analysis (Figure S3A,B of the Supporting Information). Both SB15001 and SB15005 were fermented under established conditions for LTM production¹⁸ with the *S. amphibiosporus* wild-type strain as a positive control. While the wild-type strain produced **1**, with **13**–**15** as minor metabolites (Figure 2B, trace II), HPLC analysis revealed that SB15001 and SB15005 completely abolished the production of **1** and the three minor cometabolites **13**–**15** (Figure 2B, traces III and IV).

Inactivation of *ltmK* Abolishing **1 Production and Leading to the Accumulation of **14** as the Nascent Polyketide Intermediate.** The *ltmK* gene encodes the sole post-PKS tailoring enzyme in **1** biosynthesis, and inactivation of *ltmK* would therefore allow one to identify the nascent polyketide intermediate and explore the exact role of *LtmK* in tailoring the penultimate intermediate to **1** (Figure 3B). The *ltmK* gene was inactivated by replacing the entire gene with the *aac(3)IV-oriT* cassette through λ -RED-mediated PCR targeting mutagenesis,³² and the genotype of the resulting SB15002 strain was confirmed by Southern analysis (Figure S3C of the Supporting Information). To complement the $\Delta ltmK$ mutation in SB15002, pBS15007, a construct in which the expression of *ltmK* is under control of the constitutive *Erme** promoter,²⁹ was introduced into SB15002 by conjugation to afford SB15003. Finally, to investigate the effect of overexpression of *ltmK* on the production of **1**, pBS15007 was also introduced into the *S. amphibiosporus* wild-type strain by conjugation to yield SB15004. SB15002, SB15003, and SB15004 were subjected to fermentation, under the previously established condition for **1** production,¹⁸ with the *S. amphibiosporus* wild-type strain as a positive control. HPLC analysis indicated that SB15002 completely lost its ability to produce **1**, as well as **13** and **15**, but rather exclusively accumulated a compound with a retention time identical to that of **14** (Figure 2B, trace V). This compound was isolated and characterized by ¹H and ¹³C NMR, confirming its identity as **14** (Figure S6 of the Supporting Information).¹⁸ Production of **1**, together with **13** and **15**, as

well as **14**, was restored to levels comparable to that of the *S. amphibiosporus* wild-type strain in SB15003 (Figure 2B, trace VI). Finally, SB15004 produced an amount of **1** similar to that in the *S. amphibiosporus* wild-type strain (Figure 2B, trace VII), indicating that the *LtmK*-catalyzed tailoring step from **14** to **1** was not rate-limiting in **1** biosynthesis. Collectively, these data unambiguously established **14** as the nascent product for the LTM AT-less type I PKS (Figure 3). On the basis of the iso-MGS biosynthetic paradigm, we now propose a similar model for LTM biosynthesis that features the LTM AT-less type I PKS (Figure 3).

Expression of *mgsK* or *mgsJK* in the $\Delta ltmK$ Mutant Resulting in the Production of Novel Analogues. We have previously established that the three tailoring enzymes *MgsIJK* possessed remarkable substrate promiscuity, accounting for the formation of the various congeners of **2** known to *S. platensis* NRRL 18993 (Figure S2C of the Supporting Information).^{27,28} We thus reasoned that *MgsK* could recognize **14** as a substrate, catalyzing regio- and stereoselective C-8 hydroxylation, the product of which could be further methylated by *MgsJ* to produce an 8-methoxy analogue (Figure 3D). Two plasmids, pBS15010 and pBS15011, in which the expression of *mgsK* or *mgsJK* is under the control of the constitutive *Erme** promoter, were constructed and introduced into the $\Delta ltmK$ mutant SB15002 (Figure S4 of the Supporting Information). The resultant recombinant strains SB15006 and SB15007 were fermented under the same conditions previously established for production of **1**, with SB15002 as a control. Upon HPLC analysis, it was apparent that **14** was essentially all converted into a new product (**16**) in SB15006 (Figure 2B, trace VIII), which was then further converted into another new product (**17**) in SB15007 (Figure 2B, trace IX). The two new products were isolated from large-scale fermentations of SB15006 and SB15007, and their structures were elucidated by a combination of spectroscopic analysis. Compounds **16** and **17**, featuring the 17R-hydroxyl group as in **1** and the 8R-hydroxyl or 8R-methoxyl groups as in **2**, are novel analogues that cannot be produced by either the LTM or iso-MGS biosynthetic machinery alone, once again highlighting the power of combinatorial biosynthesis in natural product structural diversity.

