RESEARCH ARTICLE

Functional Characterization of a Ketoreductase-Encoding Gene med-ORF12 Involved in the Formation of a Stereospecific Pyran Ring during the Biosynthesis of an Antitumor Antibiotic Medermycin

Qiang He1,2, Le Li2, Tingting Yang1,2, Ruijuan Li1, Aiying Li1,2*

1 State Key Laboratory of Microbial Technology, School of Life Sciences, Shandong University-Helmholtz Institute of Biotechnology, Shandong University, Jinan, China, 2 School of Life Sciences, Central China Normal University, Wuhan, China

* aiyl@126.com

Abstract

Medermycin, a polyketide antibiotic, possesses strong bioactivity against a variety of tumors through a novel mechanism and is structurally featured with a pyran ring containing two chiral centers (3S and 15R). By far the biosynthetic origin of such enantiomerical conformations still remains obscure. In the present study, we reported the functional characterization of a proposed ketoreductase Med-ORF12 encoded by medermycin biosynthetic cluster and revealed its involvement in the stereochemical control at C3 center of medermycin. Firstly, bioinformatics analysis of Med-ORF12 suggested that it belongs to a group of stereospecific ketoreductases. Next, a Med-ORF12-deficient mutant was obtained and LC/MS measurements demonstrated that medermycin production was completely abolished in this mutant. Meanwhile, it was found that two shunt products were accumulated at the absence of Med-ORF12. Finally, the reintroduction of Med-ORF12 into this mutant could restore the production of medermycin. In conclusion, these data supported that Med-ORF12 is essential for the biosynthesis of medermycin and performs its role as a stereospecific ketoreductase in the tailoring steps of medermycin biosynthetic pathway.

Introduction

Streptomyces are great sources of abundant natural products with potential pharmaceutical and agricultural applications, including antibiotics, immune-modulators, antitumor agents and so on [1–2]. Among these natural products, aromatic polyketides (PKs) are valued impressively in human medicine and agriculture, due to their highly structural and bioactive diversity.

Over thirty years, taking actinorhodin (ACT, 1) as a model in the family of benzoisochromanquinones (BIQs, also referenced as pyranonaphthoquinones) [3–5] (Fig 1A), the
Fig 1. Examples of polyketide antibiotics with stereospecific centers and proposed biosynthetic pathways of medermycin and actinorhodin. A: Three members of BIQ family share opposite configuration at C3 and C15 (either 3S15R or 3R15S). Two well-known antibiotics, daunorubicin and avermectin, have more than one stereospecific center and genes for stereochemical control at C13 and C5 for these two antibiotics respectively will be discussed in this study. B: Biosynthetic pathways of MED 2 and ACT 1 were proposed to share common earlier stages to produce a bicyclic intermediate 4. The formation of this intermediate was catalyzed by polyketide synthases (minimal PKS: KS, CLF and ACP) and related enzymes (KR, ARO and CYC). It could be converted into two shunt products (DMAC 5 and aloeasaponarin II 6) in the actVI-ORF1-deficient mutant strain. The keto-reduction at C-3 of this bicyclic intermediate 4 in ACT 1 pathway was performed by ActVI-ORF1, probably by Med-ORF12 in MED 2 pathway, followed by spontaneous cyclization and dehydration, leading to the production of DNPA 7 with 3S configuration.

doi:10.1371/journal.pone.0132431.g001
biosynthesis of aromatic polyketide antibiotics has been extensively studied, especially at the earlier stages: A multifunctional enzyme complex (namely minimal polyketide synthases (minimal PKS, composed of ketosynthase (KS), chain length factor (CLF) and acyl carrier protein (ACP)) is responsible for the formation of the polyketide chain with a certain length, followed by ketoreduction, aromatization and cyclization catalyzed by ketoreductase (KR), aromatase (ARA) and cyclase (CYC) in a serial of reactions, leading to the formation of aromatic polyketide skeletons [6]. Subsequent tailoring modification steps, including methylation, dimerization, oxidation, glycosylation and so on, result in the accumulation of final products with a high diversity in their structures and bioactivity [7]. Compared to the earlier stages, most of these tailoring modification steps still have yet to be investigated, even for the biosynthetic pathway of well-studied ACT [4].

Medermycin (MED, 2), another member in the BIQ family [3], possesses strong bioactivity against many types of tumor cells through a novel mechanism, besides its inhibitory activity against Gram positive bacteria [8–10]. It is featured with a fused three-ring structure composed of a benzene ring, a quinone and a stereospecific pyran ring.

Stereochemical control is a distinguished characteristic in BIQ family, such as 3\textsuperscript{S}15\textsuperscript{R}-conFig uration for ACT 1 and MED 2 and 3\textsuperscript{R}15\textsuperscript{S}-conFig uration for granaticin (GRA 3) (Fig 1A) [3–5]. Stereochemical control is also a common and important tailoring modification during the biosynthetic pathways of many other polyketide antibiotics, including well-known antitumor agent daunorubicin (Fig 1A) [11]. Generally, the stereochemistry of a functional group influences remarkably on the bioactivity of natural products [12–13], hence the biosynthetic problem concerning the formation of this pyran ring in MED 2 has been an important issue [12].

The 29-gene-containing cluster for the biosynthesis of MED 2 was cloned and sequenced in 2003 [14]. Comparative studies between med and act gene clusters proposed that the biosynthetic pathways of ACT 1 and MED 2 share similar earlier stages to produce a bicyclic intermediate 4 (Figs 1B and 2A). Many homologous genes are present on both of these two gene clusters, some of which are proposed for post-PKS modification of ACT 1 [14]. For example, actVI-ORF1 and med-ORF12 share 65/57% of similarity/identity (Fig 2A) [14], implying they share a common role in the pathways of these two compounds.

