Microbial Biotechnology

Sulfane sulfur-activated actinorhodin production and sporulation is maintained by a natural gene circuit in *Streptomyces coelicolor*

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Summary

Sulfane sulfur, including polysulfide and persulfide, is a newly identified cellular component present in microorganisms; however, its physiological functions are unclear. *Streptomyces coelicolor* M145 is a model strain of actinomycetes, which produces several polyketides, including actinorhodin. Herein, we found that both exogenously added and endogenously generated sulfane sulfur increased the actinorhodin production and accelerated spore formation of *S. coelicolor* M145. This bacterial species carries a natural gene circuit containing four genes that encode a CsoR-like transcription factor (ScCsoR), persulfide dioxygenase (ScPDO), rhodanese and a sulfite transporter, which were shown to be responsible for sensing and removal of excessive sulfane sulfur. ScCsoR was observed to bind to the promoters of the four genes, thus repressing their transcription. Sulfane sulfur modified Cys37 of ScCsoR, and the modified ScCsoR did not bind to the promoters, thereby activating the transcription of ScPDO. The deletion of ScCsoR decreased cellular sulfane sulfur, while the deletion of ScPDO increased its levels. The increased sulfane sulfur promoted actinorhodin production and sporulation. This study unveiled a natural gene circuit for maintaining sulfane sulfur homeostasis in bacteria. Further, we identified the trigger effect of sulfane sulfur on actinorhodin production, presenting a new approach for activating polyketide gene clusters in actinomycetes.

Introduction

Sulfane sulfur, presenting as persulfide (HSSH and RSSH) and polysulfide (HSSₙH, Sₙ, RSSₙH, RSSₙR, n ≥ 2), is a common component found in mammalian cells, (Fukuto et al., 2018). It plays important physiological roles in cytoprotection, anti-inflammatory activities and angiogenesis (Lau and Pluth, 2019) and is produced by various enzymes. Sulfide: quinone oxidoreductase (SQR) oxidizes H₂S to hydrogen polysulfide (H₃SₙH). Cystathionine beta-lyase (CBS), cystathionine gamma-lyase (CSE), 3-mercaptopyruvate sulfurtransferase (3-MST) and cysteiny-l-tRNA synthetase 2 (CAR2) produce sulfane sulfur from cysteine and its derivatives. Sulfane sulfur can be oxidized by persulfide dioxygenase (PDO; Hildebrandt and Grieshaber, 2008; Liu et al., 2014; Shen et al., 2015) or be reduced to sulfide by cellular thiols, thioredoxin and glutaredoxin (Hou et al., 2019).

Sulfane sulfur and its metabolizing enzymes have been studied in microorganisms as well, demonstrating that it is a key intermediate of H₂S oxidation pathway (Luebke et al., 2014; Li et al., 2017b; Shimizu et al., 2017). Sulfane sulfur also modifies some gene regulators, a process now termed as protein S-sulfhydration or persulfidation, to activate the expression of SQR and/or other H₂S metabolizing enzymes. Four types of such gene regulators (CstR, BigR/SqrR, FisR and OxyR) have been identified (Giedroc, 2017; de Lira et al., 2018; Hou et al., 2019; Shimizu and Masuda, 2020). A recent research indicates that sulfane sulfur is involved in the regulation of pathogenicity in *Staphylococcus aureus* (Peng et al., 2017), suggesting that it has other physiological functions in bacteria.

Actinomycetes are remarkable producers of polyketide drugs, including erythromycin A, rifamycin, chromomycin and tetracyclines (Robertsen and Musiol-Kroll, 2019). Advancements in microbial genomics indicate that earlier discoveries cite only a small fraction of polyketides in actinobacteria, representing approximately 10% of their biosynthetic capacity (Nett et al., 2009; Shen, 2015; Blin...
et al., 2019). Additionally, programmes related to isolation of natural products have significantly decreased since the end of the 21st century (Li and Vederas, 2009; Wright, 2017). One of the reasons responsible for the dearth of research on this subject is the fact that in actinobacteria, many biosynthetic gene clusters are regulated via unknown mechanisms. Their activation is on an ‘as-needed’ basis; however, identifying the need is difficult (Bibb, 2013; Barka et al., 2016).

Streptomyces coelicolor M145 is a model strain for studying actinomycetes. It produces several polyketide compounds (Bentley et al., 2002; Hopwood, 2007), including blue-pigmented actinorhodin (ACT; Bystrykh et al., 1996), red-pigmented undecylprodigiosin and yellow-pigmented coelomycin P2 (Liu et al., 2013; Chen et al., 2016). Among them, ACT has served as an outstanding example for genetic and biochemical investigations of polyketide metabolism, including the seminal work on polyketide biosynthetic gene clusters and the generation of hybrid polyketides (Strohl and Connors, 1992; Khosla et al., 1993). Although transcription regulation of the ACT biosynthetic gene cluster has been intensively studied, the intracellular signal that activates ACT production is still unclear (Liu et al., 2013; Mak and Nodwell, 2017).

