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RNA-Seq Analysis Reveals a Six-Gene SoxR Regulon in *Streptomyces coelicolor*

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Abstract



The redox-regulated transcription factor SoxR is conserved in diverse bacteria, but emerging studies suggest that this protein plays distinct physiological roles in different bacteria. SoxR regulates a global oxidative stress response (involving > 100 genes) against exogenous redox-cycling drugs in *Escherichia coli* and related enterics. In the antibiotic producers *Streptomyces coelicolor* and *Pseudomonas aeruginosa*, however, SoxR regulates a smaller number of genes that encode membrane transporters and proteins with homology to antibiotic-tailoring enzymes. In both *S. coelicolor* and *P. aeruginosa*, SoxR-regulated genes are expressed in stationary phase during the production of endogenously-produced redox-active antibiotics. These observations suggest that SoxR evolved to sense endogenous secondary metabolites and activate machinery to process and transport them in antibiotic-producing bacteria. Previous bioinformatics analysis that searched the genome for SoxR-binding sites in putative promoters defined a five-gene SoxR regulon in *S. coelicolor* including an ABC transporter, two oxidoreductases, a monooxygenase and an epimerase/dehydratase. Since this *in silico* screen may have missed potential SoxR-targets, we conducted a whole genome transcriptome comparison of wild type *S. coelicolor* and a *soxR*-deficient mutant in stationary phase using RNA-Seq. Our analysis revealed a sixth SoxR-regulated gene in *S. coelicolor* that encodes a putative quinone oxidoreductase. Knowledge of the full complement of genes regulated by SoxR will facilitate studies to elucidate the function of this regulatory molecule in antibiotic producers.

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Data Availability: The authors confirm that all data underlying the findings are fully available without restriction. The data supporting the results of this work are available in Supporting Information and in NCBI's Gene Expression Omnibus and accessible through GEO series accession number GSE57268 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc = GSE57268).

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Introduction

The redox-regulated transcription factor SoxR is present in a diverse range of Proteobacteria and Actinobacteria and homologs are highly similar at the amino acid level [1]. SoxR homologs function as homodimers and have a conserved amino-terminal helix-turn-helix DNA binding domain, suggesting that these proteins bind to and regulate transcription from similar operator sequences. This has been confirmed in organisms where SoxR has been biochemically characterized [2-7]. SoxR homologs also share a conserved sequence (CysX2CysXCysX5Cys) in the carboxy-terminus that has been shown to be necessary for coordinating [2Fe-2S] centers in SoxR proteins from Escherichia coli, Pseudomonas aeruginosa, and Streptomyces coelicolor [5,8,9]. These [2Fe-2S] clusters are central to SoxR's ability to detect changes in the cellular redox environment and regulate gene expression in response. SoxR was originally discovered in the enterobacterium E. coli where it promotes resistance to redoxcycling drugs like paraquat and menadione [10,11]. Subsequent studies revealed that in this microorganism SoxR mediates its effects in a two-step process. Upon sensing redox stress via its [2Fe-2S] clusters, SoxR activates the expression of a second transcription factor, soxS [12,13]. SoxS, an AraC-type regulator then recruits RNA polymerase to the promoters of >100 genes (the SoxRS regulon), whose protein products cumulatively restore redox homeostasis and repair oxidant-induced cellular damage [14].

The E. coli SoxRS regulon is conserved in other enterobacteria where it functions to confer generalized protection against exogenous redox-cycling compounds. Various lines of research conducted in the past decade indicate that this function may be limited to members of the Enterobacteriaceae, and that the SoxR regulatory network is different in other bacteria. A comprehensive bioinformatic survey of sequenced bacterial genomes revealed that while a soxR homolog is detected in 176 genomes, a soxS homolog is present only in enteric bacteria where it appears to be the solitary gene directly regulated by SoxR [1]. The same study showed that in non-enterics (all of which lack soxS), SoxR is predicted to directly regulate a small number of genes. In further contrast to the apparent function of SoxR in enteric bacteria, none of the putative SoxR targets in non-enterics encode proteins that are typically involved in oxidative stress detoxification and repair. Instead these genes encode membrane transporters and enzymes with homology to proteins that modify small molecules, including antibiotics. The absence of a soxS homolog and the predicted SoxR regulons in non-enterics has given rise to the notion that SoxR does not regulate a generalized oxidative stress response in the majority of bacteria. This has been corroborated for *Pseudomonas putida*, *P. aeruginosa*, and *S. coelicolor*, where deletion of *soxR* does not result in increased sensitivity to redoxcycling drugs when compared to the parental strain [5,9,15,16]. While the function of SoxR in *P. putida* remains unknown, SoxR regulates gene expression in response to redox-active endogenously produced antibiotics in both *P. aeruginosa* and *S. coelicolor*, indicating that SoxR plays a role that is intimately tied in with the physiology of these organisms [5,6,9,17,18].