Structural Elucidation of **16 as 8,9-Dihydro-8R-hydroxy-LTM and **17** as 8,9-Dihydro-8R-methoxy-LTM.** Both **16** and **17** were isolated as colorless oils (Materials and Methods). High-resolution electrospray ionization (HRESI) MS analysis of **16** afforded an $[M + Na]^+$ ion at m/z 498.2461, establishing its molecular formula as $C_{26}H_{37}NO_7$ (calcd $[M + Na]^+$ ion at m/z 498.2466). The ¹H, ¹H–¹H COSY, and ¹³C NMR spectra of **16** are very similar to those of **15** (Figures S7 and S8 of the Supporting Information),¹⁸ except that the signals at δ_H 5.27 (dd, $J = 15.5, 8.0$ Hz, H-7), δ_H 4.45 (m, H-8), and δ_H 2.15 (m, H-10) in **15** are shifted to δ_H 5.49 (dd, $J = 15.6, 3.8$ Hz, H-7), δ_H 4.14 (m, H-8), and δ_H 1.79 (m, H-10) in **16**, respectively, in the ¹H NMR spectrum and that signals at δ_C 73.4 (C-8) and δ_C 39.4 (C-9) in **15** are shifted to δ_C 68.9 (C-8) and δ_C 42.2 (C-9) in **16**, respectively, in the ¹³C NMR spectrum (Table S3 of the Supporting Information). The correlations of H-8 with C-6, C-7, and C-9 and H-9 with C-8 in the HMBC spectrum of **16** established the hydroxyl group attachment at the C-8 position (Figure 4A). Thus, the only difference between **16** and **15** is the absolute configuration at C-8, and the coupling constant values between H-7 and H-8 are diagnostic in determining the absolute configuration at C-8.

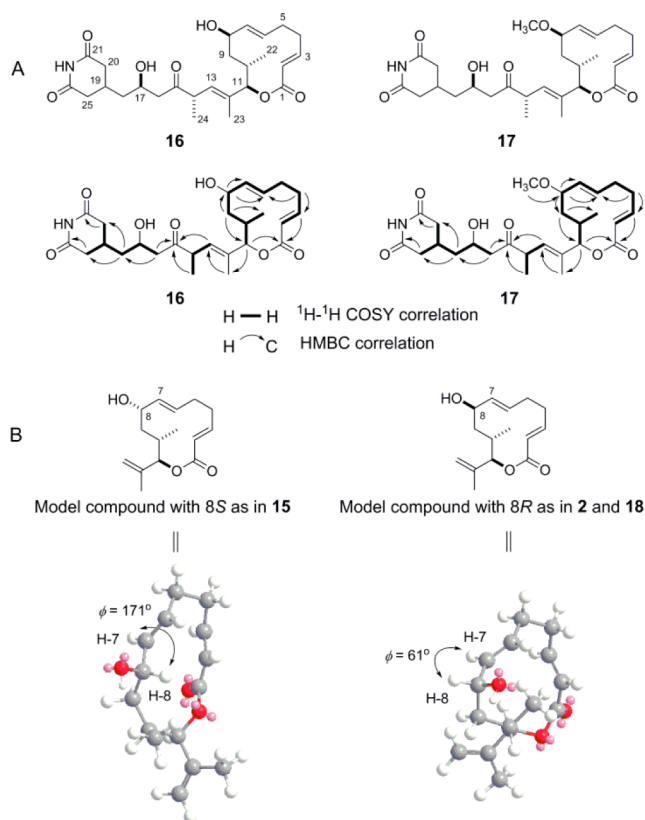


Figure 4. Structures of the two novel analogues 8,9-dihydro-8R-hydroxy-LTM (**16**) and 8,9-dihydro-8R-methoxy-LTM (**17**). (A) Key ^1H – ^1H COSY and HMBC correlations of **16** and **17** supporting their structural assignments. (B) MM2 energy-minimized configurations of model 12-membered macrolides with an 8S- or 8R-hydroxyl group, depicting the predicted H-7/H-8 dihedral angles that support the 8R absolute configurations for **16** and **17** on the basis of the H-7/H-8 coupling constants.

