

Unveiling the Post-PKS Redox Tailoring Steps in Biosynthesis of the Type II Polyketide Antitumor Antibiotic Xantholipin

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SUMMARY

Xantholipin from *Streptomyces flavogriseus* is a curved hexacyclic aromatic polyketide antitumor antibiotic. The entire 52 kb xantholipin (*xan*) biosynthetic gene cluster was sequenced, and bioinformatic analysis revealed open reading frames encoding type II polyketide synthases, regulators, and polyketide tailoring enzymes. Individual in-frame mutagenesis of five tailoring enzymes lead to the production of nine xantholipin analogs, revealing that the xanthone scaffold formation was catalyzed by the FAD binding monooxygenase XanO4, the δ -lactam formation by the asparagine synthetase homolog XanA, the methylenedioxy bridge generation by the P450 monooxygenase XanO2 and the hydroxylation of the carbon backbone by the FAD binding monooxygenase XanO5. These findings may also apply to other polycyclic xanthone antibiotics, and they form the basis for genetic engineering of the xantholipin and similar biosynthetic gene clusters for the generation of compounds with improved antitumor activities.

INTRODUCTION

Polycyclic xanthone antibiotics feature an angular fused hexacyclic skeleton with a xanthone structure (orange in Figure 1). They are active against fungal or Gram-positive bacteria, including methicillin-resistant *Staphylococcus aureus* and vancomycin-resistant *Enterococcus faecalis* (Peoples et al., 2008), or against several cancer cell lines (Kunimoto et al., 2003). Xantholipin (**1**) is a polycyclic xanthone antibiotic produced by *Streptomyces flavogriseus*. It possesses potent cytotoxicity against the leukemia cell line HL60 (IC₅₀ < 0.3 μ M) and the oral squamous carcinoma cell line KB (IC₅₀ < 2 nM; Terui et al., 2002), and it was recently identified as a lead drug for the treatment of fibrotic diseases (Yuichi et al., 2003). The total chemical synthesis of another polycyclic xanthone antibiotic, FD-594, has been achieved, but it required many steps and did not produce sufficient amounts for medical testing (Masuo et al., 2009). Genetic engineering of the gene cluster may be suitable for this purpose.

Biosynthetic studies and isotope-labeled precursor feeding experiments of polycyclic xanthone antibiotics, including simao-micin α , lysolipin I, and FD-594, proved that these molecules are derived from a single polyketide chain, which is synthesized by a type II polyketide (iterative) synthase (PKS) composed of three monofunctional peptides (Bisang et al., 1999; Bockholt et al., 1994; Carter et al., 1989; Kondo et al., 1998; Kudo et al., 2011; Lopez et al., 2010). The polyketide chains then undergo various post-PKS modifications to produce a variety of the polycyclic xanthone antibiotics. Of particular interest are the formation of the xanthone and the methylenedioxy bridge. Studies on lysolipin I indicated that a Baeyer-Villiger (ketone \rightarrow ester) oxidation is involved in the formation of the xanthone structure, because the single xanthone oxygen originates from molecular oxygen. Also, the two oxygens of the methylenedioxy group are from molecular oxygen and the carbon 27 in between originates from methionine (Bockholt et al., 1994). However, no microbial enzymes for the formation of the two structure features have been identified.

Here we report the cloning and sequencing of the entire biosynthesis gene cluster of **1**. Metabolite analysis of five blocked pathway mutants identified a FAD binding monooxygenase involved in the xanthone formation, a P450 monooxygenase for the generation of the methylenedioxy bridge, an asparagine synthetase homolog for amide ring formation, and a putative polyketide hydroxylase for the C4 hydroxylation, resulting in the proposal of a possible biosynthetic pathway for **1**. Additionally, we characterized two precursors of **1** chemically and for their anticancer activities.

RESULTS AND DISCUSSION

Identification, Cloning, and Sequencing of the *xan* Gene Cluster and Determination of Its Boundaries

Xantholipin (**1**) is similar to lysolipin (LLP) for which the complete biosynthesis gene cluster has been recently sequenced (Lopez et al., 2010). Using conserved regions of the *lIpH* halogenase, we designed primers for the polymerase chain reaction (PCR) amplification of a 300 bp fragment from *S. flavogriseus*, the producer of **1**. The predicted amino acid sequence of the PCR fragment was 73% identical to that of *lIpH*, suggesting that we may have amplified the halogenase gene, *xanH*, from the *xan* gene cluster. These primers were then used to screen a *S. flavogriseus* fosmid library (3,000 clones; 36 kb average

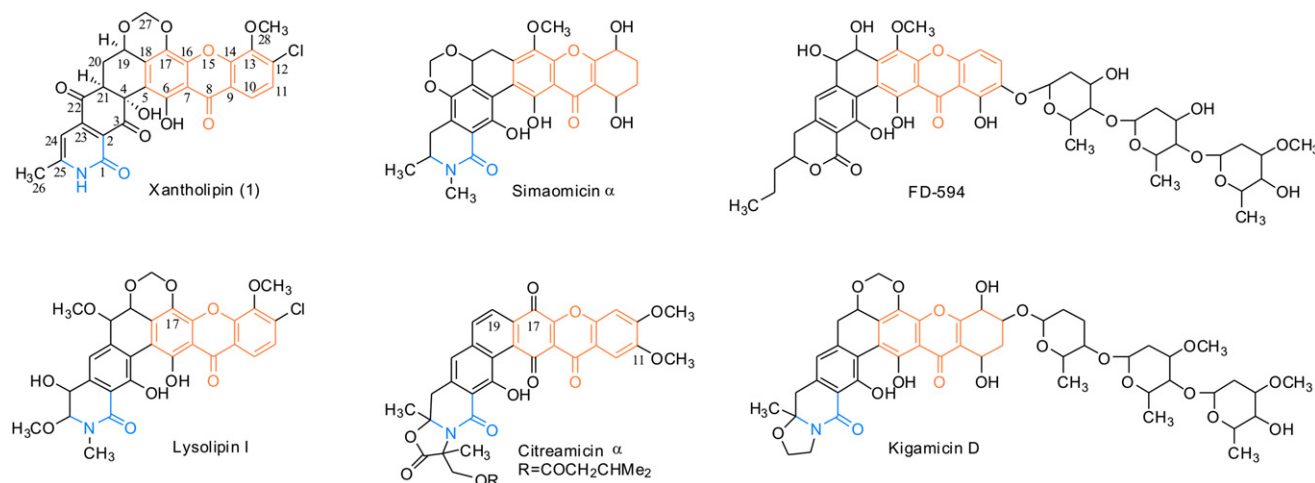


Figure 1. Structures of Xantholipin and Derivatives

Orange, xanthone; blue, cyclic amide.

insert size) using sib selection (McCormick, 1987). Sixteen overlapping fosmid clones were identified. To determine the relevance for the biosynthesis of **1**, a 40 kb DNA fragment was deleted from the *S. flavogriseus* genome by gene replacement using the remains of fosmid 28B8 after *Xho*I digestion and religation (Figure 2; Figure S1A available online). Two independently generated mutants no longer produced **1**, suggesting that the *xan* genes may have been deleted (Figure S1D). Fosmid 22E4, containing a central halogenase gene, was sequenced. It contained several putative antibiotic tailoring genes but, surprisingly, no PKS gene. Chromosome walking using PCR probes 2 and 3 (Figure 2, top) resulted in a minimal 92 kb contig flanked by fosmids 26G2 and 12A9. The 92,445 bp sequence contained 49 open reading frames (ORFs) that were likely to be involved in the biosynthesis of **1** (Figure 2 and Table 1). This was based on functional assignment (XanY1–Y5) and the presence of likely orthologs in the *llp* (lysolipin) and *pnx* (FD-594) gene clusters (Kudo et al., 2011; Lopez et al., 2010). Deletion of ORF1–5 or ORF6–9 from the *S. flavogriseus* genome produced normal looking colonies that still produced **1** (Figures 2 and S1), indicating that these ORFs were located outside the *xan* cluster.