The SoxR regulon in P. aeruginosa is induced during the production of phenazine antibiotics in stationary phase, and consists of two membrane transporters (encoded by PA4205-PA4208 and PA3718) and a monooxygenase enzyme (PA2274) [15,17]. The S. coelicolor SoxR regulon is also induced in stationary phase during the production of the benzochromanequinone blue-pigmented antibiotic actinorhodin (Act), and is similar to the *P. aeruginosa* regulon in encoding an ABC-type membrane transporter (SCO7008) and four redox enzymes (SCO1178, SCO1909, SCO2478, SCO4266) [5,6]. It is noteworthy that two of these genes encode products that are similar to enzymes that catalyze tailoring steps in the Act biosynthetic pathway. Specifically, SCO4266 is similar to the oxidoreductases ActVI-ORF2 (37% identity; 49% similarity) and ActVI-ORF4 (34% identity; 44% similarity), and SCO1909 resembles the monooxygenase ActVA-ORF6 (38% identity; 55% similarity) [19]. SCO1909 is also similar to PA2274 (37% identity; 45% similarity), the SoxR-regulated monooxygenase in P. aeruginosa [19]. An independent study had also described four of the SoxRtargets in S. coelicolor as genes whose expression is temporally coordinated with Act (eca), and whose levels are reduced in an Actnon producing strain: SCO7008 (ecaA), SCO1909 (ecaB), SCO1178 (ecaC), SCO4266 (ecaD) [20]. The composition of the P. aeruginosa and S. coelicolor SoxR regulons, and the fact that they are induced by endogenously produced redox-active antibiotics, suggests that SoxR evolved to sense endogenous metabolites and activate machinery to process and transport them in these two phylogenetically divergent bacteria. This notion is further supported by the observation that both P. aeruginosa and S. coelicolor soxR mutants display de-regulated antibiotic production/secretion [1.6].

Both the SoxR regulon in *P. aeruginosa*, and the SoxRS regulon in *E. coli* were characterized using microarray-based expression profiling [14,15]. The five-gene SoxR regulon in *S. coelicolor*, on the other hand, was identified by conducting an *in silico* search of the genome for SoxR-binding sites located upstream of predicted open reading frames (ORFs) [1,5,6]. Given the morphological complexity and large genome size of *S. coelicolor* (8.7 megabases), it is possible that the bioinformatic screen for SoxR-regulated genes may have missed potential targets. This

analysis would also have failed to identify genes that are indirectly regulated by SoxR (i.e. via an intermediate transcription factor as in *E. coli*). In this study we conducted a whole genome transcriptome comparison of the wild type *S. coelicolor* strain M145 and a *soxR* null mutant using RNA-Seq with the goal of gaining a comprehensive picture of the SoxR regulatory network in this medically important antibiotic-producer. This analysis revealed a sixth SoxR-regulated gene in *S. coelicolor*, in addition to the five confirmed targets.

Materials and Methods

Bacterial strains, culture conditions and RNA isolation

The S. coelicolor strains used in this study are listed in **Table 1**. For RNA-Seq and qRT-PCR experiments, about 10⁸ spores were cultured on R2YE plates [21] that were overlaid with cellophane and grown at 30°C for either 24 h (prior to the production of Act) or 72 h (when Act-proficient cells produce Act), with biological replicates obtained for the 72 h cultures. To harvest, cells were incubated with RNAprotect bacterial reagent (Qiagen) for 5 min at room temperature, scraped off the cellophane, pelleted by centrifugation for 10 min at 5,000×g, and frozen at -80° C. Cells were lysed by incubation for 15 min at 30°C in TE buffer containing 15 mg/mL lysozyme, followed by 30 s of sonication on ice. Total RNA was extracted with an RNeasy plant minikit (Qiagen) according to manufacturer's instructions. Contaminating DNA was removed by treating for 1 h at 37°C with 5 units of RNase-free DNase I (Qiagen). The RNA preparation was subject to one extraction with acidified phenol-chloroform, followed by an extraction with chloroform-isoamyl alcohol. RNA was ethanol precipitated overnight at -80° C, washed with 80% ethanol and resuspended in nuclease-free water. RNA purity and concentration were determined using a Nanodrop or Qubit spectrophotometer. RNA quality was assessed by agarose gel electrophoresis or a Nano Bioanalyzer. The absence of contaminating DNA was confirmed by the absence of product following a 30-cycle PCR reaction using RNA as template and *hrdB* primers (**Table S1**).

Library construction and RNA sequencing

Library construction and sequencing of RNA transcripts was performed by Fasteris SA (Switzerland). Briefly, RNA samples were treated to reduce ribosomal RNA levels using the Ambion MICROBExpress kit. RNA transcripts were fragmented using a buffered zinc solution. The first cDNA strand was synthesized by reverse transcription using random primers in the presence of dUTPs. After second strand synthesis and adapter ligation, the first cDNA strand was digested with uracil-DNA glycosylase. The remaining fragments were PCR amplified and 150–200 bp amplicons were selected by polyacrylamide gel electrophoresis. The resulting cDNAs underwent high-throughput sequencing in

Table	1. Streptomyces	coelicolor	strains	used i	in this	study.

Strain	Genotype or description	Reference
M145	SCP1 ⁻ SCP2 ⁻ derivative of A3(2)	[21]
M511	Δact II-ORF4 derivative of M145	[32]
M145-1A	Markerless $\Delta soxR$ derivative of M145	[5]
145/pSET152	M145 transformed with pSET152	[5]
$\Delta sox R/pSET152$	M145-1A transformed with pSET152	[5]
$\Delta sox R/pSox R$	M145-1A transformed with pSET152::soxR	[5]

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Table 2. Genes identified by RNA-Seq as SoxR-dependent in stationary phase^a.