Evidences in vitro and in vivo proved that actVI-ORF1 encodes a stereospecific ketoreductase responsible for the chirality of C3 in ACT 1 [15–16]. Since two chiral centers (C3\textsuperscript{S} and C15\textsuperscript{R}) in the pyran ring of MED 2 are identical to those in ACT 1 (Fig 1A), med-ORF12 in the med cluster was proposed to determine the chirality of C3 in MED 2 [12, 14].

In the earlier stages of ACT 1 pathway, the bicyclic intermediate 4 could be converted into two shunt products, DMAC (3,8-dihydroxy-1-methylanthraquinone-2-carboxylic acid, 5) and aloeasaponarin II (3,8-dihydroxy-1-methyl-9,10-anthaquinone, 6) at the absence of ActVI-ORF1 (Fig 1B), while ActVI-ORF1 was convincingly verified to convert the bicyclic intermediate 4 into (S)-DNPA (4-dihydro-9-hydroxy-1-methyl-10-oxo-3\textsuperscript{H}-naphtho-[2,3-c]-pyran-3-(S)-acetic acid, 7), which was the first chiral intermediate in ACT 1 pathway (Fig 1B) [4, 15, 17–18].

Our previous studies revealed that med-ORF12 could restore ACT 1 production in the actVI-ORF1-deficient mutant strain [12]. In addition to that, a co-expression system composed of 6 act earlier genes (KS-, CLP-, ACP-, KR-, ARO- and CYC-encoding genes) together with med-ORF12 could result in the production of (S)-DNPA 7, proposing that Med-ORF12 could catalyze a stereospecific ketoreduction at C3 of the bicyclic intermediate 4, formed by 6 act earlier genes [12, 19]. However, there have been no direct evidences to reveal the role of med-ORF12 in the biosynthetic pathway of MED 2.
Hence, in the present study, we performed gene knock out and complementation experiments besides bioinformatics analysis, and proved definitely its involvement in the stereochemical control in MED pathway.

**Materials and Methods**

**Strains, plasmids, media and reagents**

*Streptomyces coelicolor* CH999 (CH999) is an actinorhodin-cluster-deficient strain, and used as host for heterologous expression of the *med* cluster here [20]. The strain CH999/pIK340 was a
medermycin-producing recombinant strain [14], here acting as wild type strain (WT). *Streptomyces* sp. AM-7161 was a medermycin-producing wild type strain [14]. *Escherichia coli* ET12567/pUZ8002 was used for intergeneric conjugation [20]. pIK340 is a plasmid carrying an entire medermycin biosynthetic gene cluster [14]. pRM5 is an expression plasmid containing of 6 act earlier genes (KS-, CLP-, ACP-, KR-, ARO- and CYC-encoding genes) [19]. pT7Blue (Novagen) is used as cloning vector, while pH7, a conditional suicide vector derived from pHZ1358, is used here for knock-out of med-ORF12 [21]. pIJ8600 is an inducible vector used for the expression of streptomyces genes [20].

Media for streptomycete cultivation include YEME, GYM, 2×YT, R4, R5, R5MS (R5 without sugar) and MS media [20, 22]. *E. coli* cells were cultivated on LA agar or liquid medium [23]. Recombinant strains derived from *Streptomyces* or *E. coli* were selected using ampicillin (100 μg/ml as working concentration), apramycin (50 μg/ml) or thiostrepton (25 μg/ml).

**DNA manipulation and general genetic manipulations**
Isolation, restriction enzyme digestion and transformation of DNAs and plasmids were performed according to standard protocols [20, 23]. Plasmids were introduced into the *Streptomyces* strains by intergeneric conjugation between *E. coli* and *Streptomyces* hosts [20].

**Bioinformatics analysis**
Sequence alignment between Med-ORF12 and its homologies was conducted using the Clustal X program version 2 [24–25]. The phylogenetic tree was established based on sequence alignment using distance/neighbor joining programs with the Poisson correction distance model in MEGA software package version 4.0 [26]. The interior branch length supports were from 1000 replicates. The role of genes or gene cluster was predicated using online programs of CDD [27]. The structural modeling of Med-ORF12 was performed using PyMOL and modeled with NADPH using Autodock.

**Construction of the suicide plasmid for knock-out of med-ORF12**
For in-frame deletion of med-ORF12, firstly, a 1.5 kb fragment (namely “left arm”) containing the upstream region of med-ORF12 was amplified using pIK340 as template and a pair of primers (med12-L1/med12-L2, listed in Table 1). The PCR reaction was performed using KOD-Plus polymerase from TOYOBO and under the conditions composed of 25 recycling reactions as: 1 min at 55°C for denaturation, 45 sec at 65°C for annealing and 1 min at 68°C for extension. Similarly, using another primer pair (med12-R1/med12-R2, in Table 1), a 1.3 kb downstream fragment (right arm) was amplified (Fig 3A).

After double digestion using *HindIII* and *XbaI* (for left arm) and *XbaI* and *EcoRI* (for right arm) respectively, these two PCR products were cloned onto pT7Blue generating two plasmids designated as pHSL121 and pHSL122, respectively. After being sequenced, two arms were linked together on pT7Blue via restriction sites (*HindIII*, *XbaI* and *EcoRI* sites, specifically-designed in the primers in Table 1) to give a plasmid as pHSL123, followed by ligating these two linked regions with the suicide vector pH7. The final plasmid was designated as pHSL124. It was introduced into *E. coli* host ET12567/pUZ8002 for subsequent intergeneric conjugation.
In-frame knock-out of med-ORF12

The suicide plasmid pHSL124 was transferred from *E. coli* ET12567/(pUZ8002+pHSL124) into the medermycin-producing wild type strain WT via intergeneric conjugation [20]. After selection for apramycin- and thiostrepton- resistance, six conjugants were picked out randomly and verified by PCR with a primer set as med12-qc1/med12-qc2 (Table 1). Next, two rounds of cultivation of conjugant spores on GYM medium without supplementation of either thiostrepton or apramycin were performed for double cross-over between two arms and their homologous regions on the genome of the wild type strain WT. Several proposed med-ORF12 deficient mutants (designated as MS) were proved by antibiotic-resistance selection and PCR amplification using two primers (med12-qc1/med12-qc2).