Putative functions were assigned to most of the remaining 49 ORFs as shown in Figure 2 and Table 1. Six *xan* genes (black) encode three cyclases (C1–C3) and a type II minimal PKS consisting of the ketosynthase KS_{α} (XanF), chain length factor (CLF) KS_{β} (XanE), and acyl carrier protein (ACP; XanD). Five dispersed genes (green) were suspected for post-PKS tailoring: a halogenase (XanH) for the chlorination of C12, an asparagine synthetase homolog (XanA) for the formation of the amide ring, two *O*-methyltransferases (XanM1 and XanM2) for the methylation of the C13 and C17 hydroxyl groups, and an additional unassigned methyltransferase (XanM3). Consistent with the highly oxidized structure of **1**, 19 redox genes (pink and red) were identified. In addition to the biosynthetic machinery, five genes (*xanY1–Y5*, dark blue; not present in the *llp* and *pnx* gene clusters) are for the *S*-adenosyl methionine (SAM) recycling, and three genes (*xanB1–B3*, light blue; also without orthologs in the *llp* or *pnx* clusters) are for the synthesis of

malonyl-CoA extender units. There are also three putative regulatory and three resistance genes (yellow) and five ORFs without known functions (gray, *xanJ*, *xanP*, *xanG*, *xanL*, and *xanW*), also without orthologs in the *llp* or *pnx* clusters.

Biosynthesis of the δ -Lactam

Most polycyclic xanthones feature a cyclic amide, which is otherwise rare among aromatic polyketides (blue in Figure 1; FD-594 has a δ -lactone instead). In addition, there are various substituents on the amide nitrogen in the different polycyclic xanthones, such as a methyl group in lysolipin, an amino group in simaomicin, and more complicated heterocyclic structures in citreamicin α and Kigamicin D. The *xan* gene cluster contains an asparagine synthetase homolog, XanA, which strongly resembles the class II asparagine synthetase homolog LlpA (lysolipin) and the amide synthetase FdmV (Chen et al., 2010; fredericamycin A, not a xanthone), including the conserved active site and ligand binding site residues Arg50, Glu80, Asp103, Ser264, and Gly376 (Larsen et al., 1999). FdmV has been characterized to catalyze the formation of a hexacyclic amide, which is similar to the amide ring of **1**. XanA probably converts the polyketide-derived C1 carboxyl group to an amide, followed by the possibly spontaneous elimination of water to form ring F (Figure 3, conversion of compound **8a** to **9**).

To test this hypothesis in vivo, *xanA* was replaced by an apramycin resistance gene in the *S. flavogriseus* genome (Figure S1E). The $\Delta xanA$ mutant accumulated compound **8a** containing a chlorine and a xanthone structure but was devoid of nitrogen (Figures 3 and 4). From 14 l fermentation broth, 85 mg of compound **8a** was purified. ESI-Q-TOF MS, 1D and 2D nuclear magnetic resonance (NMR) spectra (^1H , ^{13}C , DEPT, HMBC, HSQC), provided the molecular formula $\text{C}_{26}\text{H}_{19}\text{ClO}_9$ and the structure (Figures 3, 4, S2, and S3). Different from our expectation, there was no spontaneous cyclization producing the α -pyrone derivative of **8a**; instead, a carboxyl group remained at C1, which was probably a more suitable substrate for XanA. The purified compound **8a** was then crystallized and X-ray diffraction was performed to further confirm the position

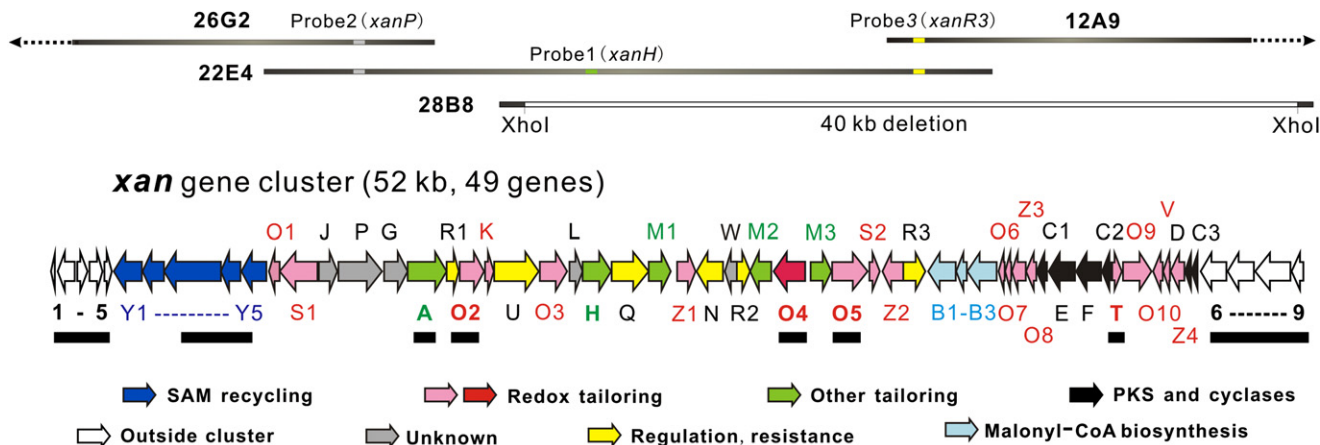


Figure 2. The Organization of the Xantholin Biosynthetic Gene Cluster

The gene cluster spans 52 kb and contains 49 ORFs. Genes are organized according to their proposed roles (see also Table 1); the underlined genes were individually inactivated, except for *orf1*–*5*, *orf6*–*9*, and *xanY3*–*Y5*, which were deleted together. The 40 kb gray part of fosmid 28B8 is the region that was deleted in strain ZWK1. The construction and verification of the deletion mutants are shown in Figure S1.

of chlorine at C12 of **8a**. The X-ray spectra clearly revealed the chlorine bound to C12; surprisingly, a lactone derivative **8b** was generated during the crystallization of **8a** from methanol (Figures 3 and S2).

Since *xanA* is within a group of 14 genes that might form an operon, inactivation of downstream genes by polar effects on transcription might have contributed to the phenotype of the $\Delta xanA$ mutant. Complementation expressing *xanA* from the high copy vector pJTU1278 partially (40%) restored production of **1** (Figure S1P), suggesting that deletion of *xanA*, and not inactivation of a downstream gene, was responsible for the production of **8a**. Therefore, XanA was required for the amidation of **8a**.

Bioinformatic Search for Genes for the Baeyer-Villiger Oxidation

We next addressed the biosynthesis of the xanthone ring, which is the characteristic feature of all polycyclic xanthone antibiotics and has not been elucidated genetically before. Studies on lysolipin I indicated that a Baeyer-Villiger oxidation was involved in the formation of xanthone (Bockholt et al., 1994), and epoxidation followed by a Baeyer-Villiger oxidation was proposed to be involved in the xanthone formation of the polycyclic xanthone antibiotic FD-594 (Kudo et al., 2011). Therefore, we selected four redox enzymes (O2, O4, O5, and T), which had orthologs in the FD-594 and lysolipin gene clusters, as possible candidates that might catalyze these reactions or, as we found out later, were involved in other steps for the biosynthesis of **1**.