In this study, we found that exogenous sulfane sulfur could function as a signal to activate ACT production in S. coelicolor M145. Endogenous sulfane sulfur was detected in this bacterium, and a natural gene circuit controlling its level was identified. Genetically editing the gene circuit changed the intracellular sulfane sulfur levels, ACT production and morphological development of S. coelicolor M145. The gene circuit contains elements that can sense and control intracellular sulfane sulfur level, to maintain it in a homeostatic state like a thermostat. This study unveiled new physiological roles of sulfane sulfur that may provide a new strategy for activating cryptic polyketide gene clusters in actinomycetes.

Results

Treating S. coelicolor with sulfane sulfur promoted its ACT production and spore formation

We used both sulfide (in the form of sodium hydrosulfide [NaHS]) and sulfane sulfur (in the form of S₈) to treat S. coelicolor M145. These chemicals were mixed with the yeast-beef-peptone (YBP) agar and poured on a culture plate. Next, S. coelicolor M145 spores were spread on the surface of the plate, and the plate was incubated at 30°C for 6 days. Addition of sulfide marginally increased ACT production, while addition of S₈ significantly increased the production in the bacterial cells compared with that of untreated control cells (Fig. 1A and B). The highest increase was observed with the addition of 50 μM S₈, about fourfold higher than the control. The addition of S₈ also influenced morphological development of S. coelicolor M145, as evidenced by more spores being produced on the S₈-containing plate (Fig. 1C). These results demonstrated that sulfane sulfur can act as a stimulus to affect ACT production and spore formation of S. coelicolor M145.

Endogenously produced sulfane sulfur also promoted ACT production and spore formation

Persulfide dioxygenase oxidizes glutathione persulfide (GSSH) and can lower cellular sulfane sulfur (Xia et al., 2017). The S. coelicolor M145 genome contains a gene coding for a hypothetical protein, SCO0618, containing an N-terminal PDO-like domain and a C-terminal rhodanese domain (Fig. 2A). This indicates that SCO0618 might be a type III PDO. We compared it with three representative type III PDOS, Zunongwangia profunda SM-A87 (ZpPDOIII, ADF52140.1), Staphylococcus aureus (SaPDOIII, WP_000465474.1) and Bacillus cereus ATCC 10876 (BcPDOIII, EEK49737.1) by using ClustalW (Fig. S1). To test whether this protein catalyses sulfane sulfur oxidation, we cloned its ORF into a pET15b vector and transformed the recombinant plasmid into E. coli BL21(DE3) for expression. The enzyme thus obtained was purified with an N-terminal His-tag using affinity chromatography and was confirmed via SDS-PAGE (Fig. S2). Subsequent enzymatic assays indicated that it oxidized sulfane sulfur. When glutathione persulfide (GSSH) was used as a substrate, its Vₘₐₓ reached 1.19 ± 0.2 μmol O₂ min⁻¹ mg⁻¹ and Kₘ value was 94.25 ± 54.61 μmol (Fig. 2B and C). Hereafter, we refer to this enzyme as ScPDO.

We deleted the Scpdo gene in S. coelicolor M145. S. coelicolor M145 wild type (wt) and the mutant ΔScpdo did not display a significant difference in growth on agar plates. We subsequently detected its intracellular sulfane sulfur levels using a fluorescence-based chemical probe SSP4 using a procedure described by Chen and colleagues (2013). The mutant ΔScpdo accumulated a higher amount of sulfane sulfur, especially at the early exponential phase, than did wt (Fig. 2D). To confirm these results, we used an HPLC-based method to quantify the intracellular sulfane sulfur in both ΔScpdo and wt (Ran et al., 2019). The mutant accumulated 16.06 ± 0.41 μM·OD₄₅₀⁻¹ intracellular sulfane sulfur at 24 h, while wt accumulated 6.41 ± 0.34 μM·OD₄₅₀⁻¹ (Fig. 2E). We also constructed a complemented strain (ΔScpdo:: Scpdo) by introducing an Scpdo-expression plasmid into ΔScpdo, in which the Scpdo expression was controlled by a constitutive promoter, kasOp⁺ (Bai et al., 2015). This strain produced similar amounts of intracellular sulfane sulfur as wt. These results indicated that ScPDO was indeed responsible for sulfane sulfur
Fig. 1. Using NaHS and S₈ to treat *S. coelicolor* M145.
Different concentrations of NaHS or S₈ (20, 50 and 100 µM) were added to agar plates before inoculation of *S. coelicolor* M145 spores. Control was the plate without addition of the chemicals. After inoculation, the plates were incubated at 30 °C for 6 days and then subjected to analysis.
A. Total ACT production of *S. coelicolor* M145. Data were from three repeats.
B. S₈ increased ACT production of *S. coelicolor* M145. Images were captured from the reverse side of the plates.
C. S₈ affected spore formation of *S. coelicolor* M145. Images were captured from the front side of the plates.

Fig. 2. ScPDO had PDO activity, and its deletion increased intracellular sulfane sulfur.
A. ScPDO had two hypothetical domains, PDO and RHOD.
B and C. ScPDO displayed persulfide dioxygenase activity by oxidizing GSSH into sulfite. (B) Persulfide dioxygenase activity was confirmed by detecting O₂ consumption. ScPDO (1 µM) and 0–500 µM GSSH were mixed in 3 ml KPi buffer (100 mM, pH 7.4) at 25 °C, and O₂ consumption was examined using an Orion RDO meter. Black line, reaction mixture with ScPDO; grey line, reaction mixture without ScPDO. (C) Catalysing kinetics of ScPDO was calculated using the Michaelis–Menten equation as reported previously (Liu et al., 2014).
D and E. Detection of intracellular sulfane sulfur of wt, ΔScpdo and ΔScpdo::Scpdo strains with SSP4 and HPLC. High SSP4 fluorescence indicated high sulfane sulfur concentration in the cells. For B–D, data are expressed from three independent repeats and shown as mean ± SD.