SCO number	Decrease in $\Delta soxR$ v	ersus WT on Day 3 ^b	Increase in WT on	Day 3 vs. Day1 ^c	Predicted function ^e
	Fold change	q-value ^d	Fold Change	q-value ^d	
0319	84	2e-28	>100	9e-12	Hypothetical protein
0320	79	3e-21	52	5e-10	Quinone oxidoreductase
1178 ^f (ecaC)	52	2e-21	49	1e-9	NAD-dependent epimerase/dehydratase
4266 ^f (ecaD)	30	2e-28	14	4e-7	Oxidoreductase
7688 ^g	28	8e-2	1	1e0	Hypothetical protein
1177	23	7e-18	32	2e-7	GntR-family transcriptional regulator
0321	18	7e-18	26	9e-8	Carboxylesterase
1909 ^f (ecaB)	18	1e-22	11	7e-6	Monooxygenase
1734	15	4e-11	24	2e-5	Secreted cellulose binding protein
4021	11	6e-2	2	8e-1	Two-component histidine kinase
2478 ^f	7	3e-1	>100	1e-3	Flavoprotein reductase
7682 ^g	6	4e-1	<1	8e-1	Non-ribosomal peptide synthase
4157	6	4e-1	4	4e-1	Protease
4020	6	3e-3	2	8e-1	Two-component response regulator
6165	6	3e-1	>100	8e-4	Hypothetical protein
1697 ^h	5	8e-3	1	1e0	SoxR
2878 ⁹	5	4e-1	<1	5e-1	Hypothetical protein
7008 ^f (ecaA)	5	3e-9	37	2e-9	ABC transporter

^aGenes are organized in decreasing order of SoxR-dependence as determined by RNA-Seq.

^bRNA for this comparison was obtained from 3-day old WT or *\DeltasoxR* cultures, both of which were blue-pigmented.

^cRNA for this comparison was obtained from 3-day old WT (blue-pigmented) or 1-day old WT (unpigmented) cultures.

^dFalse discovery rate adjusted q-values were calculated according to the Benjamini-Hochberg procedure [27].

^ePredicted functions of genes were obtained from StrepDB (http://strepdb.Streptomyces.org.uk).

^fConfirmed SoxR-targets that are directly regulated by SoxR in response to Act production [5,6].

⁹These genes are unlikely to be Act-dependent since they were not differentially expressed between 3-day old versus 1-day old WT samples, and were not considered for further analysis.

^hSCO1697 is soxR which is constitutively expressed over the course of development, and is not autoregulated [5].

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an Illumina HiSeq 2000 sequencer to obtain single reads of 100 bp using the forward sequencing primer.

was performed to identify related sequences and elucidate potential gene function.

Read mapping and differential expression

The sequenced reads were mapped on to the *S. coelicolor* genome [22] using BWA v0.5.9 [23] and read counts for each annotated transcript were compiled using BEDtools without regard to strand [24], on the web-based platform Galaxy [25–27]. For both the wild type and $\Delta soxR$ libraries, ~97% of total reads were mapped to the *S. coelicolor* genome. Differential gene expression was assessed using DESeq with library sizes normalized by the median count ratios across transcripts and false discovery rate adjusted q-values calculated according to the Benjamini-Hochberg procedure [28]. The data supporting the results of this work are available in NCBI's Gene Expression Omnibus and accessible through GEO series accession number GSE57268 (http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc = GSE57268).

Re-annotation of SCO0319, SCO0320, SCO0321 region

To identify alternative potential transcripts in the region surrounding *SCO0319-SCO0321* the *S. coelicolor* genome region spanning *SCO0318* to *SCO0321* (318329–322255) was scanned for open reading frames (ORFs) using Geneious (v.7.1.4, Biomatters). Large ORFs (>500 bp) that spanned the region of *SCO0319* to *SCO0320* were selected and a BLAST search [29]

Quantitative RT-PCR

The cDNA templates for qRT-PCR were generated from total RNA with iScript (Bio-Rad). The primers used for qRT-PCR (Integrated DNA Technologies) were designed using Primer3 software [30], with a melting temperature of 60°C, length of ~20-nt, and amplicon length of ~100 bp (**Table S1**). Each qRT-PCR reaction (20 μ L) contained 25 ng cDNA, 250 nM each of forward and reverse primer, and 10 μ L Power Sybr green PCR master mix (Applied Biosystems). qRT-PCR reactions were carried out in a StepOne PCR machine (Applied Biosystems) with the following reaction parameters: 10 min at 95°C; 40 two-step amplification cycles with 15 s denaturation at 95°C and 1 min annealing and extension at 60°C; final dissociation stage for 15 min to generate a melting curve and verify specificity of amplification products. Samples were assayed in duplicate and the target signal standardized to the level of the housekeeping sigma factor *hrdB*.

Electrophoretic mobility shift assay (EMSA)

A gel mobility shift assay was used to assess the binding of purified SoxR to promoter DNA. The method of Dela Cruz *et al.* [5] was used to purify histidine-tagged SoxR and generate DIG-labeled DNA probes. Primers used to PCR amplify the promoter DNA fragments are listed in **Table S1**. For the Table 3. Validation of RNA-Seq results by quantitative RT-PCR^a.