Construction of the expression plasmid for complementary analysis of med-ORF12

The 975 bp med-ORF12 gene in a full-length was amplified using a primer set (med12-A/med12-B, Table 1) and plasmid pIK340 as a template for PCR. The resultant product was cloned onto the *Hind*III-*BamHI* sites of pT7Blue to give a plasmid pHSAY18. After being sequenced, the insert was cleaved out from pT7Blue and ligated with the integrating vector pIJ8600, generating a recombinant plasmid pHSAY19, which was transferred into the med-ORF12-deficient mutant strain MS for complement analysis. The obtained transformant strains were proved by PCR using two primers (med12-A/med12-B, Table 1)

Metabolite analysis of streptomycetes strains by LC/MS

To prepare crude extracts from streptomycete cultures, 100 μl of fresh *Streptomyces* spores (1.0x10^6 spores/μl) were inoculated into 10 ml of TSB liquid medium and grown at 30°C for 48 h. Then, 2 ml of the resultant cultures were added to 100 ml of R5MS medium [22] in a 500-ml Erlenmeyer flask. After 5 d incubation on a shaker (200 rpm) at 30°C, supernatants were collected by centrifugation at 6000 rpm for 10 min, followed by extraction with EtoAc [15].

Subsequently, the crude extracts were dissolved in 1 ml of EtoAc and subjected to LC/MS analysis on the system of Agilent 1100 HPLC/Brucker Esquire HCT under the following conditions: column: TSK gel ODS-100S (5 μm, 15.0 cm×4.6 mm, TOSOH); column temperature:
Fig 3. Structure-based sequence alignment between Med-ORF12 and its homologies and the modeled structure of Med-ORF12. A: The primary sequence of Med-ORF12 is aligned with the ActVI-ORF1, JHAD, Sa10 and WP_019763579. The sequences are annotated with corresponding secondary structures in JHAD. Arrows represent β-sheets and helices indicate α-helices. The similar residues are shaded and identical residues are highlighted in black. The key catalytic residues are marked by ☆. The numbers indicating amino acid positions of each entry refer to the nucleotide sequence deposited in the databases. Sequences retrieved from GenBank were aligned using Clustal X. B: Homology model of Med-ORF12 using one chain of JHAD from human heart as template. Left: The homology modeled structure of Med-ORF12 obtained using PyMOL. β-strands and α-helices are shown in yellow and red, respectively. Right: The NADH molecule is modeled in active sites of Med-ORF12 by the docking program Autodock. The NADPH is shown in cyans.
40°C; eluent A: H₂O + 0.1% HCOOH; eluent B: CH₃CN + 0.1% HCOOH; gradient elution: 0–35 min (10–95% B), 35–40 min (95% B), 40–45 min (95–10% B); flow rate: 1.0 ml/min; detection wavelength: 254 nm and 434 nm; ionization mode: APCI(-) [14].

LC/HRMS (High Resolution Mass Spectrum) analysis of two shunt products

Crude extracts from the liquid culture of the med-ORF12-deficient strain were subjected to LC/HRMS measurement on the Bruker Compact (Qq-TOF LC-MSMS mass spectrometer system) in a full scan in a negative ionization mode (APCI-). During HPLC analysis, two shunt products were detected at 254 nm and eluted with a gradient as described as above (section: Metabolite analysis of streptomyces cultures by LC/MS). During subsequent HRMS analysis, the parent ions with m/z 297.0405 for compound DMAC 5 (C₁₆H₉O₆⁻ ([M-H]⁻)) and with m/z 253.0506 for compound aloesaponarin II 6 (C₁₅H₉O₄⁻ ([M-H]⁻)) were monitored and then fragmented.

Results

Bioinformatics analysis of Med-ORF12

Conserved domain search suggested that Med-ORF12 is a member of 3-hydroxyacyl-CoA dehydrogenase protein family (3HAD). 3HADs (EC 1.1.1.35) are related with the fatty acid metabolism and reversibly catalyze the β-oxidation of the hydroxyl group of L-3-hydroxyacyl-CoA into a keto group [28](Fig 2C).

Blast search showed that many members in 3HAD family possess a quite high identities to Med-ORF12, and are widely distributed in many bacteria, most of which are Actinobacteria, such as Streptomyces, Frankia, Nocardiopsis, Kibdelosporangium and Amycolatopsis, while Streptomyces make up a largest portion among these bacteria.

Many homologues in the streptomyces species are involved in the secondary metabolism in Streptomyces: for example, ActVI-ORF1 from S. coelicolor [16] was proved functionally to be an stereospecific ketoreductase involving in actinorhodin biosynthesis [15–16]. Sa10 annotated as a putative 3-hydroxyacyl-CoA dehydrogenase gene (having an identity of 74% to Med-ORF12) was encoded by the indigoidine/auricin hybrid cluster from Streptomyces aureofaciens [29–30]. It is noteworthy that a hypothetical protein (WP_019763579) from Streptomyces sp. Wigar10 has a highest identity (91%) with Med-ORF12. Using antiSMASH program online, we found that it was located within a secondary metabolite gene cluster, which shows extremely high resemblance to the gene cluster for medermycin biosynthesis. So, we deduced WP_019763579 to be involved in the biosynthesis of an unknown natural product [31].