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oxidation, preventing the accumulation of sulfane sulfur inside the cells. Notably, in all three strains, sulfane sulfur accumulation showed a similar trend; the level of sulfane sulfur was the highest at the exponential phase, then started to decrease and finally stabilized at the stationary phase (Fig. 2D and E).

We observed that \( \Delta \text{Scpdo} \) produced more ACT than wt, especially from 60 to 96 h (Fig. 3A and B). Quantitative analysis indicated that the ACT production by \( \Delta \text{Scpdo} \) was about two- to sevenfold higher than that of wt depending on the cultivation time. In addition, \( \Delta \text{Scpdo} \) generated spores significantly earlier than wt and \( \Delta \text{Scpdo}: \text{Scpdo} \) (Fig. 3C). However, the \( \Delta \text{Scpdo}: \text{Scpdo} \) strain exhibited a similar trend of ACT production as wt.

Identification of a transcription factor governing intracellular sulfane sulfur levels

Genome background analysis indicated that there is a hypothetical transcription factor gene (sco0620) adjacent to \( \text{Scpdo} \), which encodes a CsoR-like_DUF156 protein. Hereafter, we refer to this protein and its encoding gene as ScCsoR and \( \text{Sccco} \) respectively. To test whether ScCsoR is involved in the regulation of intracellular sulfane sulfur levels, we constructed a \( \text{Sccco} \) deletion strain \( \Delta \text{Sccco} \) and a complemented strain \( \Delta \text{Sccco}: \text{Sccco} \). SSP4 test showed that \( \Delta \text{Sccco} \) accumulated slightly less sulfane sulfur than wt, whereas \( \Delta \text{Sccco}: \text{Sccco} \) produced slightly more sulfane sulfur than wt. The production differences were more apparent with the HPLC test (Fig. 4A). Moreover, \( \Delta \text{Sccco} \) produced less ACT than wt and its sporulation was also significantly delayed (Fig. 4B).

To test whether ScCsoR functions via regulating the expression of \( \text{Scpdo} \), we analysed the \( \text{Scpdo} \) transcription levels in both \( \Delta \text{Sccco} \) and wt using RT-qPCR. The level of \( \text{Scpdo} \) mRNA in \( \Delta \text{Sccco} \) was much higher than that in wt, with \( \Delta \text{Sccco} \) exhibiting 100- to 480-fold higher levels than wt depending on the growth time (Fig. 5). These results suggested that ScCsoR negatively controlled the expression of \( \text{Scpdo} \).

\text{ScCsoR} is a transcription repressor that binds to the upstream sequence of \( \text{Scpdo} \) promoter

To investigate the mechanism by which ScCsoR controls the expression of \( \text{Scpdo} \), we expressed \( \text{Sccco} \) gene in \( \text{E. coli} \) BL21(DE3) and obtained the purified ScCsoR protein. A 5‘-rapid amplification of cDNA ends (RACE) experiment indicated that \( \text{Scpdo} \) transcription started at \(-12 \text{ bp relative to the start codon GTG}. \) Subsequently, a 256-bp DNA fragment containing the promoter and partial ORF sequence of \( \text{Scpdo} \) was synthesized (Fig. 6A). Purified ScCsoR showed significant affinity to this DNA fragment, as judged by an EMSA experiment (Fig. 6B). To determine the exact binding site of ScCsoR, we synthesized four short DNA fragments (f1-4), with each fragment containing a segment of the 256-bp DNA (Fig. 6A). EMSA experiments indicated that ScCsoR preferred to bind f2 and f3 fragments (Fig. 6C). We used the Multiple EM for Motif Elicitation (MEME; http://meme-suite.org/) toolbox to analyse the f2 and f3 fragments and identified a 16-bp segment containing a pair of palindromic sequences (ATAACn6GGTAT). The 16-bp segment showed a high similarity to the binding sites of Geobacillus thermodenitrificans CsoR (Coyne and Giedroc, 2013; Chang et al., 2014) and Mycobacterium tuberculosis RicR (Festa et al., 2011; Shi et al., 2014; CsoR-like Transcription factor; Fig. 6D), suggesting that it is the binding site of ScCsoR. There are two ScCsoR binding sites in the \( \text{Scpdo} \) promoter. We designated them as bs1 and bsII, located at \(-48 \text{ and } \sim 9 \text{ sites, respectively, from the transcription start site}. \) For further confirmation, we examined the binding affinity of ScCsoR to a 79-bp DNA fragment containing bs1, bsII, as well as an intergenic sequence located between them. A competitor DNA that did not contain these sequences was used as the control. ScCsoR bound to the 79-bp DNA but not to the competitor DNA (Fig. 7A); in addition, when the 79-bp DNA fragment was mixed with 50-fold (mole ratio) of the competitor DNA, ScCsoR still bound specifically to this fragment.