<i>SCO</i> Number	Fold decrease in $\Delta sox R$ versus WT ^b	Fold decrease in Δact versus WT ^c	Predicted function
Potential SoxR-t	argets ^d		
0319	4	24	Hypothetical protein
0320	86	34	Quinone oxidoreductase
0321	59	61	Carboxylesterase
1177	9	11	GntR-family transcriptional regulator
1734	8	3	Secreted cellulose binding protein
4020 ^e	1	1	Two-component response regulator
4021	10	2	Two-component histidine kinase
4157 ^e	1	2	Protease
6165 ^e	1	<1	Hypothetical protein
soxR and confirm	med SoxR-targets		
1697	47	1	SoxR
1178 (ecaC)	60	29	NAD-dependent epimerase/dehydratase
1909 (ecaB)	30	30	Monooxygenase
2478	31	15	Flavoprotein reductase
4266 (ecaD)	29	59	Oxidoreductase
7008 (ecaA)	15	30	ABC transporter

^aRNA for qRT-PCR validation was obtained from independent biological samples.

^bDifferential gene expression in WT and *\DeltasoxR* in 3-day old cultures assessed by qRT-PCR. Gene expression was standardized to the housekeeping sigma factor, *hrdB*, and normalized to WT.

^cDifferential gene expression in WT and the Δact strain (M511) in 3-day old cultures assessed by qRT-PCR. Gene expression was standardized to the housekeeping sigma factor, *hrdB*, and normalized to WT.

^dGenes are arranged in order of increasing SCO number

^eThese genes did not demonstrate SoxR- or Act-dependent expression in the qRT-PCR validation assay and were eliminated from further analysis.

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binding reaction, 6 fmol of DNA probe was incubated with SoxR (0 to 20 nM) for 15 min at 25°C in binding buffer (10 mM Tris-HCl, pH 8, 75 mM KCl, 0.1 mM dithiothreitol, 10% glycerol, 2 mM MgCl₂, 0.1 μ g poly(dI)-poly(dC), 2 mM dAMP) in a total volume of 30 μ L. For the competition assay, 3,000 fmol of unlabeled probe (specific competitor) was added to the reaction mixtures. Protein-bound and uncomplexed DNA products were separated on a 5% polyacrylamide gel (Tris-HCl, pH 8, 3.3 mM sodium acetate, pH 7.9, 1 mM EDTA, 2% glycerol) that was run at 6°C and 180 V for 90 min. The DNA was transferred to nylon membranes (Roche), cross linked by UV, and detected using a DIG gel shift 2nd-generation kit (Roche) according to manufacturer's instructions.

Reverse Transcription PCR (RT-PCR)

RT-PCR reactions were carried out on total RNA using the OneStep RT-PCR kit (Qiagen) according to manufacturer's instructions. All RT-PCR reactions consisted of 30 cycles using primers listed in **Table S1**. Reactions were analyzed on TAE: agarose gels and nucleic acids were visualized using ethidium bromide.

Bioinformatic analysis of Streptomyces genomes for potential SoxR-regulated genes

Whole genome sequences of *Streptomyces* species downloaded from the NCBI genomes database (ftp://ftp.ncbi.nih.gov/ genomes/) were searched for potential homologs of the genes of the *Streptomyces coelicolor* SoxR regulon using tblastn [29], retaining only the best matching genomic region for each gene. If two of the regulon genes matched to the same region of the genome, only the best matching gene was reported. For each retained matching sequence, the region lkb upstream was searched for the soxbox binding motif [1] using a position-specific scoring matrix applied via the BioPython toolkit [31]. The nucleotide frequencies for each target genome were used to calculate expected background match rates, and matches with a log-odds score for the soxbox motif greater than 10 are reported.

Results

RNA-Seq analysis reveals several novel genes as potential SoxR targets

The previously described five-gene SoxR regulon in S. coelicolor was identified bioinformatically [5,6]. In order to identify other potential SoxR-regulated genes that might have been missed by this bioinformatic approach, we conducted RNA-Seq analysis to detect genes differentially expressed between wild type and a $\Delta soxR$ mutant strain. RNA substrates were extracted from hyphae grown on R2YE medium. Because Act (or a precursor) is a known physiological activator of SoxR in S. coelicolor [5,6], samples were collected three days post-inoculation when cells were actively producing this blue-pigmented antibiotic. As a control, samples were also collected 24 h post-inoculation, when no pigmented antibiotics were visible (and SoxR is quiescent). To identify genes that were potentially both SoxR- and Act- dependent, we focused on those that met two criteria: (i) were differentially expressed between wild type and $\Delta soxR$ in 3-day old cultures (Act produced); (ii) were also differentially expressed between 3-day old (blue) and 1-day old (unpigmented) wild type cultures. As predicted, the five established members of the SoxR regulon (ecaA-ecaD, SCO2478) were all significantly overexpressed in wild type compared to the