It is well known that ketoreductases (KR) are widely distributed in the pathways of aromatic polyketide natural products. For example, the keto-reduction by KR in the nascent polyketide chains is a common and critical step for the formation of aromatic polyketide skeletal core in the earlier stages [6, 19]. The stereochemistry at C3 of MED 2 via a stereospecific keto-reduction might be controlled by also a ketoreductase, either Med-ORF6 or Med-ORF12 (only two possible candidates encoded by med cluster). So, we established a phylogenetic tree using proposed ketoreductases in BİQ pathways and several closer homologies using MEGA 4.09. Fig 2B showed that these homologies involving microbial secondary metabolism could be divided into three groups, though PSI-Blast revealed that they were divided into two families: Members in
Group I and II belong to the family of short-chain alcohol dehydrogenases (SDR-family) [32] while members in Group III belong to that of L-3-hydroxyacyl-CoA dehydrogenase (3HAD-family) (Fig 2B).

Even we used more homologues from *streptomyces* species and a non-streptomyces species for phylogenetic analysis and found they could be categorized into group III (Figure A in S1 File). Blast search for the downstream and upstream genes of these homologues in their genomes supported that most of them might be related to the biosynthesis of secondary metabolites, though which structures remain unknown [33–36].

Several closer homologies in Group III and Med-ORF12 were multiply aligned with Clustal X, showing that some conserved sites commonly exist in these enzymes, such as His141/Glu153 for Med-ORF12 and His129/Glu141 for ActVI-ORF1. Secondary structures analysis revealed that eight β-sheets and eleven helixes are aligned in a similar mode in all examples (Fig 3A).

Subsequently, we performed structural modeling of Med-ORF12 using 3HAD from human heart as reference protein template (Med-ORF12 has 27% identity to 3HAD) [28], revealing that Med-ORF12 shares a similar structure to 3HAD (Fig 3B). Similar to the subunit of 3HAD, the first domain of Med-ORF12 is composed of alpha/beta dinucleotide folds and an unusual helix-turn-helix motif extending from the central beta-sheet, forming a Rossman fold, which should be for NADPH-binding. The second domain at C-terminal is composed of primarily alpha-helical, and might mediate subunit dimerization. Fig 3B also showed that these conserved sites are located in the proposed catalytic center of Med-ORF12.

**Inactivation of Med-ORF12**

To verify the role of Med-ORF12 in the biosynthetic pathway of MED2, we conducted knock-out experiments: a conditional suicide plasmid pHSL124 was constructed using pYH7 as vector (Fig 4A). pYH7 is an automatically-replicating vector under the selection of thiostrepton, but acts as a suicide plasmid without thiostrepton selection [21]. An artificially-designed sequence (ATG TCTAGA TGA) was designed to replace in-frame the full gene of *med*-ORF12. Subsequently, pHSL124 was introduced into the wild type strain WT.

We picked out several apramycin- or thiostrepton-sensitive colonies and confirmed them to be *med*-ORF12-deficient strains (designated as MS) (Fig 4B) by PCR amplification.

The supernatants of mutant strains cultivating on R5MS liquid medium were analyzed by HPLC for MED2 production, using the wild type strain as positive control. Red-brownish pigmentation in the MED-producing strain acted as indicator for MED2 production [14]. Fig 5A showed *med*-ORF12 deficiency caused pigmentation disappearance in the supernatants of the mutant strain grown on liquid medium, in a sharp contrast to the wild type strain, implying the disappearance of MED2 production. HPLC measurements (Fig 5B and Figure B in S1 File) demonstrated clearly that MED2 production was abolished completely in *med*-ORF12-deficient mutant strain.

**Complementary analysis of Med-ORF12**

In order to characterize further the role of *med*-ORF12, we performed complementary experiments to investigate if the expression of *med*-ORF12 could restore *med*-ORF12-deficiency. An expression plasmid pHSAY19 was constructed derived from an integrative vector pIJ8600, which was designed for expression of *med*-ORF12 under the control of the inducible thiostrepton promoter, tipAp [20]. After introducing it into the mutant strain MS, we verified the integration of *med*-ORF12 in the chromosome of the host MS by PCR amplification (Fig 4B).

After 5 d expression induced by thiostrepton supplemented into the liquid culture of the complementary strain MS/pHSAY19, the supernatant showed obvious restoration in
pigmentation, on the contrary to the mutant strain MS/pIJ8600 (Fig 5A). HPLC measurement suggested that complementary strain could produce MED with a comparable yield as in the wild type strain (Fig 5B).

**Accumulation of two shunt products at the absence of med-ORF12**

In a sharp contrast to the wild type strain, LC/MS measurement (Fig 6A) showed that two compounds indicated as two peaks at 16.7 and 22.6 min were accumulated significantly in the
med-ORF12 deficient mutant strain (Figure B in S1 File) also showed several other new peaks in a lower amount between 15–25 min besides these two major ones, in a contrast to the wild type strain).

CH999/pRM5 was a producer of DMAC\(^5\) and aloesaponarin II\(^6\), two shunt products in ACT\(^1\) pathway (Fig 1B). Using CH999/pRM5 as a positive control, it was observed that two compounds by the med-ORF12-deficient mutant strain showed identical retention time and mass spectra to these shunt products by CH999/pRM5 (Fig 6). HRMS analysis of these two shunt products further suggested this mutant strain could accumulate DMAC\(^5\) and aloesaponarin II\(^6\) (Figure C and Figure D in S1 File).