These observations agreed well with MM2 energy-minimized configurations for the model 12-membered macrolides with an 8S- or 8R-hydroxyl group, where the dihedral angle (ϕ) for H-7/H-8 is 171° (hence, a large coupling constant) or 61° (hence, a small coupling constant), respectively (Figure 4B). Thus, as exemplified by **15** that contains an 8S-hydroxyl group, the observed H-7/H-8 coupling constant is 8.0 Hz.¹⁸ In contrast, for **2** or 17R-hydroxy-8-desmethyl-iso-MGS (**18**), each of which contains a methoxyl and hydroxyl group at C-8 with an opposite configuration to **15** (Figure 1), the observed H-7/H-8 coupling constant value is 3.5 or 4.0 Hz, respectively.^{18,28} The observed H-7/H-8 coupling constant in **16** was 3.8 Hz (Table S3 of the Supporting Information), thereby assigning the 8R absolute configuration to **16**. Thus, **16** was established as 8,9-dihydro-8R-hydroxy-LTM, and this structure is consistent with

MgsK catalyzing the same regio- and stereoselective hydroxylation as it does for **2** and its congeners.²⁸

The structure of **17** was established in a manner similar to that used for **16**. Thus, HRESI-MS analysis of **17** afforded an $[\text{M} + \text{Na}]^+$ ion at m/z 512.2617, establishing its molecular formula as $\text{C}_{27}\text{H}_{39}\text{NO}_7$ (calcd $[\text{M} + \text{Na}]^+$ ion at m/z 512.2623). The ^1H , ^1H – ^1H COSY, and ^{13}C NMR spectra of **17** are very similar to those of **16** (Figures S9 and S10 of the Supporting Information), except that the signals at δ_{H} 5.49 (dd, $J = 15.6$, 3.8 Hz, H-7) and δ_{H} 4.14 (m, H-8) in **16** are shifted to δ_{H} 5.27 (dd, $J = 15.8$, 4.4 Hz, H-7) and δ_{H} 3.54 (m, H-8) in **17**, respectively, in the ^1H NMR spectrum and that signals at δ_{C} 138.4 (C-7), δ_{C} 68.9 (C-8), and δ_{C} 42.2 (C-9) in **16** are shifted to δ_{C} 134.5 (C-7), δ_{C} 78.4 (C-8), and δ_{C} 40.7 (C-9) in **17**, respectively, in the ^{13}C NMR spectrum (Table S3 of the Supporting Information). Compound **17** also has extra signals attributed to a methyl group at δ_{H} 3.29 (s) and δ_{C} 56.5. These data indicate that **17** is an O-methylated product at the C-8 hydroxyl group of **16**, and the correlations of H-8 with C-6, C-7, and C-9 and $\text{H}_3\text{C-O}$ with C-8 in the HMBC spectrum clearly established the methoxyl attachment at C-8 in **17** (Figure 4A). Finally, **17** was established as 8,9-dihydro-8R-methoxy-LTM, the 8R absolute stereochemistry of which was similarly assigned on the basis of the relatively small H-7/H-8 coupling constant of 4.4 Hz (Table S3 of the Supporting Information).

Cytotoxicity Assay of the Two New Analogues **15** and **16** in Comparison with **1**, **2**, and Selected Congeners.

The SAR emerging from the preliminary screening of a previously constructed glutarimide-containing polyketide library has revealed that (i) 12-membered macrolides, as exemplified by **1** and **2**, have more potent cytotoxicity against select cancer cell lines than their 14-membered and ring-open congeners, such as **3** and **4**–**6**, respectively, and (ii) congeners containing a C-17 hydroxyl group are generally more potent than their C-16/C-17 dehydrated or saturated counterparts (Figure 1).^{20,22} The new analogues **16** and **17**, featuring both the 17R-hydroxyl group as in **1** and the 8R-hydroxyl or 8R-methoxyl group as in **2**, a combination of functional groups that have not been evaluated previously, presented an opportunity to expand the SAR beyond the natural **1** and **2** congeners. Thus, **16** and **17** were assayed for cytotoxicity against three cancer cell lines, MDA-MB-231 breast cancer cells, HeLa cervical carcinoma cells, and Jurkat T lymphoma cells, and their activities were compared with the activities of **1**, **2**, **13**, **15**, and **18** (Materials and Methods).^{20,22} The seven compounds are all 12-membered macrolides and, with the exception of **2**, all feature the C-17 hydroxyl group yet have varying substitutions at positions C-8 and C-9 (Figure 1).