Figure 1. Complementation analysis to confirm SoxR-dependence of genes identified by RNA-Seq. qRT-PCR was performed on RNA isolated from WT/pSET152, *ΔsoxR*/pSET152, and a *ΔsoxR* strain complemented with wild-type *soxR* (pSoxR), that were grown on R2YE plates for 3 days. The expression levels of all genes were standardized to the level of the constitutively expressed housekeeping sigma factor, *hrdB*, and normalized to expression in WT/pSET152. The results represent the means and standard deviation (bars; some are not visible on this scale) of four independent experiments. doi:10.1371/journal.pone.0106181.g001

 $\Delta soxR$ mutant (**Table 2**). Each showed a ≥4-fold differential expression in wild type versus $\Delta soxR$, and a q-value of ≤0.5. Using these parameters (≥4-fold differential expression and a q-value of ≤0.5), twelve novel genes were discovered to be potentially upregulated by SoxR in stationary phase (**Table 2**). Of these twelve genes, three (*SCO7688, SCO7682, SCO2878*) did not display differential expression between 1-day and 3-day old wild type cultures (**Table 2**). These genes are thus unlikely to be Act-dependent and were eliminated from further analysis.

Quantitative real-time PCR confirms SoxR- and Actdependence in a subset of genes identified by RNA-Seg

To validate the RNA-Seq results, the SoxR-dependence of the nine newly identified genes was analyzed by quantitative real time PCR (qRT-PCR). RNA was obtained from independent biological samples (wild type and $\Delta soxR$) following growth for three days on complex agar (R2YE) medium. Six of the nine newly identified genes were significantly overexpressed in wild type as compared to $\Delta soxR$ (\geq 4-fold difference) by qRT-PCR analysis: *SCO0319*, *SCO0320*, *SCO0321*, *SCO1177*, *SCO1734*, *SCO4021* (**Table 3**). The other three genes that were tagged as SoxR-

dependent by RNA-Seq (SCO4020, SCO4157, SCO6165) did not demonstrate differential expression between wild type and $\Delta soxR$ in the qRT-PCR assay and were eliminated from further analysis (**Table 3**).

Since SoxR is transcriptionally active only in Act-producing cells, SoxR-dependent genes should also demonstrate Act-dependence. This is true for the five previously confirmed SoxR targets (**Table 3**). To determine if the six new candidate SoxR-targets demonstrate Act-dependence, their expression levels were compared in 3-day old wild type and the Act-deficient strain M511 (a strain with an in-frame deletion of the pathway-specific regulator of Act biosynthesis, *actII*–ORF4) [32]. Four of the six new candidate SoxR targets (*SCO0319, SCO0320, SCO0321,* and *SCO1177*) were significantly under-expressed (\geq 11-fold difference) in M511 compared with wild type, while *SCO1734* and *SCO4021* showed a more modest (2–3 fold) difference in expression levels between the two backgrounds (**Table 3**).

To further confirm that reduced expression of the six new potential SoxR-targets in the $\Delta soxR$ background was due to SoxR-deficiency, we conducted a complementation experiment. Using qRT-PCR, the expression levels of the relevant genes (along with



Figure 2. RNA-Seq reads visualized on Integrative Genomics Viewer. Reads were obtained from RNA isolated from 3-day old wild type and *AsoxR* samples grown on R2YE medium. Red indicates the positive strand; blue indicates the negative strand; scale bar indicates the chromosomal position in the *S. coelicolor* M145 genome. A) *SCO0319-SCO0321;* B) *SCO1177-SCO1179;* C) *SCO1908-SCO1910;* D) *SCO2477-SCO2479;* E) *SCO4265-SCO4267;* F) *SCO7007-SCO7009.* doi:10.1371/journal.pone.0106181.g002

known controls) were analyzed in the complemented $\Delta soxR$ strain that expresses soxR from a chromosomally integrated plasmid pSET152 (Table 1). Figure 1 shows that expression of SCO1697 (soxR), and its five confirmed targets (ecaA-ecaD, SCO2478) was restored in the soxR null strain complemented with a wild type copy of soxR (soxR/pSoxR). Furthermore, expression of three genes that showed significant Act-dependence (**Table 3**), SC00320, SC00321 and SC01177, was also rescued in the soxR-complemented strain (Figure 1). By contrast, SCO0319, SCO1734 and SCO4021 failed the complementation test. In this experiment, SC00319 showed similar expression levels in the $\Delta soxR$ and the complemented $\Delta soxR$ backgrounds, while SCO1734 and SCO4021 were similarly expressed in wild type, $\Delta soxR$ and the complemented $\Delta soxR$ backgrounds (Figure 1). Given these results SC00319, SC01734 and SC04021 were eliminated as SoxR-targets.

In summary, RNA-Seq analysis followed by qRT-PCR validation identified eight genes that are both SoxR- and Actdependent. Five of these are previously confirmed members of the *S. coelicolor* SoxR regulon [5,6]. Of the newly identified SoxRinduced genes, it appeared that *SCO0320* (homologous to the carboxy-terminal half of *SLI_0274*, a quinone oxidoreductase in *Streptomyces lividans*, **Figure S2**) and *SCO0321* (carboxylesterase) may be transcriptionally coupled. As redox-associated enzymes, *SCO0320* and *SCO0321* functionally cluster with the known SoxR-targets, *ecaB*, *ecaC*, *ecaD* and SCO2478. SCO1177 encodes a putative GntR-family transcriptional regulator. The GntR family members (so named for the *Bacillus subtilis* repressor of the gluconate operon) normally act as transcriptional repressors, and regulate gene expression in response to nutritional and/or other environmental signals [33].