**Discussion**

**Essential role of Med-ORF12 in the biosynthesis of medermycin**

Stereochemical control is a common and important tailoring modification during the biosynthetic pathways of many polyketide antibiotics, including aromatic polyketide antibiotics daunorubicin and ACT\(^1\) (Fig 1A), and well-known macrolide antibiotic avermectin (Fig 1) [37]. Understanding the stereochemistry in these compounds is of highly chemical, biological and pharmacological importance for generating hybrid natural products with improved pharmacological profiles via metabolic engineering in a combinatorial fashion.

MED\(^2\) possesses 7 stereospecific centers (Fig 1). In the present study, our investigation focused on the conFig uration at C3 center of its pyran ring. We characterized functionally a ketoreductase-encoding gene med-ORF12 located in med cluster by knock-out and complementary experiments, and found that its deficiency caused disappearance of MED\(^2\).
Meanwhile, introduction of \textit{med}-ORF12 into this mutant strain could restore the production of MED 2. Our data here offered the direct evidences to reveal its critical role in the biosynthesis of MED 2 for the first time.

**Involvement of \textit{med}-ORF12 in the stereochemical control**

In the present study, we also observed two shunt products were accumulated at the absence of \textit{med}-ORF12 (Fig 6). As shown in Fig 1B, at the presence of 6 \textit{act} genes for earlier stages of ACT pathway.
The bicyclic intermediate 4 was produced, but further entered into a shunt pathway if actVI-ORF1 is absent [15]. So, actVI-ORF1-deficient strain could accumulate two shunt products DMAC 5 and aloesaponarin II 6. The recombinant strain S. coelicolor CH999/pRM5 was also able to produce these two shunt products because pRM5 carried 6 genes encoding KS, ACP, CYC, KR, ARO and CYC from ACT pathway (Fig 1B), though CH999 is an entire-ACT-cluster-deficient mutant from S. coelicolor [15,19].

Our previous studies proved that Med-ORF12 could convert the bicyclic intermediate 4 into (S)-DNPA 7 with a configuration (3S) at C3, when working together with these 6 act early biosynthetic genes encoding KS, ACP, CYC, KR, ARO and CYC [12]. This suggested that Med-ORF12 could recognize the bicyclic intermediate 4 as substrate and play a similar role to ActVI-ORF1. Hence, it is predicted that two shunt products DMAC 5 and aloesaponarin II 6 could be accumulated in the med-ORF12-deficient strain. LCMS and HRMS measurements in the present study proved such a speculation, using CH999/pRM5 as a control (Fig 6 and Figure C and Figure D in S1 File).

All these evidences supported that Med-ORF12 could play an essential role in the MED 2 pathway, as a ketoreductase stereospecific for the formation of the chiral center at C3 of MED 2.

Three groups of ketoreductases in the biosynthetic pathways of polyketides

Ketoreductases belong to the family of oxidoreductases and widely distributed in the pathways of aromatic polyketide natural products. Here, our phylogenetic analysis revealed that these ketoreductases from Streptomyces strains and even from non-Streptomyces strains could be divided into three groups and PSI-Blast search further demonstrated that group I and II belong to SDR-family and group III belongs to 3HAD-family (Fig 3B and Figure A in S1 File).

Therefore, the closer evolutionary distance between Group I and II allowed us to deduce that both of them might originate from a common ancestor, though they undergo different evolutionary routes to acquire different function: Group I represents region-specific ketoreductases (KR), exemplified by ActIII for keto-reduction at C-9 of ACT 1, determining the precise position for the folding of a 16-membered polyketide chain before aromazation and cyclization [3–5]. All the conserved and characteristic core motifs and amino-acid residues of the domains are present in all the members in group I.

Among the members in Group III and II, only ActVI-ORF1, Gra-ORF6 and Med-ORF12 have been characterized functionally.

Though the members in Group II (SDR family) have quite low identities to those in Group III (3HAD family), previous evidences suggested that both of them might be involved in the enantiomeric control in the pathways of polyketides, as stereo-specific ketoreductases:

1. Gra-ORF6 in group II and ActVI-ORF1 in Group III were revealed to function as stereo-specific ketoreductases in the pathways of GRA 3 (for 3R conFig uration) and ACT 1 (for 3S conFig uration) respectively (Fig 1A) [15–16]. Consistently, 3HAD catalyzes the fatty acid metabolism as stereo-specific ketoreductase (Fig 2C).

2. AveF and DnrU in group II were proposed to be ketoreductases for stereochemical control at C-5 in avemectin and at C13 in daunorubicin respectively [13–14], though no direct experimental evidences support the functions of them.

We also found the highest homologue WP_019763579 (identity: 91%) might be involved in the biosynthesis of a mediromycin-like compound (though it was annotated as a hypothetical protein in GenBank), because antiSMASH analysis supported that WP_019763579 is located within a med-like gene cluster in the genome of Streptomyces sp. Wigar10 [31]. Additionally,
based on the phylogenetic tree in Figure A in S1 File, we proposed more proteins classified into group III might have a similar role to Med-ORF12 and ActVI-ORF1.

Examples in these two groups might perform different mechanisms: previous biochemical analysis *in vitro* and *in vivo* suggested Gra-ORF6 and ActVI-ORF1 control completely opposite chirality in GRA 3 and ACT 1 pathways respectively, though they could recognize an identical substrate (the common bicyclic intermediate 4) [15–16]. It is noteworthy that AveF and DnrU also control different opposite chirality in avemectin (C5R) and daunorubicin (C13S) pathways, but both of them are present in a same group. Hence, the mechanisms of members in Group II and III need to be further clarified.