As summarized in Table 2 (also see Figure S5 of the Supporting Information), while **1** was most potent against all cell lines, compounds **13** and **15**–**17**, which contain 9R-hydroxyl, 8S-hydroxyl, 8R-hydroxyl, and 8R-methoxyl groups, respectively, were more potent than **18**, which contains two

Table 2. EC₅₀ Values (nanomolar) of the Novel Analogues **16** and **17** against Selected Cancer Cell Lines in Comparison with Those of **1** and **2** and Selected Congeners **13**, **15**, and **18**^a

| cancer cell line | 2 | 18 | 15 | 13 | 16 | 17 | 1 |
|------------------|----------|--------------------|-----------|-----------|-----------|-----------|----------|
| MDA-MB-231 | 464 | 1.24×10^3 | 47.6 | 172 | 157 | 87.4 | 11.9 |
| HeLa | 110 | 346 | 11.1 | 35.1 | 30.7 | 13.4 | 2.50 |
| Jurkat | 688 | 1.30×10^3 | 51.8 | 175 | 112 | 115 | 18.0 |

^aSee Figure 1 for structures.

hydroxyl groups at 8R and 9R, and **2** and **18** were the least potent among all the compounds tested. These findings reinforce the significance of the C-17 hydroxyl group but now reveal new insights into the C-8 and C-9 substitution. (i) While a double bond at C-8 and C-9 as found in **1** is the most potent, a single substitution at C-8 or C-9 such as in **13** and **15–17** retains significant activity, whereas double substitutions at C-8 and C-9 such as in **2** and **18** significantly reduce activity. (ii) While there is no significant difference between C-8 and C-9 substitution as in **16** and **13**, the 8S-hydroxyl group, as in **15**, appears to be more potent than the 8R-hydroxy counterpart as in **16**. (iii) O-Methylation of the 8R-hydroxyl group also increases, albeit modestly, the potency, as in **15** and **16**. These findings should facilitate the design, engineering, and preparation of additional analogues of **1** and **2** for anticancer drug discovery and development, and **15** and **17**, which have low nanomolar potency on par with that of **1**, warrant further mechanistic and *in vivo* evaluation.

DISCUSSION

LTM (**1**) is a potent inhibitor of tumor cell migration and eukaryotic protein translation. Cloning and sequencing of the *ltm* biosynthetic gene cluster from *S. amphibiosporus* ATCC 53964 revealed that it is highly similar to the *mgs* gene cluster from *S. platensis* NRRL 18993 that encode iso-MGS (**2**) biosynthesis. The *ltm* cluster consists of nine genes that encode an AT-less type I PKS (LtmBCDEFGHL) for biosynthesis of the nascent glutarimide-containing polyketide intermediate **14** and a post-PKS tailoring enzyme (LtmK) that converts **14** to **1**. Inactivation of *ltmE* or *ltmH* completely abolished the production of **1**, while inactivation of *ltmK* yielded a mutant strain in which **14** accumulated. Further, the production of **1** can be restored in the Δ *ltmK* mutant by expressing *ltmK* in trans. These results support a model for **1** biosynthesis that features the LTM AT-less type I PKS that is very similar to the iso-MGS AT-less type I PKS. AT-less type I PKSs harbor unusual domains and PKS architecture and often violate the colinearity rule that is a hallmark for type I PKSs. Comparative studies of the LTM and iso-MGS PKSs now provide an excellent opportunity to probe how AT-less type I PKSs select and control reductive modifications during polyketide chain elongation.