SoxR binds to the promoter of only one of its new putative targets (with a twist)

As mentioned before, SoxR homologs from different bacteria have highly conserved DNA binding domains, and thus bind to similar operator sequences (soxbox) in the promoters of their target genes. We previously demonstrated that SoxR directly binds to the promoters of its five known targets in S. coelicolor (all of which share a similar promoter architecture) to directly activate their transcription [5]. However, visual inspection of the DNA region upstream of the newly identified potential SoxR targets, SC00320, SC00321, and SC01177 failed to reveal potential SoxR docking sites. Nevertheless, we decided to empirically determine if SoxR binds to the promoter regions of its three new putative target genes by electrophoretic mobility shift assays (EMSA). Towards this end we incubated purified SoxR with DIGend-labeled DNA fragments that span $\sim 170-200$ bp upstream of each predicted ORF. While these assays showed that SoxR bound at the expected dissociation constant of 5 nM to the promoter of



Figure 3. Reannotation of the *SCO0319-SCO0321* **region in** *S. coelicolor.* (A) *SCO0319* is annotated in StrepDB as transcribed divergently from *SCO0320* and *SCO0321*. (B) Reannotation of the same region using Geneious (v.7.1.4 Biomatters) predicts that *SCO0320* initiates further upstream within the *SCO0319* ORF; the reannotated ORF is renamed *SCO0320ext*. A conserved SoxR-binding site (soxbox) is positioned upstream of *SCO0320ext* is aligned with the *E. coli soxS* promoter and the promoters of five confirmed SoxR target genes in *S. coelicolor*. SoxR binding sites are indicated in bold type, and the inverted arrows depict the sequence of dyad symmetry. The asterisks indicate conserved nucleotides within the SoxR binding site. The transcriptional start site of *E. coli soxS* is labeled +1, and the -10 and -35 sequences are indicated. The number of nucleotides to the predicted start codons of the different genes is shown. doi:10.1371/journal.pone.0106181.a003

ecaD (a known target), we were unable to detect binding to any of the DNA probes for the new potential targets, even at the highest concentration tested (20 nM; data not shown).

Given these puzzling findings, we examined the RNA-Seq reads for all SoxR-regulon members more closely on Integrative Genomics Viewer (**Figure 2**) [34,35]. The five known SoxR targets (*ecaA-ecaD*, *SCO2478*) appeared as expected, displaying strong expression on the annotated DNA strand in the wild type background, and very low (to negligible) reads in the $\Delta soxR$



Figure 4. SoxR binds to the promoter of *SCO0320ext in vitro*. A DIG-end-labeled DNA fragment spanning the putative promoter region of *SCO0320ext* was incubated with increasing amounts of purified histidine-tagged SoxR protein. Protein-bound complexes (C) and free DNA (F) were separated on a 5% native polyacrylamide gel. The specificity of SoxR binding was demonstrated by the addition of a 500-fold molar excess of unlabeled competitor probe. doi:10.1371/journal.pone.0106181.g004

background (Figure 2B-F). An examination of the RNA-Seq reads for the SCO0319-SCO0321 region, however, revealed a few unexpected features. According to the annotation in StrepDB (http://strepdb.Streptomyces.org.uk), SCO0320 and SCO0321 are transcribed from the same strand, while SCO0319 is divergently transcribed from the complementary strand (Figure 2A and 3A). However, the RNA-Seq reads for all three genes mapped on the same strand, and furthermore, large numbers of reads mapped to the intergenic regions between the three genes (Figure 2A). To investigate the possibility that these reads derived from a previously unannotated transcript, we reannotated and manually curated all possible ORFs within this region. An alternate ORF that spans the region of what is currently annotated as SCO0319 and SCO0320 was identified, which we have named SCO0320ext (corresponding to chromosomal positions 319614-320569; Figure 3B). A BLAST search [29] of the translated amino acid sequence from SCO0320ext revealed that its Nterminal region is homologous to SLI_0274, a quinone oxidoreductase in S. lividans, the C-terminus of which had previously been identified as homologous to SCO0320 (Figure S2). Intriguingly, the DNA region upstream of SCO0320ext revealed a potential SoxR binding site and a promoter architecture that resembles the promoters of the five confirmed SoxR targets in S. coelicolor (Figure 3C).

To test if SoxR binds to the putative promoter of *SCO0320ext*, we conducted EMSA with purified SoxR and a DNA fragment that spans the region upstream of *SCO0320ext*. **Figure 4** shows that SoxR binds to this region with high affinity; the amount of



Figure 5. Transcription of *SCO0320ext* **results in read-through into the** 5' **region of** *SCO0321***.** (A) RNA-seq reads of the *SCO0319-SCO0321* region obtained from RNA obtained from three-day old M145 wild type cells. (B) Schematic representation of reannotated *SCO0319-SCO0321* region. Predicted ORFs are indicated by block arrows; the box labeled "SB" indicates the SoxR-binding site upstream of *SCO0320ext*. The primers used for RT-PCR analysis are indicated; primer sequences are reported in Table S1. (C) RT-PCR analysis was carried out on RNA isolated from 3-day old wild type cultures grown on R2YE plates. The *hrdB* gene encodes the house-keeping sigma factor, and was included as a control. *SCO0320ext* was amplified using primers 0321N-F and 0321N-R; the 3' region of *SCO0321* was amplified with primers 0321N-F and 0321N-R; the 3' region of *SCO0321* was amplified with 2005 marker (New England Biolabs).