XanO4 was between 29% and 31% identical to the atypical Baeyer-Villiger monooxygenases (BVMOs), including MtmOIV (mithramycin), JadH (jadomycin), and GilOI (gilvocarcin), all of which lack the typical sequence motif of type I BVMOs and are not related to the type II BVMOs (Beam et al., 2009; Gibson et al., 2005; Kharel et al., 2007; Rix et al., 2005). The similarities included all three conserved motifs (GxGxxG for binding the ADP part of FAD, the sequence DG for binding the pyrophosphate of FAD or NADPH, and GD for binding the riboflavin of FAD). XanO4 also resembled the atypical BVMO GrhO5 (42% identity) which,

together with GrhO1, has been characterized through in vivo experiments to catalyze three C–C bond cleavages for the formation of the spiroketal center of griseorhodin (Yunt et al., 2009).

Phylogenetic analysis of putative BVMOs showed two groups, the type I BVMOs, and the atypical BVMOs, three of which (XanO4, LlpOVIII, and PnxO4) were involved in the biosynthesis of polycyclic xanthone antibiotics (Figure S4B). Interestingly, these three proteins, together with GrhO5, formed a distinct subgroup, separated from the group containing MtmOIV (mithramycin). These sequence comparisons suggested that XanO4 may catalyze the formation of xanthone, using a mechanism similar to that of the atypical BVMO GrhO5. However, no homolog of GrhO1 was encoded in the *xan* cluster; therefore, XanO4 may act alone or with other unidentified protein.

XanO2 was the only P450 monooxygenase in the *xan* gene cluster. Several plant P450 monooxygenases have been characterized to catalyze the formation of methylenedioxy bridges between a methoxy group and a nearby hydroxyl group (Ikezawa et al., 2007; Ono et al., 2006). XanO2 was thus a candidate for methylenedioxy bridge formation, except that FD-594, which does not have a methylenedioxy bridge, also contained P450 homologs. Alternatively, since many P450s catalyze epoxidations (Guengerich, 2003), XanO2 might also catalyze the epoxidation for xanthone formation.

XanO5 was similar to TcmG and GrhO9, which have been proved to be involved in the tailoring hydroxylation of the aromatic type II polyketide antibiotics tetracenomycin and griseorhodin, respectively (Decker et al., 1993; Yunt et al., 2009). XanO5 may thus be responsible for hydroxylating a precursor of **1** or even be an alternative to XanO4 for xanthone formation (XanO5 and XanO4 are 41% identical).

XanT, from the SchA/CurD superfamily, was conserved in the three polycyclic xanthone antibiotic gene clusters but was rare among the other aromatic PKS gene clusters. The only XanT homolog was FdmK, which was proposed to be involved in C–C bond cleavage for the formation of the spiro carbon center of fredericamycin (Wendt-Pienkowski et al., 2005). XanT

contains a conserved NAD(P) binding site like XanO4, suggesting they use the same cofactor. XanT was proposed to be the partner protein supporting XanO4, but inactivation of *xanT* in *S. flavogriseus* did not cause accumulation of **6** (see below).

S. Flavogriseus Mutant Derivatives Lacking Redox Genes

The four genes *xanO2*, *xanO4*, *xanO5*, and *xanT* (Figure 2, underlined with black bars) had characteristics that suggest the likely involvement in xanthone formation. They were deleted separately in *S. flavogriseus* and replaced by an apramycin resistance cassette (Figure S1). LC-MS showed that each of the four mutants accumulated compounds that were different from **1** (Figure 4). Q-TOF MS was used to determine the molecular formulae of the main components in each extract (Figures 4 and S3).

The XanO4 blocked mutant accumulated large amounts of the red (483 nm) compound **6** (**1** was yellow). By optimizing the fermentation conditions and the isolation procedure, we obtained 48.5 mg of **6** from a 7 l culture of the $\Delta xanO4$ strain. Evaluation of the ^1H , ^{13}C , DEPT, HMBC, HSQC NMR spectra (Figure S2 and Table S1) produced the structure of **6** (Figure 3). The NMR spectra resembled those of **8a**, but the signals for the ether bond of xanthone were missing, and a methoxylated anthraquinone was identified instead. (Note that C15 of **6** is replaced by O in **7**.)

The $\Delta xanO2$, $\Delta xanO5$, and $\Delta xanT$ mutants produced insufficient amounts of compounds for NMR analysis, and it was interesting to note that all of the three mutants accumulated more than one product, which implied that the primary reaction products were probably metabolized by other enzymes or underwent spontaneous reactions. XanT and XanO4 probably acted first, because the products from the deletion mutants did not contain chlorine or nitrogen. XanO2 and XanO5 probably performed late steps, because the accumulated products contained one chlorine and one nitrogen, like compound **1** (Figures 3 and 4). From the high-resolution molecular masses, the proposed functions of XanO2 and XanO5 in the biosynthetic pathway of **1** (see above), and the structures of **8a/8b** and **1**, compounds **11a** (accumulated by the $\Delta xanO2$ mutant) and **15a** (accumulated by the $\Delta xanO5$ mutant), probably had the structures shown in Figure 3. Structure **7** was proposed from the comparison of the known structures **6** and **8a/8b**, and **9** was proposed from the comparison between **8a/8b** and **1**. The proposed structures of the other compounds are explained below.

Introducing an integrative plasmid expressing *xanO2* from the *ermE*^{*} promoter (McDaniel et al., 1993) into the $\Delta xanO2$ mutant only partially restored **1** production (Figure S1Q), possibly because the promoter was not suitable for XanO2 expression in *S. flavogriseus* or because the expression of the downstream *xan* genes was reduced.

XanO4 was purified from an *E. coli* clone. In the presence of XanO4, it oxidized NADPH to NADP and transferred the electrons to FAD. Compound **6**, the proposed substrate of XanO4, was not converted to compound **7** using NAD(P)H and FAD as cofactors (Figure S5). Therefore, XanO4 probably requires a partner protein to convert **6** to **7**. Unfortunately, XanT, the proposed partner protein of XanO4 (see above), could not be obtained in soluble form for enzymatic testing.

Bioactivity of Xantholin and Putative Biosynthetic Intermediates

To determine how the oxidative modification during **1** biosynthesis affects bioactivity, we performed in vitro cytotoxicity assays of **6**, **8a**, **1**, and doxorubicin as a positive control. Table 2 shows that **1** was 10-fold more effective against a lung cancer cell line, 3-fold more effective against a colon cancer cell line, and equally effective against a leukemia than the commonly used anticancer drug doxorubicin. The precursors **6** and **8a** were, however, about 10^2 - to 10^3 -fold less potent. These data indicated that the post-PKS modifications were important for bioactivity.

Proposal for a Biosynthetic Pathway for **1**

The proposed biosynthesis pathway of **1** (Figure 3) probably begins with the synthesis of malonyl-CoA from carboxylation of acetyl-CoA by the enzymes XanB1–B3 (probable acetyl-CoA carboxylase; Figure 2, light blue arrows; Table 1). These enzymes probably duplicate enzymes from the primary metabolism, which supply malonyl-CoA for fatty acid biosynthesis. Similar clusters occur in other antibiotic biosynthesis gene clusters, including that of pradimicin (Kim et al., 2007), but interestingly, there are no orthologs in the *llp* and *pnx* gene clusters.

Condensation of 13 malonate units by the minimal type II PKS (XanDEF; acyl carrier protein, ACP; chain length factor, CLF or KS $_{\beta}$; ketosynthase, KS $_{\alpha}$) probably produces the hypothetical, enzyme-bound, and fully oxidized polyketide **2**. The three PKS subunits contained all the conserved active-site residues (Bisang et al., 1999). A phylogenetic analysis of KS $_{\beta}$ s of type II PKSs showed that, as expected for the C26 polyketide chain of **1**, XanE clustered with the longer chain (C24–30) KS $_{\beta}$ s (Figure S4A). Unusual for type II PKS gene clusters, including *llp* and *pnx*, *xanD* (ACP) was separate from the KS genes.