LtmCDEFG consist of the LTM AT-less type I PKS with LtmH acting iteratively in trans to load malonate onto each of the 11 modules (Figure 3A). Although the exact mechanism is not established, the LtmD amidotransferase likely catalyzes the formation of the signature glutarimide moiety by transferring the amine group from glutamine to the LtmC-tethered malonate on the loading module, in a manner akin to that of MgsD,²⁷ SmdH,¹¹ and ChxD.¹⁴ After one cycle of elongation by LtmE module-2, the formation of the glutarimide moiety invokes an unusual PKS chemistry, requiring LtmE module-3 to catalyze a Michael addition of the C2 unit to the α,β -unsaturated thioester (Figure 3C). While LtmE module-3 is characterized with a typical KS and an ACP domain, it features the newly discovered B domain.^{11,14,35,36} Recent structural and biochemical characterization of the KS-B didomain for rhizoxin biosynthesis from *Burkholderia rhizoxinica* has confirmed their role in a PKS-mediated Michael addition, and the KS-B-ACP domain architecture has been proposed as a hallmark to search for PKS modules that catalyze β -branching in polyketide biosynthesis.³⁶ Previously, this domain has been annotated as an unknown domain in iso-MGS biosynthesis²⁷ and as an X

domain in 9-methylstreptimidone biosynthesis.¹¹ On the basis of the recent structural and biochemical evidence of the KS-B domain for rhizoxin biosynthesis, we have now adopted the B domain nomenclature for the glutarimide-containing polyketides.¹⁴ The revised proposal for glutarimide biosynthesis sets the stage for characterizing the novel biochemistry *in vitro* (Figure 3C).

Following the assembly of the glutarimide starter unit, eight additional rounds of extensions catalyzed by PKS modules 4–11 produce the full length LTM polyketide backbone, which is finally cyclized and released as the nascent PKS product **14**. Given the structural similarity between **1** and **2** (Figure 1) and the sequence and architectural homology between the LTM and iso-MGS AT-less type I PKSs (Figure 3, Table 1, and Figure S2 of the Supporting Information), it is therefore remarkable that the two PKSs synthesize distinct nascent polyketides, **14** for the LTM type I PKS and **12** and **13** for the iso-MGS type I PKS (Figure 3B and Figure S2C of the Supporting Information). Thus, on the basis of the collinear model for type I PKSs, the LTM PKS would minimally lack a MT domain in module-5, a KR, a DH, and an enoylreductase (ER) domain in module-8, and a KR and ER domain in module-10. In contrast, the iso-MGS PKS would minimally lack a DH domain in module-4, a MT domain in module-5, a KR domain in module-8, and a KR and ER domain in module-10. Intriguingly, while the loading module and modules 2–7 of the LTM and iso-MGS PKSs are essentially identical, the LTM PKS affords a single polyketide product with a hydroxyl group at C-17 and the iso-MGS PKS yields two products, the major one of which features a double bond at C-16 and C-17. There are also distinct domain architectural differences between module-8 and -9 for the LTM and iso-MGS PKSs; however, how these differences account for the structural variations in the resultant polyketide products in **1** and **2** biosynthesis remains unknown (Figure 3A,B and Figure S2B,C of the Supporting Information). AT-less type I PKSs are rich in unusual domains and PKS architecture, and they often violate the colinearity rule most known for type I PKSs.^{44,45} Comparative studies of the LTM and iso-MGS PKSs, as well as PKSs that direct the biosynthesis of other glutarimide-containing polyketides,^{1,2,10,11,27,28} should provide opportunities to probe some of these properties, for example, how AT-less type I PKSs select and control reductive modifications during polyketide chain elongation.

Accumulation of **14** as the sole metabolite in SB15002 (Figure 2B, trace V) and restoration of the production of **1** and the two minor metabolites **13** and **15** in SB15003 (Figure 2B, trace VI) suggested that the LtmK P450 enzyme acts as a desaturase that catalyzes regio- and stereospecific desaturation of **14** to **1**, with **13** and **15** as minor products (Figure 3B). P450 enzymes that act as desaturases are known, but the majority are associated with liver or lung microsomes that catalyze various metabolic biotransformation events in eukaryotes.^{46–48} Only a few P450 desaturases of bacterial origin have been described,^{49–52} for example, CYP102A1 of *Bacillus megaterium*,^{49–51} CYP101A1 of *Pseudomonas putida*,⁴⁹ and CYP199A4 of *Rhodospseudomonas palustris* HaA2.⁵² P450-catalyzed desaturation proceeds via abstraction of hydrogen from the substrate to yield transient free radical intermediates, which undergo elimination of a hydrogen radical to afford the olefin product or recombination with a hydroxyl radical to form the hydroxylated products. Regardless of a prokaryotic or eukaryotic origin, all known examples of P450-catalyzed