ScCsoR detaches from \( \text{Scpdo} \) promoter in the presence of sulfane sulfur

We performed an EMSA experiment in the presence of hydrogen polysulfide (HS\(_n\)H). The binding affinity of ScCsoR was significantly decreased by HS\(_n\)H (Fig. 7B). Thus, these results indicated that sulfane sulfur reacted with ScCsoR, leading to its detachment from the DNA binding site.
Previous studies have demonstrated that transcription factors CstR, BigR/SqrR, FisR and OxyR all use their cysteine residues to react with sulfane sulfur (Giedroc, 2017; Li et al., 2017b; Hou et al., 2019). ScCsoR contains two cysteine residues, Cys37 and Cys66. To explore which one of these is required to sense sulfane sulfur, we treated the purified ScCsoR with HSnH and analysed the treated protein with LC-MS/MS. As a reference, DTT-treated ScCsoR was also analysed following the same protocol. Two peptides were found in the DTT-treated sample: peptide 1 contained Cys66, and peptide 2 contained Cys37 (Fig. 7C; Figs S3 and S4). Their thiol groups were directly blocked by acetamide (CAM), indicating that these two peptides were not modified. Peptide 1 was also found in the HSnH-treated sample, and a third peptide, peptide 3, which contained Cys37-SS-CAM modification, was found in the HSnH-treated sample but not in the DTT-treated sample (Fig. S5), indicating that Cys37-SH could be modified to Cys37-SSH when ScCsoR was treated with sulfane sulfur.

We then constructed three ScCsoR mutants: ScCsoR-C37S, ScCsoR-C66S and ScCsoR-C37S-C66S, in which the cysteine residue was mutated to serine. EMSA results indicated that these three mutants could still bind to the probe DNA; however, HS₃-S could not detach them from the probe DNA (Fig. S6). These results indicated that both Cys37 and Cys66 were involved in the sulfane sulfur sensing process of ScCsoR.

ScCsoR is a regulator of a sulfane sulfur oxidation gene circuit

We found that the 16-bp binding site is also present in the intergenic region of SccsoR and sco0621 promoters...
To investigate whether a similar gene circuit is present in other Streptomyces species, we analysed all the sequenced *Streptomyces* genomes deposited at the website https://patricbrc.org/, which contained 1777 genomes as ascertained on Nov 26, 2019. ScPDO homologous proteins were found in 1248 genomes. Among them, 773 genomes also contained the genes coding for ScCsoR homologous proteins (CsoR-like TF) close to their *pdo* genes (within 3-ORF distance). We further analysed the genomes of 16 strains that were widely studied due to the production of important antibiotics and various polyketide drugs (Table S1), including the neomycin producer *S. fradiae*, validamycin producer *S. hygroscopicus*, lincomycin producer *S. lincolnensis* and natamycin producer *S. lydicus*. They were all observed to harbour a similar gene circuit containing PDO, TauE, RHOD and CsoR-like TF as that of ScPDO (Fig. 9). These results suggested that the phenomenon of cellular sulfane sulfur affecting polyketide biosynthesis might be widely present in the *Streptomyces* species.

**Distribution of sulfane sulfur oxidation gene circuit in other Streptomyces species**

To investigate whether a similar gene circuit is present in other species of *Streptomyces*, we analysed all the...
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(A) –188

\[ \text{GTACCCAACTTCGGCGTCGTACACCGGAGTTGCCCAGGTCTCGCGGACACAGGC} \]

\[ \text{f1} \]

\[ \begin{array}{c}
\text{–107} \\
\text{GCAGCCGGGCGCGACACTCGGCCCAGGACGCGCGGGGCGGTAGACAGACG} \\
\text{f2}
\end{array} \]

\[ \begin{array}{c}
\text{–76} \\
\text{ScCsoR binding site bsl} \\
\text{ACGACACACCTCTCAATACCTCTGTGGTATATCTGTACGTTGCGCCGGAAG} \\
\text{f3}
\end{array} \]

\[ \text{Transcription start} \]

\[ \text{Scepdo} \]

\[ \begin{array}{c}
\text{ATACCCAGGGGTATTTCCGGAAGGAGACGTGTCGGTACGTTCTCTTCGACC} \\
\text{f4}
\end{array} \]

32

\[ \text{TCCTGGAGACCGAGTCACTGGGCAAACCCGAGCCTATCTGGC} \]

(B)  ScCsoR (µM)

| ScCsoR (µM) |
|-------------|
| 2.2         |
| 8.8         |
| 15.4        |

DNA fragment  
(–188 bp to 68 bp)