Three carbonyl reductions at C11, C17, and C19 were postulated to produce the hypothetical compound **3**, based on isotope-feeding experiments of lysolipin and citreamicin (Guy et al., 1991). They are probably performed by the 3-oxoacyl-ACP reductases XanZ3 and XanZ4, which have orthologs in the *llp* and *pnx* gene clusters (about 70% identity) and are similar to the characterized C19 ketoreductase (KR) BenL (benastatin; Lackner et al., 2007). C19 KRs are known exclusively from angucycline antibiotics, and usually there is only a single C19 KR in each cluster (Lackner et al., 2007). The *xan*, *llp*, *pnx*, and *grh* (griemorhodin) gene clusters encode two probable C19 KRs, which were different from most angucycline antibiotic gene clusters (Li and Piel, 2002). Since no C11 KR has been, to our best knowledge, reported before, a phylogenetic analysis of the C17 KRs and the C19 KRs was performed. It revealed that XanZ3 clustered with LlpZl, PnxG, GrhT, GrhO10, and known C19 KRs and XanZ4 clustered with LlpZlll and PnxW but not with known C17 KRs (Xu et al., 2005; Figure 5). XanZ3 was thus the likely C19 KR, and XanZ4, LlpZlll, and PnxW may belong to a new family of C11 KRs, which are exclusive to polycyclic xanthone antibiotics.

Guided by the C11 and C19 hydroxyl groups, the cyclases XanC1–C3 then form five C–C bonds (intramolecular aldol condensations) to form the angular 5-ring aromatic structure **4**. XanC1 resembled the first and second ring cyclase TcmN (tetraenomycin; McDaniel et al., 1995; Shen and Hutchinson, 1996).

Table 1. Deduced Functions of ORFs in the Xantholipin Biosynthetic Gene Cluster

Xan Proteins		Protein Homologs	
Gene (aa)	Proposed Function	Protein, Origin	Identity/Similarity (%)
<i>xanY1</i> (472)	Adenosylhomocysteinase	SSDG_05037, <i>S. pristinaespiralis</i>	95/97
<i>xanY2</i> (300)	Methylenetetrahydrofolate reductase	SSDG_05038, <i>S. pristinaespiralis</i>	84/89
<i>xanY3</i> (1165)	Methionine synthase	SSCG_05781, <i>S. clavuligerus</i>	89/93
<i>xanY4</i> (330)	Carbohydrate kinase	SsfX2, <i>S. sp. SF2575</i>	70/79
<i>xanY5</i> (406)	SAM synthetase	SSCG_03814, <i>S. clavuligerus</i>	85/91
<i>xanO1</i> (145)	Monooxygenase	SanK, <i>S. sp. SANK 61196</i>	33/51
<i>xanS1</i> (407)	Dehydrogenase	LipS, <i>S. tendae</i>	59/74
<i>xanJ</i> (242)	Unknown	ORF6, <i>S. antibioticus</i>	42/56
<i>xanP</i> (539)	Unknown	SBI_09782, <i>S. bingchenggensis</i>	69/78
<i>xanG</i> (391)	Unknown	LipG, <i>S. tendae</i>	60/71
<i>xanA</i> (614)	δ -lactam formation	LipA, <i>S. tendae</i>	72/80
<i>xanR1</i> (227)	Transcriptional regulator	LipRIII, <i>S. tendae</i>	49/61
<i>xanO2</i> (400)	Cytochrome P450	LipOIV, <i>S. tendae</i>	73/84
<i>xanK</i> (89)	3Fe-S ferredoxin	LipK, <i>S. tendae</i>	67/77
<i>xanU</i> (709)	Antibiotic transporter	ORF27, <i>S. globisporus</i>	57/73
<i>xanO3</i> (392)	FAD-binding monooxygenase	Krac_4468, <i>Ktedonobacter racemifer</i>	44/59
<i>xanL</i> (186)	Unknown	BSG1_19969, <i>Bacillus sp. SG-1</i>	33/52
<i>xanH</i> (471)	Halogenase	LipH, <i>S. tendae</i>	70/85
<i>xanQ</i> (533)	Transporter	FdmT, <i>S. griseus</i>	46/60
<i>xanM1</i> (339)	O-methyltransferase	LipMIV, <i>S. tendae</i>	63/72
<i>xanZ1</i> (285)	Reductase	LipZII, <i>S. tendae</i>	59/73
<i>xanN</i> (431)	Antiporter	Veg3, uncultured soil bacterium	57/76
<i>xanW</i> (162)	Unknown	SACTE_4627, <i>S. sp. SirexAA-E</i>	48/67
<i>xanR2</i> (153)	Transcriptional regulator	PnxR1, <i>S. sp. TA-0256</i>	34/53
<i>xanM2</i> (340)	O-methyltransferase	LipMIII, <i>S. tendae</i>	63/74
<i>xanO4</i> (540)	FAD-binding monooxygenase	LipOVIII, <i>S. tendae</i>	70/81
<i>xanM3</i> (344)	O-methyltransferase	LipMII, <i>S. tendae</i>	64/77
<i>xanO5</i> (531)	FAD-binding monooxygenase	GrhO9, <i>S. sp. JP95</i>	43/58
<i>xanS2</i> (228)	Dehydrogenase	LipU, <i>S. tendae</i>	56/69
<i>xanZ2</i> (302)	Oxidoreductases	SSAG_00456, <i>S. sp. Mg1</i>	80/87
<i>xanR3</i> (335)	Transcriptional regulator	SACTE_0424, <i>S. sp. SirexAA-E</i>	83/87
<i>xanB1</i> (447)	Biotin carboxylase	PdmP1, <i>Actinomadura hibisca</i>	63/74
<i>xanB2</i> (174)	BCCP	PdmP2, <i>Actinomadura hibisca</i>	51/59
<i>xanB3</i> (564)	Carboxyl transferase	ZhuF, <i>S. sp. R1128</i>	59/68
<i>xanO6</i> (103)	Monooxygenase	LipOII, <i>S. tendae</i>	59/68
<i>xanO7</i> (115)	Monooxygenase	LipOIII, <i>S. tendae</i>	58/71
<i>xanZ3</i> (250)	3-oxoacyl-ACP reductase	LipZI, <i>S. tendae</i>	70/83
<i>xanO8</i> (153)	Monooxygenase	LipB, <i>S. tendae</i>	65/77
<i>xanC1</i> (155)	Cyclase	RubF, <i>S. collinus</i>	68/78
<i>xanE</i> (428)	Chain length factor	LipE, <i>S. tendae</i>	68/77
<i>xanF</i> (425)	Ketosynthase	LipF, <i>S. tendae</i>	80/89
<i>xanC2</i> (144)	Cyclase	LipCII, <i>S. tendae</i>	72/84
<i>xanT</i> (128)	CurD homolog	LipT, <i>S. tendae</i>	71/84
<i>xanO9</i> (472)	Oxidase	Aur2I, <i>S. aureofaciens</i>	47/62
<i>xanO1</i> (148)	Monooxygenase	LipQ, <i>S. tendae</i>	61/72
<i>xanV</i> (126)	CurD homolog	LipV, <i>S. tendae</i>	72/84
<i>xanZ4</i> (251)	3-oxoacyl-ACP reductase	LipZIII, <i>S. tendae</i>	66/75
<i>xanD</i> (86)	ACP	SsfC, <i>S. sp. SF2575</i>	48/67
<i>xanC3</i> (112)	Cyclase	LipCIII, <i>S. tendae</i>	69/84