protein needed to bind 50% of the DNA was between 5 and 10 nM (**Figure 4**), which is comparable to the affinity of SoxR for its other target genes in *S. coelicolor* [5]. Specificity of binding was demonstrated by addition of 500-fold excess of unlabeled competitive DNA that resulted in displacement of SoxR from the labeled probe (**Figure 4**). This result, combined with the qRT-PCR assays described above, strongly suggests that *SCO0320ext* is a direct SoxR target, and the sixth member of the SoxR regulon in *S. coelicolor*.

SCO0321 and SCO1177 are not SoxR targets, but artifacts of transcriptional read-through

These results helped explain how SoxR regulates transcription of SCO0320ext. However, we were left with two other potential SoxR targets (SC00321 and SC01177) that do not appear to harbor a SoxR-binding site in their putative promoters, leaving the mechanism of their regulation unclear. Given that the RNA-seq reads for SCO0321 are concentrated within the 5' end of the gene (Figures 2A and 5A), we propose that SCO0321 is not a real SoxR-target, but likely an artifact of transcriptional read-through from SCO0320ext. The primers that were used in the previously described qRT-PCR validation assays for SCO0321 (0321N-F and 0321N-R; Figure 5B) bind within the 5' region of the gene that demonstrated high RNA-Seq reads (Figure 5A). However, when we used a different set of primers (0321C-F and 0321C-R; **Figure 5B**) that bind within the 3' region of the gene, we did not observe an RT product (Figure 5C). This result is consistent with the lack of RNA-seq reads within the 3' region of SCO0321 and indicates that the complete sequence of SCO0321 is not independently transcribed under our experimental conditions.

An examination of the RNA-Seq reads for SCO1177 similarly suggested that this gene is not a SoxR-target but an artifact of transcriptional read-through from the adjacent gene, SCO1178 (ecaC), a confirmed SoxR regulated gene. As annotated in StrepDB, SCO1177 and ecaC are convergently transcribed from

opposite strands, however all RNA-Seq reads for *SCO1177* map to the sense strand for *ecaC* (**Figure 2B**). Moreover, the reads for *SCO1177* cluster towards the 3' end of the gene that is closest to *ecaC*, and could readily be explained by transcriptional readthrough of *ecaC*. Here, *SCO1177* was detected as a SoxR-target in our initial RNA-Seq analysis because read counts were compiled without regard to strand specificity. Furthermore, the primers used in the qRT-PCR validation assays for *SCO1177* happen to prime within the 3' region of this gene that demonstrated high RNA-Seq reads (**Figure 2B**). It should be noted that we did not observe significant reads in the region corresponding to the sense strand of *SCO1177* in either wild type or $\Delta soxR$ strains at any of the time points sampled (day 3 reads shown in **Figure 2B**; day 1 data not shown). Thus, *SCO1177* also does not appear to be expressed under the growth conditions used in this work.

Discussion

In this work we expanded the SoxR regulon in *S. coelicolor* by comparing the transcriptomes of wild type and *soxR* null mutant strains in stationary phase using RNA-Seq. This regulon is composed of six genes induced by SoxR in response to the redoxactive antibiotic Act that is produced in stationary phase. Five of these genes (*ecaA-ecaD*, *SCO2478*) were previously identified as SoxR-dependent using a bioinformatics approach designed to identify genes based on the presence of a SoxR-binding site in putative promoter regions [5,6]. The Act-dependent expression of four of the SoxR-targets (ecaA-ecaD) was independently described by Huang and co-workers [20]. The sixth S. coelicolor SoxR-regulon member identified in this work, *SCO0320ext*, was missed by the bioinformatic search for SoxR-targets, and was also not previously identified as an "*eca*" gene.

SCO0320ext encodes a putative quinone oxidoreductase with homology to SLI_0274 in S. lividans and SAV_4018 in Streptomyces avermitilis (**Table 4**). Both the S. lividans and S. avermitilis

Predicted function		S. coelic	<i>olor</i> homolog in ^b				
	sco	SAV	SCLAV	SGR	SLI	SCAB	SVEN
Quinone oxidoreductase	0320ext	4018	None	None	0274	None	None
Monooxygenase	1909 (ecaB)	None	1111	5610	2221	70221	1541
NAD-dependent epimerase/dehydratase	1178 (ecaC)	None	p0595	3374	1455	5581	4172
Flavoprotein reductase	2478	5665	1679	5059	2813	62351	2266
Oxidoreductase	4266 (ecaD)	3956	3239	4037	4501	None	4021
ABC transporter	7008 (ecaA)	7218	0059	6589	7210	12171	None

genes whose promoters contain conserved SoxR binding sites: SAV_1623 (putative transketolase, and SAV_4017 (putative TetR family transcriptional regulator). The S. avermitilis genome harbors two other doi:10.1371/journal.pone.0106181.t004