(C)  ScCsoR (µM)

| ScCsoR (µM) |
|-------------|
| 15.4        |

| f 1          |

(D)  ATACCC  GGGGGTAT

\[ \begin{array}{c}
\text{bs I} \quad \text{ATACCCAGGGGTAT} \\
\text{CsoR.1} \quad \text{ATACCCAGGGGTAT} \\
\text{bs II} \quad \text{ATACCCAGGGGTAT} \\
\text{CsoR.1} \quad \text{ATACCCAGGGGTAT} \\
\text{RicR.1} \quad \text{ATACCCAGGGGTAT} \\
\text{RicR.2} \quad \text{ATACCCAGGGGTAT} \\
\text{RicR.3} \quad \text{ATACCCAGGGGTAT} \\
\text{RicR.4} \quad \text{ATACCCAGGGGTAT} \\
\text{RicR.5} \quad \text{ATACCCAGGGGTAT} \\
\end{array} \]
PDO. SQR oxidizes H₂S to sulfane sulfur, which is sensed by the gene regulator to elevate the expression of sqr and pdo. Bioinformatic analyses indicate that both H₂S oxidation and sulfane sulfur oxidation gene circuits are widely present in heterotrophic bacteria (Xia et al., 2017; de Lira et al., 2018). Our study indicates that the regulation mechanisms of H₂S oxidation and sulfane sulfur oxidation are different. We observed that all the TFs in the H₂S-oxidizing gene circuits are constitutively expressed at low levels and that these circuits are very sensitive even to low levels of H₂S (< 10 μM; Xia et al., 2017; Li et al., 2017a), suggesting that bacteria containing H₂S-oxidizing gene circuits cannot tolerate the presence of H₂S. However, the expression of TF of the sulfane sulfur oxidation gene circuit is dynamic, which can confer wide amplitude of expression on the genes that it controls – from being completely silenced to be fully turned-on. This makes the strain tolerant to certain levels of intracellular sulfane sulfur – the gene circuit is turned on by a threshold level.

We found that sulfane sulfur affects ACT production as well as spore formation in S. coelicolor M145. Further investigations are needed to illustrate its exact functioning mechanism. For ACT production, hundreds of genes are involved in this process (Xu et al., 2017; Xu et al., 2019). Among them, ActII-ORF4 has been identified as a determining regulator and its expression is regulated by at least eight regulatory proteins (Liu et al., 2013). Our preliminary study indicated that sulfane sulfur cannot directly react with ActII-ORF4, suggesting that sulfane sulfur possibly reacts with a regulatory protein that controls ActII-ORF4 (data not shown). For the generation of

Fig. 6. Characterization of the scpdo promoter.
A. DNA sequence of scpdo promoter. The transcription start site of scpdo was identified using 5'-RACE. GTG is the start codon. The DNA fragments used for EMSA analysis were denoted as f1 to f4.
B. EMSA analysis of the binding affinity of ScCsoR to scpdo promoter. DNA probe (1 nM) was incubated with different amounts of ScCsoR (0, 2.2, 8.8, 15.4 μM). Black arrow indicates the free DNA probe, and red arrow indicates the shifted DNA probe. 50-fold excess of unlabelled competitor DNA and its mixture with probe DNA were also tested.
C. EMSA analysis of the ScCsoR (in reduced form) binding affinity to different parts of scpdo promoter. The DNA probe used in (B) was divided into four DNA fragments (f1 to f4), and other conditions were the same as in (B). Only f2 and f3 fragments exhibited obvious band shifts after incubation with ScCsoR, suggesting that ScCsoR was bound to the promoter region of scpdo gene.
D. Multiple EM for Motif Elicitation analysis of scpdo promoter. The binding sequences of CsoR from Geobacillus thermocatenifllicans and RicR from Mycobacterium tuberculosis, and sequences of f2, f3 fragments were used as inputs for MEME analysis. A 16-nt consensus sequence with reverse palindrome was identified (ATACCn6GGTAT).

Fig. 7. Hydrogen polysulfide (HSₙH) affected the binding affinity of ScCsoR to scpdo promoter.
A. ScCsoR exhibited binding to the probe DNA containing bs1 and bsII, even in the presence of competitor DNA. DNA probe (1 nM) was incubated with different amounts of ScCsoR (0, 2.2, 8.8, 15.4 μM). Black arrow indicates the free DNA probe, and red arrow indicates the shifted DNA probe. 50-fold excess of unlabelled competitor DNA and its mixture with probe DNA were also tested.
B. HSₙH detached ScCsoR from the probe DNA. DNA probe (1 nM) was incubated with different concentrations of ScCsoR (0, 0.5, 1.0, 2.0, 4.0, 8.0, 8.0, 8.0, 8.0 μM). Different amounts of HSₙH (0, 1, 2, 3 mM) were added to the reaction system.
C. LC-MS/MS analysis of HSₙH- and DTT-treated ScCsoR. Details of the MS data are shown in Figs S3–S5.
spores, the regulation is more complex than that of ACT production. Spore formation involves more than a dozen regulatory proteins that compose a multiple-layered regulatory network (Bush et al., 2015). Investigating which of these proteins are affected by sulfane sulfur requires systematic studies, including transcriptomics and/or proteomics approaches. Further, sulfane sulfur levels can also be controlled multiple mechanisms. We observed that sulfane sulfur levels in the DScpdo strain varied with the time of cultivation, indicating that the ScCsoR-pivoted gene cluster is not the only system that acts as an effector. The intracellular redox potential controlled by the ratio of reduced mycothiol/oxidized mycothiol could be another effector (Newton et al., 2008). Thus, the sulfane sulfur metabolism, ACT production and sporulation might be interrelated in S. coelicolor M145.