Table 1. Continued

Xan Proteins		Protein Homologs	
Gene (aa)	Proposed Function	Protein, Origin	Identity/Similarity (%)
<i>orf1</i> (71)	AbaA-like protein	SGR_1054, <i>S. griseus</i>	85/97
<i>orf2</i> (286)	DNA-binding	SGR_1055, <i>S. griseus</i>	83/89
<i>orf3</i> (163)	Regulatory protein	SGR_1056, <i>S. griseus</i>	51/65
<i>orf4</i> (543)	Unknown	SGR_1057, <i>S. griseus</i>	64/72
<i>orf5</i> (97)	Unknown	SGR_880, <i>S. griseus</i>	54/73
<i>orf6</i> (647)	Ketoacyl-ACP synthase	PctM, <i>S. pactum</i>	44/53
<i>orf7</i> (462)	Acyl-CoA synthetase	KRH_03520, <i>Kocuria rhizophila</i>	34/51
<i>orf8</i> (566)	Acetolactate synthase	Trd_0894, <i>Thermomicrobium roseum</i>	35/46
<i>orf9</i> (220)	Ketoacyl-ACP synthase	PctM, <i>S. pactum</i>	45/61

XanC1 may thus form the first two bonds at C9–C14 and C7–C16 in **1**. XanC3 was very similar to TcmI and JadI (jadomycin), which have been proved to be responsible for “kinked ring” cyclization (Shen and Hutchinson, 1993), and may thus form the bond between C4 and C21 for ring E of compound **4**. XanC2 resembled several cyclases (RubD, GrhS, FdmD, and TcmJ), which

may increase the yields of cyclized products by stabilizing the complex composed of the minimal PKS (Bao et al., 1998). There are no clues about which enzymes may perform the third and fifth cyclizations to form compound **4**.

Based on the proven structure of compound **6** (see below), the pentacyclic compound **4** needs to undergo three hydroxylation

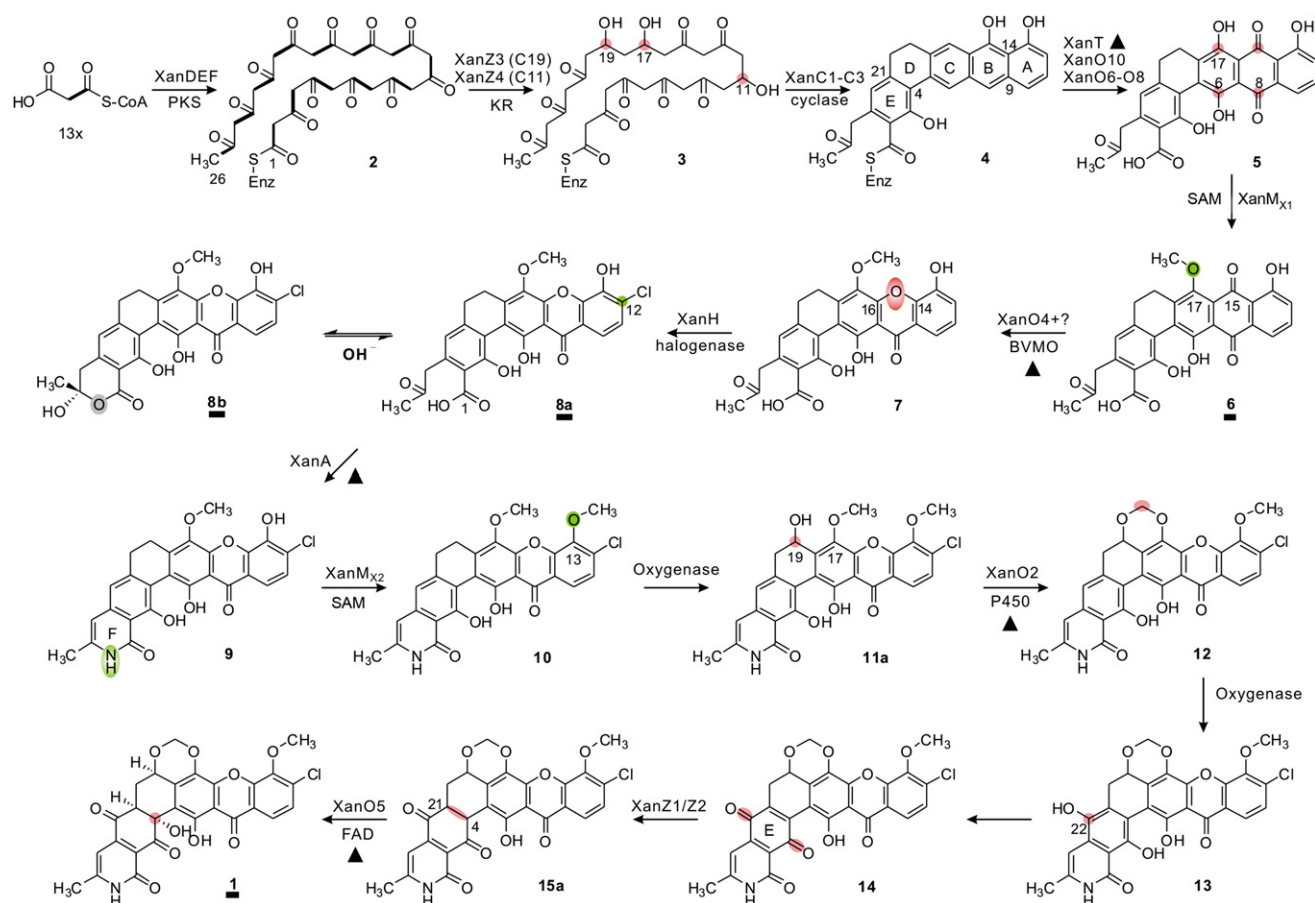


Figure 3. Proposed Xantholinin Biosynthetic Pathway

Black underline, structure confirmed using NMR or X-ray crystallography as shown in Figure S2; black triangle, gene deleted in producer strain; shading, latest chemical change. See also Figure S6.

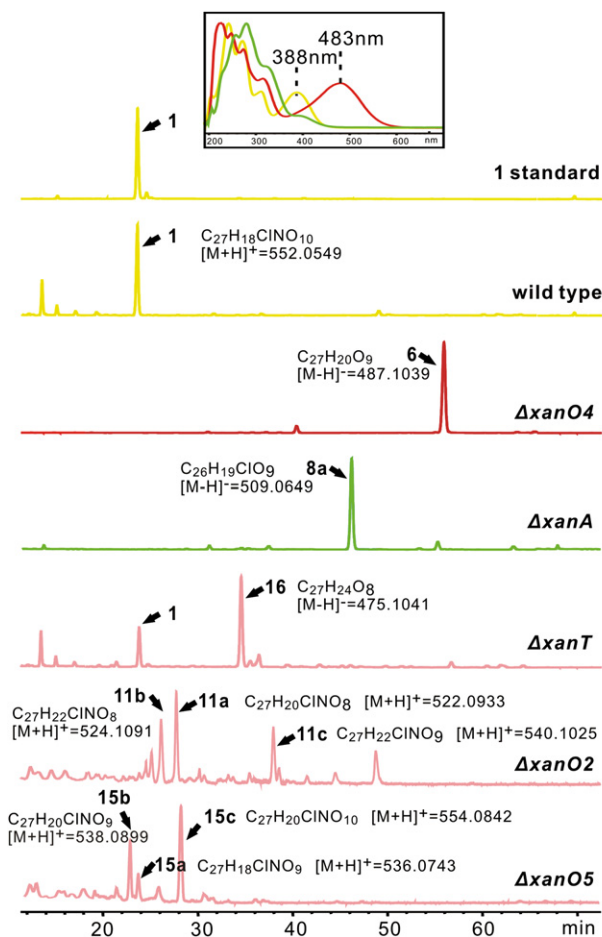


Figure 4. Analysis of Antibiotic Production by *S. Flavogriseus* and Its Mutant Derivatives

Top, UV spectra of compounds **1** (xantholipin, yellow), **6** (red), and **8a** (yellow). Bottom, HPLC traces (274 nm UV absorption). The formulas of the compounds were given by Q-TOF Mass shown in Figure S3. Note, **8b** was generated during the crystallization from methanol for X-ray analysis.