**Sequence comparison between 3HAD and members in group III**

Med-ORF12 belongs to the family of 3HAD. 3HAD could catalyze a reversible reaction between L-3-hydroxyacyl-CoA and 3-oxoacyl-CoA and mainly exist in mammals (Fig 2C). Some members of 3HAD family are also found in many microorganisms, including the examples in Group III. It is noteworthy that 3-oxoacyl-CoA, the substrate for 3HAD in mammals, showed a partially similar structure to the reactive portion of the bicyclic intermediate 4, the common substrate for ActVI-ORF1 and Med-ORF12 (Figs 1B and 2).

Molecular modeling studies and biochemical observations of 3HAD from human heart elucidated that two sites (His158 and Glu170) constituted a catalytic dyad located in the active site region [28]. As the first example characterized to be involved in microbial secondary metabolism in Group III, ActVI-ORF1 also contains two conserved residues (His129 and Glu141), which were proved to be critical for the reduction of the C-3 keto group of the bicyclic intermediate 4 [16]. In the present study, we found these two sites are highly conserved in all the members of Group III, further indicating their importance for the reducing activity of these enzymes.

Further, similar to ActVI-ORF1 and 3HAD, we found these two residues are located in the active site region in the modeled structure of Med-ORF12 (Fig 3B), as the second example characterized to be involved in microbial secondary metabolism in Group III. This implied that Med-ORF12 controls the chiral conformation at C3 of MED 2 via a similar mechanism to ActVI-ORF1 and 3HAD: In the catalytic dyad composed of Glu153 and His141, the carboxyl group of Glu153 is responsible for the activation of the proton in His141 side chain, probably via nucleophilic attack. Then the activated proton of the imidazolyl ring of His141 mediated the electron transfer from C-3 keto-group of the substrate. Accordingly, the pro-S hydride from NAD(P)H was transferred to the C-3 of the bicyclic intermediate 4, leading to the formation of S conFig uration.

We have established prokaryotic expression and purification system of Med-ORF12 [38] and hope to perform enzymatic assay *in vitro* in next steps to investigate its mechanism.

In a conclusion, stereochemical control in BIQ pathways is one of most important tailoring modification. In the present study, functional characterization by gene knockout, gene complementary, LC/MS and HRMS analysis elucidated convincingly that Med-ORF12 is essential for the production of MED 2 as a stereospecific ketoreductase and reduces the keto-group at C3 during MED 2 pathway to form a specific conFig uration (3S). These data supported further that MED 2 and ACT 1 share parts of tailoring modification steps besides their earlier stages, and will help us understand better the stereochemistry in BIQ pathways as an important step towards metabolic engineering to generate a novel pharmacophore.

**Supporting Information**

S1 File. Bioinformatics analysis of Med-ORF12 and HRMS spectra of two shunt products. Figure A: Phylogenetic tree established using more homologues of Med-ORF12. These entries
include proteins involved in biosynthesis of secondary metabolites: Med-ORF12 and Med-ORF6 (for MED2 in *Streptomyces sp AM-7161*), ActVI-ORF1 and ActIII (for ACT1 in *S. coelicolor A3*(2)), DauB (for aklaviketone in *Streptomyces sp*), Gra-ORF5 and Gra-ORF6 (for GRA3 in *Streptomyces violaceoruber*), AveF (for avermectin in *Streptomyces avermitilis*), DnrU(for daunorubicin in *Streptomyces peucetius*), Sa10 (for indigoidine/auricin in *Streptomyces aureofaciens*), 3HAD (3HAD_B) from human heart and a hypothetical protein WP_019763579 (encoded by a cluster highly in *Streptomyces sp* Wigar10, homologous to medermycin cluster). Additionally, several highly homologous proteins proposed to encode hydroxylacyl-CoA dehydrogenases were also analyzed: WP_033529872 from *Streptomyces galbus* (identity to Med-ORF12: 80%), WP_030735501 from *Streptomyces sp.* (66%), WP_043385010 from *Streptomyces mutabilis* (66%), WP_030080416 from *Streptomyces peucetius* (65%), WP_042193116 from *Kibdelosporangium sp.* (65%) and WP_030666394 from *Streptomyces cellulosae* (64%). The bar indicated the evolutionary distance. The numbers on branch nodes were percentages of 1000 sets of bootstrap supports. All homologies are divided into two families (3HAD: 3-hydroxyacyl-CoA dehydrogenase protein family; SDR: short-chain alcohol dehydrogenases family) and further into three groups. **Figure B**: Comparison of metabolite production between the wild type strain WT/pIJ8600 (A) and med-ORF12-deficient strain MS/pIJ8600 (B and C). Crude extracts isolated from the wild type and mutant strains respectively were subjected to HPLC analysis, indicated as UV absorption at 434 nm. The Y axes of B and C (both for mutant strain) were adjusted into different scales for better comparison with the wild type strain. In a contrast to the wild type strain (WT/pIJ8600, A), the mutant strain (MS/pIJ8600) could not produce MED2 due to the deficiency of med-ORF12, but could produce many intermediates or shunt products (12–25 min), which were not present in the wild type strain. Among of them, two major components (deduced to be DMAC5 and aloesaponarin II6) were further analyzed in next experiments. **Figure C**: HRMS analysis of aloesaponarin II6 from the control strain CH999/pRM5 (A and C) and med-ORF12-deficient strain MS/pIJ8600 (B and D). The parent ions were detected in A and B (HR-MS, A and B) using atmospheric pressure chemical ionization (APCI-), and then further fragmented (HR-MS2, C and D). **Figure D**: HRMS analysis of DMAC5 from the control strain CH999/pRM5 (A) and med-ORF12-deficient strain MS/pIJ8600 (B). The parent ions were detected in A and B (HR-MS, APCI-).