Figure 6. Extent of conservation of the SoxR regulon in 55 *Streptomyces* **species.** The graph shows the number of *Streptomyces* species surveyed that contain sequences homologous to *soxR* or SoxR-regulated genes from *S. coelicolor*. The black segment indicates the number of homologs with a soxbox motif within 1 kb upstream, while the white segment shows those that lack a soxbox sequence. doi:10.1371/journal.pone.0106181.g006

homologs of SCO0320ext contain putative SoxR-binding sites in their promoters suggesting that they may be under SoxR regulation in these organisms as well. It is peculiar that only the amino-terminal half of SCO0320ext is homologous to the other two proteins. As illustrated in Figure S2, there is almost complete sequence identity between the amino-terminal halves of SCO0320ext and SLI 0274, but clear divergence in the carboxy-terminal halves. An alignment of the nucleotide sequences of SLI_0274 and SCO0320ext shows a single base difference (deletion of cytosine 494) in SCO0320ext that causes a shift in the open reading frame (Figure S2A). Interestingly, SCO0320 (as annotated in StrepDB) is almost identical to the carboxy-terminal half of SLI 0274 (Figure S2). Thus it appears that S. coelicolor M145 acquired a mutation making what was originally one gene (that corresponding to SCO0320ext), appear to be two separate genes (SCO0319 and SCO0320). Analysis of the nucleotide sequence of this region in two other S. coelicolor A3(2) derivatives (M600 and J1501) showed the same base deletion found in M145 (data not shown). At this point it is unclear if SCO0320ext is functional in S. coelicolor, or if the mutation has any physiological consequences for the organism.

SoxR is typically considered a transcriptional activator. Nevertheless, we analyzed our RNA-Seq data for genes whose expression may be inhibited by SoxR. Several candidate genes were found to be overexpressed in the $\Delta soxR$ mutant compared to wild type (\geq 4-fold difference; **Table S2**). From this list, thirteen genes showed statistically significant differential expression (qvalue \leq 0.1), and were selected for validation by qRT-PCR performed on RNA isolated from independent biological samples. Of these, only six showed significantly different expression levels $(\geq 4$ -fold) in wild type and the $\Delta soxR$ mutant by qRT-PCR (**Table S3**). However, none of these six genes passed the complementation test, where we expected lower expression in the wild type and soxR-complemented backgrounds compared to the soxR-deficient background. Instead, most were overexpressed in the soxRcomplemented strain contrary to the predicted pattern (Figure S1). It can thus be concluded that SoxR does not function as a transcriptional inhibitor in S. coelicolor.

As described earlier, SoxR stimulates the production of a similar group of genes (transporters and redox enzymes) in *S. coelicolor* and *P. aeruginosa* in response to endogenously produced antibiotics; Act in the former and phenazines in the latter. This

scenario appears to be maintained in other sequenced streptomycetes. An analysis of seven Streptomyces species that are annotated in the Streptomyces database (http://strepdb.Streptomyces.org.uk) shows that all six SoxR-regulated genes in S. coelicolor have homologs in its closest relative S. lividans, while a subset of these genes are conserved in the five other Streptomyces species (Table 4). Notably, all the genes listed in Table 4 contain a SoxR-binding sequence (soxbox) in their promoters suggesting that they are under SoxR control (soxbox sequences available in Table S4). A more extensive survey of 55 sequenced Streptomyces species showed the presence of soxR in every species analyzed (**Table S4**; **Figure 6**). Interestingly, while soxR is not autoregulated in S. coelicolor, about 13 percent of soxR homologs in other Streptomyces surveyed showed a potential soxbox upstream suggesting that these homologs may be autoregulated. The suite of SoxRregulated genes in S. coelicolor is fairly well conserved in the other streptomycetes analyzed, ranging from 100 percent conservation of *ecaA* to 33 percent for *ecaC* (**Table S4; Figure** 6). When found, these genes commonly contain a potential soxbox within one kilobase pairs upstream (Table S4; Figure 6). SCO0320ext is a notable exception with only 23 percent of homologs harboring a soxbox sequence. While none of the other species aside from S. coelicolor produce Act per se, members of this genus are known to produce other biologically active secondary metabolites. These molecules could serve as signals that trigger SoxR activity in other streptomycetes. Knowledge of the individual genes regulated by SoxR will facilitate further studies to elucidate the function of this regulatory protein and its regulon in antibiotic producers.

Supporting Information

Figure S1 Complementation analysis to further assess the SoxR-dependence of genes identified by RNA-Seq. (DOCX)

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Figure S2 Sequence alignment of SCO0320ext and SCO0320 with the quinone oxidoreductase SLI_0274 from S. lividans.

(DOCX)

Table S1Primers used in this study.(DOCX)

Table S2 Genes identified by RNA-Seq as upregulated in $\Delta soxR$ compared to WT in stationary phase. (DOCX)

Table S3Validation of RNA-Seq data reported in TableS2 by quantitative RT-PCR.(DOCX)

Table S4 tblastn matches of genes of the SoxR regulon to *Streptomyces* whole genome sequences regions. Soxbox positions are listed relative to the 5' end of the blast hit. (XLSX)

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Author Contributions

Conceived and designed the experiments: MC NN JAS. Performed the experiments: MC NN JAS. Analyzed the data: MC NN JAS. Contributed reagents/materials/analysis tools: MC JAS. Contributed to the writing of the manuscript: MC NN JAS.

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