See also Figure S5.

steps at C6, C8, and C17, followed by oxidation to yield the anthraquinone structure of **5**, which was probably no longer attached to the ACP. Aromatic type II PKS gene clusters, including *xan*, do not contain a thioesterase, and it is not known how the molecules are released from the ACP (Staunton and Weissman, 2001).

Four small putative antibiotic biosynthesis monooxygenase (ABM) superfamily enzymes, XanO6, XanO7, XanO8, and XanO10, were predicted to be associated with the hydroxylations of the aromatic backbone. XanO8 was similar to FdmM (48% identity), which has been characterized to hydroxylate C6 nearest to the inside bend of fredericamycin (Chen et al., 2009), which corresponds to position C6 in **5**. Therefore, XanO8 probably hydroxylates C6. XanO6 and XanO7 were similar to PdmH, which has been proved to be responsible for quinone formation in the pentangular polyphenol backbone of pradimicin (Zhan et al., 2008). XanO10 was similar to FdmM1 (48% identity) hydroxylating C8 in fredericamycin (Chen et al.,

Table 2. Antitumor Activity of Xantholipin Precursors

Tumor Cell Lines	IC ₅₀ (ng/ml)			
	Doxorubicin	Xantholipin	6	8a
A-549 (lung)	266 ± 46	21 ± 2	3,900 ± 300	8,900 ± 600
HCT-116 (colon)	27 ± 7	7 ± 1	6,700 ± 1,000	11,900 ± 1,400
P388 (leukemia)	11 ± 1	11 ± 1	>20,000	20,000 ± 7,000

2009). The two reactions both correspond to the oxidation at C8 in **5**. XanO6, XanO7, and XanO10 were thus candidates for the hydroxylation at C8 in **4**. The oxidation of **4** at C17 may also be performed by one of the three enzymes.

An additional gene, *xanT* (SchA/CurD superfamily), also encodes a small protein similar to ABM monooxygenase and has homologs in the *llp* and *pnx* clusters. XanT was the possible enzyme that supports XanO4 in the formation of the xanthone structure. Inactivation of *xanT* resulted in the accumulation of a new metabolite **16** but did not abolish **1** production (Figure 4). Q-TOF analysis revealed that **6** and **16** contained the same number of carbons but **16** lacked an oxygen (Figure S3), suggesting that XanT acted on a less oxidized precursor of **6** before the xanthone was synthesized. The residual production of **1** in these mutants might have resulted from nonspecific activities from unidentified endogenous enzymes.

One of the three putative S-adenosyl-L-methionine-dependent methyltransferases (SAM-MTases; XanM1, 2, or 3) then transfers a methyl group to the C17 hydroxyl group of **5** to produce **6**, which was accumulated by the Δ *xanO4* mutant (see above). The hypothetical compound **10** contains two methoxy groups. XanM1 and XanM2 resembled O-methyltransferases, and XanM3 contains motifs for O- and C-methyltransferases. XanM1 and XanM2 were thus likely to be involved in the methylation of the C13 and C17 hydroxyl groups. The fully characterized compounds **6** and **8a** were O-methylated at C17 but not at C13. The C13 O-methylation may thus be specific for the hypothetical compound **9**, which contains the chlorinated xanthone and the δ -lactam.

XanY1–Y5, encoded by successive genes on the left side of the *xan* cluster, are a complete set of enzymes for the recycling of S-adenosylhomocysteine (a by-product in the SAM-dependent methylation reaction) to SAM (Haagen et al., 2006). These five genes are presumably required to ensure the supply of sufficient SAM precursor for the two methylations during the late growth phase, as has been proposed for saframycin biosynthesis (Li et al., 2008). Inactivation of XanY3–Y5 decreased the production of **1** about two-fold, which might have been caused by the insufficient supply of the methyl group from SAM (Figure S1). The residual **1** production probably utilized SAM from the primary biosynthetic pathway.

XanO4, probably together with an unidentified protein, catalyzes xanthone formation resulting in **7** because deletion of XanO4 resulted in the accumulation of **6** (see above). GrhO1 and GrhO5 (similar to XanO4) act together to catalyze three carbon-carbon bond cleavages for the biosynthesis of the

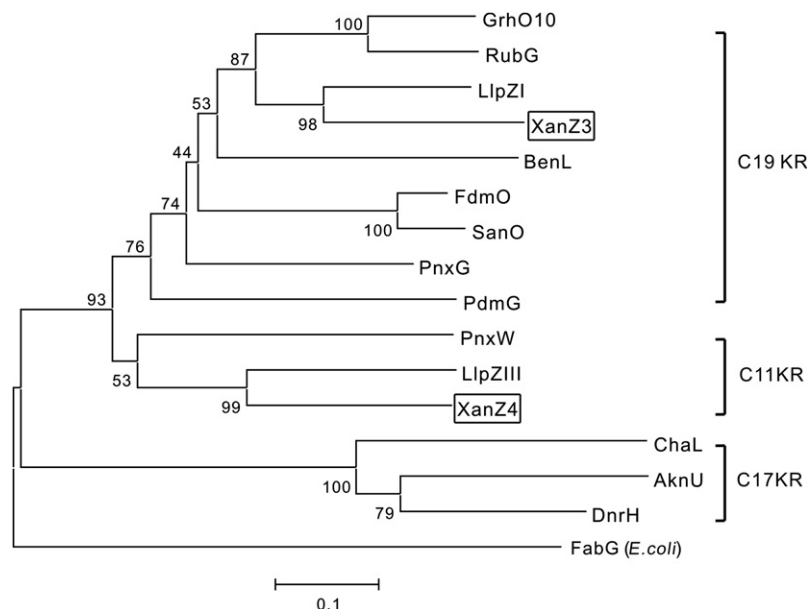


Figure 5. Phylogenetic Analysis of the XanZ3, XanZ4, and Their Homologs

PDB ID codes: LlpZI, CAM34340; GrhO10, AAM33668; RubG, AAG03071; BenL, CAM58804; FdmO, AAQ08926; SanO, ADG86325; PnxG, BAJ52686; LlpZIII, CAM34370; PnxW, BAJ52708; ChaL, CAH10174; AknU, AF264025; DnrH, AAA65202; FabG, EGT67884. See also Figure S4.

8a and the trans-complementation restored **1** biosynthesis. This indicated that XanA was most probably required for the incorporation of the nitrogen and the formation of the characteristic δ -lactam ring to produce **9**. Compound **6**, that accumulated in the $\Delta xanO4$ mutant, was not amidated by XanA, suggesting that the xanthone structure, the chlorine or both may be important for XanA activity (Figure 3). This was in line with FdmV (fredericamycin), which can utilize a wide range of nitrogen sources but is specific for the fredericamycin precursor (Chen et al., 2010). Multiple alignments of

spiroketal polyketide griseorhodin (Yunt et al., 2009). A two enzyme mechanism, including the XanO4 homolog PnxO4, was proposed for the xanthone ring formation for FD-594 (Kudo et al., 2011). Biosynthesis of the other polycyclic xanthone antibiotic lysolipin may thus also implicate a two component system, including the XanO4 homolog LlpOVIII to form the xanthone structure.