Our finding is significant with respect to the study of polyketides as well as antibiotics. There are many undeveloped polyketide biosynthetic gene clusters in actinomycetes of which the regulatory mechanisms are yet unknown. Using genetic engineering methods (most commonly by altering their promoters) to activate these gene clusters usually involves laborious work and are not always effective. Sulfane sulfur may represent a new stimulus that can activate certain polyketide biosynthetic gene clusters. Using exogenous sulfane sulfur to treat actinomycetes or altering their intracellular sulfane sulfur might help in the discovery of new polyketide drugs from such bacteria.

**Experimental procedures**

**Strains and growth conditions**

All the strains and plasmids used in this study are listed in Table S2. S. coelicolor M145 and its derivatives were cultured at 30 °C. Mannitol soya flour (MS) agar medium was used for sporulation and conjugation. YBP (yeast-beef-peptone) solid medium was used for determining ACT production, phenotypic observation, growth analysis and RNA extraction. YBP liquid medium was used for
quantifying intracellular sulfane sulfur. Antibiotics were added to the media when required. *E. coli* DH5α was used for plasmid construction, and the *E. coli* BL21 (DE3) strain was used for protein expression. *E. coli* ET12567 (pUZ8002) was used to transfer nonmethylated DNA into *S. coelicolor* M145.
amplified and downstream (1131 bp) regions of editing system (Huang et al., 2015). Upstream (1008 bp) and downstream (1131 bp) regions of Scpdo ORF were amplified from S. coelicolor M145 genomic DNA. A 20-nt NGG mode sequence (5′-ACCGGCGGTTCGCTGCTGAT-3′) was amplified using the sgRNA-guided CRISPR-Cas9 genome editing system (Table S3). The mutant, ΔScpdo, of S. coelicolor was constructed using only the medium. ΔScpdo was obtained using PCR. This DNA fragment was inserted into the integrative vector pMS82 to make a new plasmid pCom0618 (Table S2), which was then transformed into the ΔScpdo strain using a conjugation method. The ΔSccsoR complementary strain, ΔSccsoR::Scpdo, was constructed following the same protocol, except that a DNA fragment containing SccsoR ORF and its native promoter (a 507 bp fragment upstream of its ORF) was used. To construct control strains, empty pMS82 vectors were also transformed into ΔScpdo and ΔSccsoR strains.

**Construction of S. coelicolor ΔScpdo and ΔSccsoR**

All the primers used in this study are listed in Table S3. The mutant, ΔScpdo, of S. coelicolor, was constructed using the sgRNA-guided CRISPR-Cas9 genome editing system (Huang et al., 2015). Upstream (1008 bp) and downstream (1131 bp) regions of Scpdo ORF were amplified from S. coelicolor M145 genomic DNA. A 20-nt NGG mode sequence (5′-ACCGGCGGTTCGCTGCTGAT-3′) was obtained from CRISPy-web (http://crispy.secondarymetabolites.org/). A special single guide RNA (sgRNA; containing a 20-nt sequence) was amplified from pKCCas9dO. The linear pKCCas9dO vector was generated by SpeI/HindIII digestion. The upstream and downstream DNA fragments, sgRNA, and the linear pKCCas9dO vector were finally assembled to generate the plasmid pKCCas9dO-sco0618 using the ClonExpress™ II One Step Cloning Kit (TaKaRa). This plasmid was introduced into S. coelicolor M145 by conjugal transfer between E. coli ET12567 (pJZ8002) and M145. The exconjugants were selected and examined following a reported protocol (Li et al., 2017a). After incubation at 37 °C for several rounds of selection, the pKCCas9dO-Scpdo plasmid was extracted from the mutant and the final construct, ΔScpdo, was obtained. The strain ΔSccsoR was subsequently constructed using a homologous recombination method (Lu et al., 2018).

**Construction of the complementary strains**

To construct a complementary strain of ΔScpdo, namely, ΔScpdo::Scpdo, a DNA fragment containing a kasOp′ promoter (Bai et al., 2015) and Scpdo ORF was obtained using PCR. This DNA fragment was inserted into the integrative vector pMS82 to make a new plasmid pCom0618 (Table S2), which was then transformed into the ΔScpdo strain using a conjugation method. The ΔSccsoR complementary strain, ΔSccsoR::Scpdo, was constructed following the same protocol, except that a DNA fragment containing SccsoR ORF and its native promoter (a 507 bp fragment upstream of its ORF) was used. To construct control strains, empty pMS82 vectors were also transformed into ΔScpdo and ΔSccsoR strains.

**Examination of the ACT production**

For testing the effects of H2S and sulfane sulfur, S. coelicolor M145 spores (2 × 10⁶) were inoculated on YBP agar plates containing different concentrations of NaHS or Sb₂, and the plates were incubated at 30 °C for 6 days. For quantitative analysis of the ACT production, a previously reported method was used (Kieser et al., 2000). Briefly, total ACT was quantified by scraping off all the medium containing mycelia from the plate and mixing it with 1 M KOH. The reaction was conducted overnight at 25 °C, and later, 1 ml aliquot was taken out for quantification. Intracellular ACT was quantified using only the mycelia, and extracellular ACT was quantified using only the medium. ACT in the sample was quantified by measuring absorbance at 640 nm.