desaturation are accompanied by hydroxylation, with hydroxylation generally being the major course of the reaction; the P450 desaturases do not convert the hydroxylated products into olefins.⁵³ It is therefore highly significant that, under the fermentation conditions tested, LtmK catalyzes the desaturation of **14** to **1** with minimal formation of the hydroxylated products, **13** and **15** (Figure 3B). Mechanistic and structural characterization of LtmK may therefore reveal new insights into how P450 desaturases control the fate of the initial radical intermediate in partitioning between the elimination and hydroxylation pathways.

Compared to LtmK, the MgsK P450 enzyme in **2** biosynthesis catalyzes the hydroxylation at position C-8, which is further O-methylated by the MgsJ methyltransferase to furnish the methoxyl group found at C-8 in **2**. Both MgsK and MgsJ have broad substrate promiscuity, and exploitation of the tailoring steps in **1** and **2** biosynthesis therefore provides an attractive opportunity to engineer novel glutarimide-containing polyketide natural products. Indeed, introduction of *mgsK* and *mgsJK* into the $\Delta ltmK$ mutant strains resulted in the production of two novel analogues **16** and **17**, featuring the 17R-hydroxyl group as in **1** and 8R-hydroxyl or 8R-methoxyl group as in **2** (Figure 1C), a combination of structural features not seen in the natural congeners of **1** and **2**. Cytotoxicity assays of **16** and **17** against select cancer cell lines, in comparison with those of **1**, **2**, and specific natural congeners, allowed us to expand the emerging SAR of the glutarimide-containing polyketide family of natural products. Given their fascinating modes of action as inhibitors of tumor cell migration and eukaryotic protein translation and the potent cytotoxicity of **1** and **2** against cancer cell lines, these findings lay the foundation for facilitating the design, development, and preparation of additional analogues of **1** and **2** for anticancer drug discovery and development.

■ ASSOCIATED CONTENT

● Supporting Information

Strains and plasmids (Table S1), summary of the primers used in this study (Table S2), ¹H and ¹³C NMR data for **16** and **17** (Table S3), highlighting of **2** as the true natural product of *S. platensis* NRRL 18993 and **3** and **4–6** as shunt metabolites of **2** following H₂O-mediated ring expansion and ring-opening rearrangements, respectively (Figure S1), a comparison of the biosynthesis of **1** in *S. amphibiosporus* ATCC 53964 and **2** in *S. platensis* NRRL 18993 (Figure S2), studies confirming that the cloned *ltm* cluster directs **1** biosynthesis in *S. amphibiosporus* ATCC 53964 by gene inactivation of *ltmE*, *ltmH*, and *ltmK* (Figure S3), the construction of the *mgsK* and *mgsJK* expression plasmids used to produce **16** and **17** in recombinant strains SB15006 and SB15007 (Figure S4), summary of the cytotoxicity assay data for **16** and **17** against three cancer cell lines in comparison with those of **1**, **2**, and selected native congeners (Figure S5), and ¹H and ¹³C NMR spectra of **14**, **16**, and **17** (Figures S6–S10). This material is available free of charge via the Internet at <http://pubs.acs.org>.

Accession Codes

The GenBank accession number for the *ltm* gene cluster from *S. amphibiosporus* ATCC 53964 is GQ274954.

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■ ABBREVIATIONS

ACP, acyl carrier protein; AMT, amidotransferase; AT-less type I PKS, acyltransferase-less type I polyketide synthase; B, branching; DGN, dorrigin; DH, dehydratase; ER, enoylreductase; GCG, Genetic Computer Group; GTI-seq, Global Translation Initiation Sequencing; HRESI-MS, high-resolution electrospray ionization mass spectrometry; iso-MGS, isomigrastatin; KR, ketoreductase; KS, ketosynthase; LB, lysogeny broth; LTM, lactimidomycin; MGS, migrastatin; MT, methyltransferase; NCBI, National Center for Biotechnology Information; ori-T, origin of transfer; SAM, S-adenosylmethionine; SAR, structure–activity relationship; TE, thioesterase; TSB, tryptic soy broth.

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