The mechanism of xanthone formation remains uncertain. XanO4 may cleave the C14-C15 or the C15-C16 bond. We proposed two alternative paths for the formation of the xanthone scaffold from **6** (Figure S6). The two paths share a Baeyer-Villiger oxidation catalyzed by XanO4, converting the quinone of **6** to a lactone via a “Criegee” intermediate. After that, the lactone may undergo a second round of Baeyer-Villiger oxidation, and the resultant circular carbonate ester is formed, followed by a decarboxylation reaction to produce **7** (path A). Alternatively, the lactone is hydrolyzed and recycled via an intramolecular OH-attack. The resultant intermediate is hydroxylated at C7 followed by a decarboxylation and dehydration to yield **7** (path B). In both cases some additional, unidentified protein or cofactor may be required for the reaction.

Chlorination of **7** at C12 by XanH produces **8a**, which was fully characterized. XanH, the only putative halogenase of the *xan* gene cluster was 70% identical to LlpH (lysolipin biosynthesis), which is a nonheme, flavin-dependent halogenase (Lopez et al., 2010). Enzymes in this class catalyze the formation of carbon-halogen bonds at electron-rich positions, using $FADH_2$, a halide ion (usually Cl^-), and molecular oxygen (Yeh et al., 2007). The reduced flavin is provided by a partner flavin reductase, which could not be located within the sequenced 92 kb region. XanH seemed to be specific for **7** containing the xanthone structure, because it did not chlorinate **6** in the $\Delta xanO4$ mutant.

The next step is the δ -lactam (ring F) formation by the amide synthetase XanA (resembled asparagine synthetase; see above). Inactivation of XanA accumulated an expected carboxylic acid

FdmV homologs revealed that the Cys2, which is indispensable for glutaminase activity, was mutated to Ser in PdmN (pradimicin) and GrhP (griseorhodin), implying that their ability to hydrolyze Gln was lost (Chen et al., 2010). In the case of PdmN, L-Ala was still used as a nitrogen source instead of free NH_3 (Kim et al., 2007). The putative amide synthetases for citreamicin and kigamicin D feature the C2S mutation and may utilize unusual amino acids to form the heterocyclic structures.

C13 O-methylation by XanM1 or XanM2 produces **10**, which is then hydroxylated at C19 to produce **11a**. The identity of the oxidase for the C19 hydroxylation of **10** remains uncertain.

Bioinformatic analysis and the gene replacement experiment (see above) suggested that XanO2, the only P450 monooxygenase in the *xan* gene cluster, may convert **11a** to **12** (methylenedioxy bridge formation). Small amounts of a compound with the same formula as **11a** (calculated 522.0950, observed 522.0933, resulting in the same formula) was isolated from the $\Delta xanO2$ mutant. Two other compounds, **11b** and **11c**, with a UV absorption similar to compound **8a** (380nm), were probably shunt products derived from **11a** with added H_2 or H_2O , respectively. Their structures were not characterized by NMR because of insufficient yield. Compound **11a** was a suitable substrate for methylenedioxy bridge formation. Considering all the information above, XanO2 probably catalyzed a reaction between the C19 hydroxyl group and the C17 methoxy group to form the methylenedioxy bridge of **12**.

Biochemical logic suggested that four steps are needed to convert **12** to **1**; hydroxylation of **12** at C22, followed by formation of the quinone ring E and reduction of the C4-C21 double bond results in **15a**, which was the predicted substrate for C4 hydroxylation.

Hydroxylation at C22 of **12** is similar to the three earlier hydroxylations at C6, C8, and C17 of **4** and was proposed to come after the methylenedioxy bridge formation by XanO2. Analysis mentioned above assigned four possible candidate enzymes for the three hydroxylations. One of the four putative

monooxygenases, XanO6, 7, 8, or 10, may perform this step. No enzyme has been identified for the formation of quinone in ring E.

Prior to the C4 hydroxylation, the double bond between C4 and C21 has to be reduced. Two reductases, other than the 3-oxoacyl-ACP reductase, XanZ1 and XanZ2, are encoded in the *xan* cluster. XanZ1 is homologous to a F420-dependent reductase and XanZ2 belongs to the aldo/keto reductase superfamily, who use soluble NAD(P)H as electron donor. It is still uncertain which one of the two reductases is responsible for the reduction of the C4–C21 double bond in the conversion of **14** to **15a**.

Compound **15a**, accumulated by the $\Delta xanO5$ (polyketide hydroxylase) mutant, possibly has the expected substrate structure for C4 hydroxylation. Compounds **15b** and **15c** were likely shunt products derived from **15a**, with added H₂ or H₂O. Thus, XanO5, was most probably responsible for the C4 hydroxylation of **15a** to complete the biosynthesis of **1**.

SIGNIFICANCE

Xantholipin is a polycyclic xanthone antibiotic representing a class of molecules exhibiting antitumor, antibacterial, and antifungal bioactivity. The biosynthetic pathways for these compounds are of great interest, as they undergo uncommon prearomatic modifications and reveal unusual functional groups. We have identified the biosynthetic gene cluster and its boundaries, which features an unusual C11 ketoreductase for the pre-aromatic reduction of the tridecaketide backbone of xantholipin. We also obtained nine compounds accumulated by blocked pathway mutants, two of which were fully characterized. This gave insight into the biosynthesis of three main structural features of xantholipin, including a FAD binding monooxygenase for the xanthone scaffold, an amide synthetase for the amide bond formation, and, to our knowledge, the first identification of a bacterial P450 monooxygenase for methylenedioxy bridge formation. Characterization of the biosynthetic pathway of xantholipin provides the opportunity to understand unusual modifications involved in the biosynthesis of polycyclic xanthone antibiotics and generate libraries of derivative compounds for structure-activity relationship studies by genetic engineering.

EXPERIMENTAL PROCEDURES

Bacteria Strains, Plasmids, Primers, Culture Conditions, and General Techniques

Bacterial strains and plasmids are listed in Table S1, primers are in Table S2. *Escherichia coli* DH10B, BW25113, and ET12567/pUZ8002 were used for cloning, λ -Red-mediated recombination or, conjugation between *E. coli* and *Streptomyces* spp., respectively (Gust et al., 2003; Paget et al., 1999). *S. flavogriseus* SS101 was the wild-type xantholipin producer used for the generation of mutant strains. ISP-3 solid medium was used for *S. flavogriseus* sporulation and SFM solid medium (Kieser et al., 2000) for conjugation between *E. coli* and *Streptomyces*. Liquid seed medium (per liter: 10 g glucose, 25 g soluble starch, 2 g yeast extract, 5 g fish meal, 3 g pharmanedia, 3 g casein, and 2 g calcium carbonate [pH 7.2]) was used for growing mycelia for the isolation of total DNA and liquid fermentation medium (per liter: 20 g oatmeal, 10 g pharmanedia, 5 g fish meal, 10 g glucose, 25 g dextrin, 2 g yeast extract, and 3 g calcium carbonate [pH 7.2]) was used for xantholipin

production. For *Streptomyces*, thiostrepton and apramycin were used at 12.5 μ g/ml and 15 μ g/ml in solid and liquid media, respectively.

The unstable *Streptomyces* plasmids pHZ1358 (Sun et al., 2009) and its derivative plasmid pJTU1278 were used for gene inactivation. pMD18-T (TaKaRa, Dalian, China) and pBlueScript II SK(+) (Stratagene, La Jolla, CA, USA) were used for DNA sequencing, and pRSET B (Invitrogen, Shanghai, China) was used for heterologous protein expression. Recombinant DNA techniques for *E. coli* and *Streptomyces* were as described (Kieser et al., 2000; Sambrook and Russell, 2000).