**Analysis of intracellular sulfane sulfur**

To perform the SSP4 test, S. coelicolor M145 and its derivatives (10 ml) were cultured in liquid YBP medium. The mycelia were harvested at different time points, washed with and finally suspended in HEPES buffer (50 mM, pH 7.4). Cells were broken down using a high-pressure crusher SPCH-18 (STANSTED). SSP4 (10 μM) was added to each supernatant, and the mixture was incubated at 37 °C for 15 min in the dark with gently shaking (125 rpm). Fluorescence intensity was measured using the Synergy H1 microplate reader. The excitation and emission wavelengths were set at 482 and 515 nm respectively. The final data were converted to fluorescence intensity per 0.5 OD₅₄₀ of cells.

For HPLC analysis, a method previously was used (Ran et al., 2019). Briefly, cells at 1 OD₄₅₀ were collected and washed with HEPES buffer (50 mM, pH 7.4) and then resuspended in 220 μl of 50 mM Tris-HCl buffer (pH 9.5) containing 1% (v/v) Triton X-100, 50 μM DTPA, and 1 mM sulfite. The samples were incubated at 95 °C for 10 min to convert sulfane sulfur to thiosulfate. After centrifugation, 50 μl supernatant was reacted with mBBr (5 μl) in the dark for 30 min, and 100 μl of acetic acid and acetonitrile mixture (v/v, 1:9) was added...
to stop the reaction. The finally obtained sample was analysed using HPLC equipped with a fluorescence detector.

Protein mutation, expression, and purification

The ORFs of Scpdo and SccsoR were amplified from S. coelicolor M145 genomic DNA. ScCsoR mutants were constructed using the revised QuikChange™ method (Xia et al., 2015). The ORFs were ligated into pET15b plasmid (Table S2) for expression. E. coli BL21(DE3) cells containing pET15b derived plasmids were cultured in LB medium at 37 °C until OD600nm reached about 0.5, and then 0.5 mM isopropyl β-D-1-thiogalactopyranoside (IPTG) was added. The temperature was changed to 30 °C, and the cultivation was continued overnight. Cells were harvested by centrifugation and disrupted using a pressure cell homogenizer (SPCH-18) at 4 °C. ScPDO and different concentrations of GSSH (0–0.5 mM) in 3 ml KPi buffer (100 mM, pH 7.4), and the reaction was conducted at 25 °C for 15 min, followed by separation on an 8% (w/v) non-denaturing polyacrylamide gel in an ice bath. Next, the DNA and proteins were transferred from the gel to a positively charged nylon membrane, fixed, blocked, washed and finally stained using solutions of an ECL Western blotting kit (GE Healthcare), and images were captured with a FlourChemQ system (Alpha Innotech).

Assay of ScPDO activity

The persulfide dioxygenase activity of ScPDO was determined using a previously reported method (Xin et al., 2016). The reaction system contained 1 μM ScPDO and different concentrations of GSSH (0–0.5 mM) in 3 ml KPi buffer (100 mM, pH 7.4), and the reaction was conducted at 25 °C. O2 consumption was examined using an Orion RDO meter (Thermo Scientific). The kinetic parameters were calculated using the Michaelis–Menten equation as reported previously (Liu et al., 2014). GSSH was prepared as described previously (Xin et al., 2016).

RT-PCR, RT-qPCR and 5’-RACE analysis

RNA from S. coelicolor was prepared by following a previously described protocol (Lu et al., 2018). Mycelia were harvested at different time points (36, 60 and 84 h) and were ground into powder using liquid nitrogen. Total RNA was treated with TRIzol, and DNase I was used to obtain chromosomal DNA. RT-PCR was carried out using a reverse transcriptase kit (Invitrogen). RT-qPCR was performed using SYBR Premix Ex Taq (Takara). hrdB gene encoding a major sigma factor was used as control to normalize the relative quantities of cDNA. Roche LightCycler 480 thermal cycler was used to determine the melting curve and specificity of the PCR products. Three independent replicates were performed in parallel. 5’-RACE was performed employing the First-Choice RLM-RACE kit (Thermo) following the manufacturer’s instructions.

Electrophoretic mobility shift assay (EMSA)

DNA probes were labelled with 5’-Biotin, and 1 nM each of the labelled probes was mixed with differing amounts of purified ScCsoR protein in a binding buffer (20 mM Tris-HCl, 2 mM EDTA, 20 mM KCl, 0.5 mM dithiothreitol (DTT) or 1 mM–3 mM HSnH, 4% (w/v) Ficol-400, pH 8.0). HSnH was prepared by mixing H2S with Sn8 as reported previously (Xin et al., 2016). The reaction was conducted at 25 °C for 15 min, followed by separation on an 8% (w/v) non-denaturing polyacrylamide gel in an ice bath. Next, the DNA and proteins were transferred from the gel to a positively charged nylon membrane, fixed, blocked, washed and finally stained using solutions of an ECL Western blotting kit (GE Healthcare), and images were captured with a FlourChemQ system (Alpha Innotech).