(DOCX)

**Acknowledgments**

We are grateful to Prof. Koji Ichinose for the generous gift of plasmid pIK340. Thanks for financial supports by the National Natural Science Foundation of China (No. 31170050 and 30770036), the Fundamental Research Funds for the Central University (No. 2014ZD003 and CCNU10A02011), and Xinjiang Production & Construction Corps Key Laboratory of Protection and Utilization of Biological Resources in Tarim Basin (No. BRZD1401).

**Author Contributions**

Conceived and designed the experiments: AL. Performed the experiments: QH LL TY. Analyzed the data: TY RL. Contributed reagents/materials/analysis tools: AL. Wrote the paper: AL. Obtained permission for use of pIK340: KI.

**References**

1. Challis GL. Genome mining for natural product discovery. J Med Chem. 2008; 51: 2618–2628. doi: 10.1021/jm700948z PMID: 18393407
2. Zhang H, Wang H, Wang Y, Cui H, Xie Z, Pu Y, et al. Genomic sequence-based discovery of novel angucyclinone antibiotics from marine Streptomyces sp. W007. FEMS Microbiol. Lett. 2012; 332(2): 105–112. doi: 10.1111/j.1574-6968.2012.02582.x PMID: 22536997

3. Ichinose K, Taguchi T, Ebizuka Y & Hopwood DA. Biosynthetic gene clusters of benzoisochromanequinone antibiotics in Streptomyces spp.: identification of genes involved in post-PKS tailoring steps. Actinomycetologica. 1998; 12: 99–109.

4. Metsä-Ketelä M, Oja T, Taguchi T, Okamoto S & Ichinose K. Biosynthesis of pyranonaphthoquinone polyketides reveals diverse strategies for enzymatic carbon-carbon bond formation. Curr Opin Chem Biol. 2013; 17(4): 562–570. doi: 10.1016/j.cpb.2013.06.032 PMID: 23886982

5. Donner CD. The divergent asymmetric synthesis of kalafungin, 5-epi-frenolicin B and related pyranonaphthoquinone antibiotics. Tetrahedron. 2013; 69 (2013): 377–386.

6. Hopwood DA. Genetic contributions to understanding polyketide synthases. Chem Rev. 1997; 97: 2465–2497. PMID: 11851466

7. Rix U, Fischer C, Remsing LL & Rohr J. Modification of post-PKS tailoring steps through combinatorial biosynthesis. Nat Prod Rep. 2002; 19: 542–580. PMID: 12430723

8. Takano S, Hasuda K, Ito A, Koide Y, Ishii F, Haneda I, et al. A new antibiotic, medermycin. J Antibiot. 1976; 29: 765–768. PMID: 956059

9. Salaski EJ, Krishnamurthy G, Ding WD, Yu K, Insal SS, Eid C, et al. Pyranonaphthoquinone lactones: a new class of AKT selective kinase inhibitors alkylates a regulatory loop cysteine. J Med Chem. 2009; 52(8): 2181–2184. doi: 10.1021/jm900075g PMID: 19309081

10. Toral-Barza L, Zhang WG, Huang X, McDonald LA, Salaski EJ, Barbieri LR, et al. Discovery of lactonomicin and related pyranonaphthoquinones as potent and allosteric inhibitors of AKT/PI3K: mechanistic involvement of AKT catalytic activation loop cysteines. Mol Cancer Ther. 2007; 6(11): 3028–3038. PMID: 17989320

11. Madduri K & Hutchinson CR. Functional characterization and transcriptional analysis of a gene cluster governing early and late steps in daunorubicin biosynthesis in Streptomyces peucetius. J Bacteriol. 1995; 177(17): 3879–3884. PMID: 7601857

12. Li A, Itoh T, Taguchi T, Xiang T, Ebizuka Y & Ichinose K. Functional studies on a ketoreductase gene from Streptomyces sp. AM-7161 to control the stereochimery in medermycin biosynthesis. Bioorg Med Chem. 2005; 13(24): 6856–6863. PMID: 16169737

13. Ichinose K, Bedford DJ, Tomus D, Bechthold A, Bibb MJ, Revill WP, et al. The granaticin biosynthetic gene cluster of Streptomyces violaceoruber Tü22: sequence analysis and expression in a heterologous host. Chem Biol. 1998; 5(11): 647–659. PMID: 9831526

14. Ichinose K, Ozawa M, Ito K, Kunieda K & Ebizuka Y. Cloning, sequencing and heterologous expression of the medermycin biosynthetic gene cluster of Streptomyces sp. AM-7161: towards comparative analysis of the benzoisochromanequinone gene clusters. Microbiology. 2003; 149(7): 1633–1645.

15. Taguchi T, Ito K., Ebizuka Y, Malpartida F, Hopwood DA, Surti CM. Chemical characterisation of disruptants of the Streptomyces coelicolor A3(2) actVI genes involved in actinorhodin biosynthesis. J Antibiot (Tokyo). 2000; 53(2): 144–152.

16. Taguchi T, Kunieda K, Takeda-Shitaka M, Takaya D, Kawano N, Kimberley MR, et al. Remarkably different structures and reaction mechanisms of ketoreductases for the opposite stereochemical control in the biosynthesis of BIS antibiotics. Bioorg Med Chem. 2004; 12(22): 5917–5927. PMID: 15498668

17. Taguchi T, Yabe M, Odaki H, Shinozaki M, Metsä-Ketelä M, Arai T, et al. Biosynthetic conclusions from the functional dissection of oxygenases for biosynthesis of actinorhodin and related streptomyces antibiotics. Chem & Biol. 2013; 20: 510–520.