S. Flavogriseus Fosmid Library

A library of *S. flavogriseus* DNA in *E. coli* EPI300-T1^R was constructed in the vector pCC1FOS using CopyControl Fosmid Library Production Kit (Epicenter, Madison, WI, USA). Conserved motifs of nonheme halogenases involved in the biosynthesis of lysolipin were identified by sequence alignment and used to design the PCR primers probe1-F/R. The fosmid library was screened by PCR and fosmid 22E4 was isolated. Two more primer pairs, (probe2-F/R, probe3-F/R) were used to isolate fosmids 26G2 and 12A9, which cover the entire xantholipin biosynthesis gene cluster (Figure 2).

Replacements in *S. Flavogriseus*

To genetically inactivate the putative xantholipin gene cluster, fosmid 28B8 was completely digested by *Xho*I and religated to give pJTU3923, which lacks 40 kb from *xanU* to *orf9* (Figure 2, open box). Then a 3.0 kb BamHI fragment from pJTU3923, containing the remains of 28B8, was cloned into the BamHI site of pBlueScriptII SK(+) to give pJTU3852. A 1.4 kb fragment containing *aac(3)IV* was amplified from pHGF9827 using primers ZWK1-F/R and inserted into the unique *Xho*I site of pJTU3852 to produce pJTU3851. Subsequently, the 4.4 kb BamHI fragment from pJTU3851 was inserted into the BamHI site of the unstable *Streptomyces* vector pJTU1278 to give the 40 kb fragment deletion vector pJTU3853, which was then introduced into *S. flavogriseus* by conjugation to generate mutant ZWK1 by double crossover (Figure S1A). PCR using primers ZWK1-DF/DR confirmed the deletion (not shown). The determination of the boundaries of the *xan* cluster and construction of individual gene inactivation mutants is described in detail in Figure S1 and the Supplemental Experimental Procedures.

Analysis of Xantholipin Metabolites

S. flavogriseus was grown for six days in fermentation medium at 30°C, collected by centrifugation and extracted overnight with *n*-butanol. The supernatant was collected by centrifugation, dried under reduced pressure and dissolved in methanol. The methanol extract was analyzed by LC-MS (Agilent 1100 series LC/MSD Trap system, Agilent, Santa Clara, CA, USA). The HPLC was operated at a flow rate of 0.5 ml/min on a Waters SB-C18 (3.9 \times 150 mm, 2 μ m) column. Solvent A was 0.2% formic acid and 0.1% TFA, solvent B was acetonitrile. The program used a gradient from 20 to 55% solvent B over 50 min. The elution was monitored at 274 nm and analyzed using the Waters Millennium Chromatography Manager (Agilent). The ion trap mass spectrometer was operated in the positive ion mode. Drying gas flow was 8 l/min, and nebulizer pressure was 206.8 kPa. Drying gas temperature was 325°C, and the fragmentation was between 1.0 V to 1.8 V. An Agilent 6530 Accurate-Mass Q-TOF LC/MS was used for high-resolution mass measurements.

Production, Isolation, and Characterization of Compounds **6** and **8a/b**

Compounds **6** and **8a** were produced, isolated, and analyzed using method similar to each other. First, the two mutant strains *S. flavogriseus* ZWK8 and ZWK2 were grown at 30°C in 7 l and 14 l of fermentation medium, respectively; each 10 l fermenter contained 7 l culture. After six days, the mycelia were harvested by centrifugation and the compounds were extracted from the mycelium by 7 l and 10 l of methanol, respectively. The extract was then evaporated to 2 l, and the remaining liquid was re-extracted by 2 l of ethyl ethanoate. Then the organic phase was concentrated and fractionated by silica gel column chromatography (chloroform/methanol, gradient elution). Fractions containing the target compound were combined, dried, and further purified by preparative LC (C18 column, 40%–60% methanol in H₂O over 10 min for **6**, 50%–80% methanol in H₂O over 10 min for **8a**). Final purification was accomplished by preparative HPLC (C18 column, gradient for **6**; 30 to

60% acetonitrile, [pH 3.5] H₂O with acetic acid over 10 min; gradient for **8a**, 30% to 80% acetonitrile, [pH 3.5] H₂O with acetic acid over 15 min). We obtained 48.5 mg of **6** and 84.5 mg of **8a**.

The structures of **6** and **8a** were characterized using ¹H, ¹³C NMR, DEPT, HSQC, HMBC (Table S1). Single crystals of **8b** were prepared by recrystallization in methanol, and the X-ray analysis was accomplished at the Shanghai Institute of Organic Chemistry.

Expression, Purification, and Analysis of the BVMO XanO4

For the heterologous expression of XanO4 in *E. coli* BL21(DE3)/pLysE/pJTU3889 (pRSET B containing *xanO4*), a BamHI-EcoRI fragment was amplified from genomic DNA using primers XanO4-F/R (Tables S2 and S3). The transformants were grown in 1 l of LBBS (per liter: 10 g trptone, 5 g yeast extract, 10 g sodium chloride, 185.9 g D-sorbitol, and 0.31 g betaine) medium containing ampicillin and chloramphenicol at 37°C to A₆₀₀ 0.6 before addition of isopropyl-β-D-thiogalactoside (IPTG) to 0.2 mM, followed by incubation at 18°C for 24 hr. His₆-XanO4 was adsorbed to a 1 ml HisTrap HP column (GE Healthcare) and eluted at 1 ml/min⁻¹ by a gradient of imidazole from 5 mM to 500 mM over 30 min in 150 mM NaCl and 20 mM Tris-HCl (pH 7.9) using the Amersham Biosciences ÄKTA FPLC.

XanO4-catalyzed oxidation of NADPH was monitored at 30°C at 340 nm, in 50 mM PBS (pH 7.5), containing 0.5 mM NADPH, 0.2 mM FAD, 40 μM XanO4, and 80 μM **6**.

Bioactivity Assays

Cell proliferation was determined using the methyl thiazolyl tetrazolium (MTT) assay for A-594, HCT-116, and P388 cells. Two thousand cells in 100 μl media per well were placed in a 96-well plate and allowed to incubate overnight at 37°C. Then, six different concentrations of xantholipin, **6**, or **8a** in DMSO were added to the cells in triplicate. After 72 hr, 20 μl 5 mg ml⁻¹ MTT was added. After 3 hr, the supernatant was removed and 100 μl DMSO was added to each well. Cell proliferation was assayed colorimetrically using a SpectraMAX340 comparing 550 nm and 690 nm absorptions. All cells were cultured according to ATCC or DSMZ guidelines. Prism (GraphPad) was used for analysis.

Phylogenetic Analysis

Multiple sequences were aligned using ClustalW (Thompson et al., 1994), and the phylogenetic trees were generated by MEGA (version 3.1; Kumar et al., 2004) using neighbor-joining with Poisson correction and 1,000 replicate bootstrap analysis. The sequence of *E. coli* FabG was used as an outgroup to root the tree in Figure 5. To validate the tree topology, minimum evolution was also used as a distance method. Both methods gave similar tree topologies.

ACCESSION NUMBERS

The complete DNA and deduced protein sequences of the *xan* gene cluster reported in this paper have been deposited in GenBank under the accession number GQ421798.

SUPPLEMENTAL INFORMATION

Supplemental Information includes six figures, three tables, and Supplemental Experimental Procedures and can be found with this article online at doi:10.1016/j.chembiol.2012.01.016.

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