LC-MS/MS analysis of ScCsoR

The purified ScCsoR was reacted with 10-fold (molar ratio) of hydrogen polysulfide (HSnH, n ≥ 2) or DTT for 20 min at room temperature. The reacted protein was treated with a denaturing buffer (0.5 M Tri-HCl, 2.75 mM EDTA, 6 M guanidine-HCl, pH 8.0) containing 1 M iodoacetamide (IAM) for 1 h, and the sample was subsequently digested with trypsin (1:25, w/w) at 37 °C for 4 h. LC-MS/MS analysis was conducted following a previously reported protocol (Li et al., 2017b). The prominence nano-LC system (Shimadzu) equipped with a custom-made silica column (75 μm × 15 cm) packed with 3-μm Reprosil-Pur 120 C18-AQ was used for the analysis. For the elution process, a total gradient of 100 min from 0% to 100% of solvent B (0.1% (v/v) formic acid in 98% (v/v) acetonitrile) at 300 nl min⁻¹ was used; solvent A was 0.1% (v/v) formic acid in 2% (v/v) acetonitrile. The eluent was ionized and electrosprayed via LTQ-Orbitrap Velos Pro CID mass spectrometer (Thermo Scientific); the run was performed in a data-dependent acquisition mode with Xcalibur 2.2.0 software (Thermo Scientific). Full-scan MS spectra (from 400 to 1800 m/z) were detected using the Orbitrap at a resolution of 60,000 at 400 m/z.

Bioinformatics analysis

ScPDO and ScCsoR were used as queries to search homologues from all the sequenced Streptomyces genomes mentioned in the website, https://patricbrc.org. The selection criteria used were E-value < 1e⁻60.
identity ≥ 35% and coverage ≥ 70%. ClustalW was used for alignment of these candidates, and a condensed neighbour-joining tree was further built with bootstrap replications at 1000 by using the p-distance method, uniform rates and pairwise deletion with a cut-off at 50% by using the MEGA 7.0 software (Kumar et al., 2016).

Acknowledgements

The work was financially supported by grants from the National Key Research and Development Program of China (2016YFA0601103) and the National Key R&D Program of China (2018YFA0901200).

Conflict of interest

The authors declare no conflict of interest.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**Fig. S1.** SDS-PAGE analysis of recombinant ScPDO. Lane 1 is the ladder (kDa), Lanes 2–4 are ScPDO with different concentrations. The theoretical MW of His-tag-ScPDO is about 49 kDa.

**Fig. S2.** Amino acid sequence alignment analysis of ScPDO and known type III PDOs. The reported type III PDOs usually contain PDO and rhodanese domains. The representative type III PDOs amino acid sequences of *Zunongwanga profunda* SM-A87 (ZpPDOIII, ADF52140.1), *Staphylococcus aureus* (SaPDOIII, WP_000465474.1) and *Bacillus cereus* ATCC 10876 (BcPDoll, EKE94737.1) were downloaded from NCBI and were compared with ScPDO by using ClustalW. The results show that ScPDO is a typical type III PDO, which is consistent with a previous report (Xia et al., 2017).

**Fig. S3.** MS2 data of peptide 1, which was from DTT-treated ScCsoR.

**Fig. S4.** MS2 data of peptide 2, which was from DTT-treated ScCsoR.
**Fig. S5.** MS² data of peptide 3, which was only detected from polysulfide-treated ScCsoR.

**Fig. S6.** EMSA assay of ScCsoR-C37S, ScCsoR-C66S and ScCsoR-C37S-C66S mutants. The Cys-to-Ser mutation did not affect the ScCsoR binding activity to Scpdo promoter; however, it caused the loss of HSₐS sensing activity and HSₐS could not release ScCsoR mutants from Scpdo promoter. The promoter region of Scpdo (256 bp) was used as the DNA probe. 1 nM DNA probe was incubated with gradient concentration of ScCsoR mutants (0, 0.5, 1.0, 2.0, 4.0, 8.0, 8.0, 8.0, 8.0 μM), different amounts of polysulfides (0, 1, 2, 3 mM) was added to the reaction system (with equal amounts of ScCsoR mutants-8.0 μM). The black arrow indicates the freedom DNA probe, the red arrow indicates the shifted DNA probe. These results show that the cysteines on ScCsoR play an important role for sulfane sulfur sensing, and the two cysteine are indispensable.

**Fig. S7.** The intergenic region of SccsoR and Scrhod promoters. The starting codon is shown in bold black, binding site is shown in red font. The number below is the length (bp) between the starting codon and binding site.

**Fig. S8.** Sco0622 and sco0623 expression was not controlled by ScCsoR. (A) Co-transcription analysis of Scpdo-tauE, rhod-sco0622 and sco0622-23. Genomic DNA, mRNA, and cDNA as templates for lanes 1, 2, and 3, respectively. (B) EMSA analysis of ScCsoR binding activity to the promoter region of sco0622 (242bp), 1 nM DNA probe was incubated with different amounts of ScCsoR (0, 2.2, 8.8, 15.4, 22.2 μM), the black arrow indicates the unbounded DNA probe. (C) RT-qPCR analysis of the transcriptional level of sco0622 in wt and ΔSccsoR strains. RNA was harvested at 36 h, 60 h, 84 h from mycelium grown in YBP solid medium. The hrdB gene was used as the internal control to normalize the mRNA level. Three independent measurements were carried out and error bars indicate the standard deviations.

**Table S1.** The distribution of Scpdo-rhod in representative Streptomyces genomes.

**Table S2.** Strains and plasmids used in this study.

**Table S3.** Primers (5′→3′) used in this study.