18. Itoh T, Taguchi T, Kimberley MR, Booker-Milburn KL, Stephenson GR, Ebizuka Y, et al. Actinorhodin biosynthesis: structural requirements for post-PKS tailoring intermediates revealed by functional analysis of ActVI-ORF1 reductase. Biochemistry. 2007; 46(27): 8181–8188. PMID: 17579485

19. McDaniel R, Ebert-Khosla S, Hopwood DA & Khosla C. Engineered biosynthesis of novel polyketides. Science. 1993; 262: 1546–1550. PMID: 8248802

20. Kieser T, Bibb MJ, Buttnrr MJ, Chater KF & Hopwood DA. Practical Streptomyces Genetics. John Innes Foundation, Norwich, UK, 2000

21. Sun Y, He X, Liang J, Zhou X & Deng Z. Analysis of functions in plasmid pHZ1358 influencing its genetic and structural stability in Streptomyces lividans 1326. Appl Microbiol Biotechno. 2009; 82 (2): 303–310.

22. Okamoto S, Taguchi T, Ochi K, Ichinose K. Biosynthesis of actinorhodin and related antibiotics: discovery of alternative routes for quinone formation encoded in the act gene cluster. Chem Biol. 2009; 16(2): 226–236. doi: 10.1016/j.chembiol.2009.01.015 PMID: 19246012
23. Sambrook J & Russell D. Molecular Cloning, A Laboratory Manual. 3rd ed. Cold Spring Harbor Laboratory Press, New York. 2001.

24. Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H, et al. Clustal W and Clustal X version 2.0. Bioinformatics. 2007; 23: 2947–2948. PMID: 17846036

25. Thompson JD, Gibson TJ, Plewniak F, Jeanmougin F & Higgins DG. The ClustalX windows interface: flexible strategies for multiple sequence alignment aided by quality analysis tools. Nucl Acids Res. 1997; 25: 4876–4882. PMID: 9396791

26. Tamura K, Dudley J, Nei M & Kumar S. MEGA4: Molecular evolutionary genetics analysis (MEGA) software version 4.0. Mol Biol Evol. 2007; 24: 1596–1599. PMID: 17488738

27. Blin K, Medema MH, Kazempour D, Fischbach MA, Breitling R, Takano E, et al. antiSMASH 2.0—a versatile platform for genome mining of secondary metabolite producers. Nucleic Acids Res. 2013; 41 (Web Server issue): W204–212. doi:10.1093/nar/gkt449 PMID: 23737449

28. Barycki JJ, O'Brien LK, Bratt JM, Zhang R, Sanishvili R, Straus AW, et al. Biochemical characterization and crystal structure determination of human heart short chain L-3-hydroxyacyl-CoA dehydrogenase provide insights into catalytic mechanism. Biochemistry. 1999; 38: 5786–5798. PMID: 10231530

29. Novakova R, Bistakova J, Homereva D, Rezuchova B & Kormanec J. Cloning and characterization of a polyketide synthase gene cluster involved in biosynthesis of a proposed angucycline-like polyketide auricin in Streptomyces aureofaciens CCM 3239. Gene. 2002; 297(1–2): 197–208. PMID: 12384301

30. Novakova R, Odnogova Z, Kutas P, Feckova L & Kormanec J. Identification and characterization of an indigoidine-like gene for a blue pigment biosynthesis in Streptomyces aureofaciens CCM 3239. Folia Microbiol (Praha). 2010; 55(2): 119–125.

31. Klassen JL, Adams SM, Bramhacharya S, Giles SS, Goodwin LA, Woyke T. et al Draft genome sequence of Streptomyces sp. strain Wigar10, isolated from a surface-sterilized garlic bulb. J Bacteriol. 2011; 193(24): 6999–7000. doi: 10.1128/JB.06257-11 PMID: 22123757

32. Persson B & Kallberg Y. Classification and nomenclature of the superfamily of short-chain dehydrogenases/reductases (SDRs). Chem-Biol Interact. 2013; 202(1–3):111–115. doi: 10.1016/j.cbi.2012.11.009 PMID: 23200746

33. Kim HJ, Karki S, Kwon SY, Park SH, Nahm BH, Kim YK. A single module type I polyketide synthase directs de novo macrolactone biogenesis during galbonolide biosynthesis in Streptomyces galbus. J Biol Chem. 2014; 289(50):34557–34568. doi: 10.1074/jbc.M114.602334 PMID: 25336658

34. Ogasawara Y, Torrez-Martinez N, Aragon AD, Yackley BJ, Weber JA, Sundararajan A. High-quality draft genome sequence of Actinobacterium Kibdelosporangium sp. MJ126-NF4, producer of Type II polyketide azicemicins, using Illumina and PacBio technologies. Genome Announ. 2015. 3(2).

35. Fehr T, Kuhn M, Loosli HR, Ponelle M, Boelsterli JJ & Walkinshaw MD. 2-Epimutalomycin and 28-epi-mutalomycin, two new polyether antibiotics from Streptomyces mutabilis. derivatization of mutalomycin and the structure elucidation of two minor metabolites. J Antibiot (Tokyo). 1989; 42(6):897–890.

36. Li Z, Rawlings BJ, Harrison PH, Vederas JC. Production of new polyene antibiotics by Streptomyces cel-lulosae after addition of ethyl (Z)-16-phenylhexadec-9-enoate. J Antibiot (Tokyo). 1989; 42(4):577–584.

37. Omura S, Ikeda H, Ishikawa J, Hanamoto A, Takahashi C, Shinose M, et al. Genome sequence of an industrial microorganism Streptomyces avermitilis: deducing the ability of producing secondary metabolites. Proc Natl Acad Sci, USA. 2001; 98 (21): 12215–12220. PMID: 11572948

38. Sun R, Liu M, Gong C, Wang W, Zeng A & Li A. Polyclonal antiserum preparation and expression detection of med-ORF12 encoding a stereochemical ketoreductase possibly involved in medermycin biosynthesis. Acta Microbiol Sinica. 2012; 52(1): 60–68.