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Response to Copper Stress in *Streptomyces lividans* Extends beyond Genes under Direct Control of a Copper-sensitive Operon Repressor Protein (CsoR)*[§]

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Background: Balancing copper homeostasis and bioavailability is critical for morphological development in *Streptomyces lividans*.

Results: In *Streptomyces lividans*, a copper-sensitive operon repressor (CsoR) protein regulates a three-locus copper homeostasis regulon.

Conclusion: Genes regulating copper homeostasis are more extensive than the regulon under direct control of CsoR.

Significance: CsoR is part of a regulatory network controlling copper homeostasis that extends beyond the CsoR regulon.

A copper-sensitive operon repressor protein (CsoR) has been identified in *Streptomyces lividans* (CsoR^{Sl}) and found to regulate copper homeostasis with attomolar affinity for Cu(I). Solution studies reveal apo- and Cu^I-CsoR^{Sl} to be a tetramer assembly, and a 1.7-Å resolution crystal structure of apo-CsoR^{Sl} reveals that a significant conformational change is necessary to enable Cu(I) binding. *In silico* prediction of the CsoR regulon was confirmed *in vitro* (EMSA) and *in vivo* (RNA-seq), which highlighted that next to the *csoR* gene itself, the regulon consists of two Cu(I) efflux systems involving a CopZ-like copper metallochaperone protein and a CopA P₁-type ATPase. Although deletion of *csoR* has only minor effects on *S. lividans* development when grown under high copper concentrations, mutations of the Cu(I) ligands decrease tolerance to copper as a result of the Cu(I)-CsoR mutants failing to disengage from the DNA targets, thus inhibiting the derepression of the regulon. RNA-seq experiments carried out on samples incubated with exogenous copper and a Δ *csoR* strain showed that the set of genes responding to copper stress is much wider than anticipated and largely extends beyond genes targeted by CsoR. This suggests more control levels are operating and directing other regulons in copper homeostasis beside the CsoR regulon.

For many bacterial organisms, maintaining cellular metal ion homeostasis is an essential requirement for viability. Sophisti-

cated cellular machinery has evolved and consists of an extensive network of specialized proteins and transporters that respond to either metal ion deprivation or overload (1–3). Copper is an essential transition metal ion in biology and is required by many proteins and enzymes that function in electron-transfer chemistry and in the catalytic oxidation of substrates. The ability of copper to donate or accept single electrons is also detrimental as “free” cellular copper ions are complicit in a Fenton-like reaction, catalyzing the production of damaging hydroxyl radicals (4). This deleterious behavior leads to bacteria avoiding copper toxicity through buffering copper in the cytosol very tightly (2, 3, 5). Once the buffering capacity for cytosolic copper is exceeded, this is sensed by transcriptional regulatory proteins termed copper sensors that trigger the expression of genes encoding for efflux systems such as copper-exporting P₁-type ATPases and copper metallochaperones that act in tandem to return the cytosol to a buffered copper state (3, 6, 7).

The filamentous Gram-positive soil-dwelling bacterium *Streptomyces* is important to both ecological and human welfare and is also one of the model systems for bacterial morphological and physiological development. The morphological differentiation of *Streptomyces* has three characteristic development stages as follows: 1) formation of branched vegetative mycelium, 2) formation of aerial hyphae, and 3) the production of spores. Concomitant or just before the morphological switch from vegetative to aerial mycelium, the metabolic development is initiated by the production of secondary metabolites. These compounds comprise a large percentage of the clinically useful antibiotics, anti-fungals, and even some of the anti-tumor agents used to date. The bioavailability of copper ions has been shown to be of crucial importance for morphological development in certain streptomycetes (8, 9). Copper dependence is restricted to the reproductive growth phase (aerial mycelium and spores), whereas vegetative growth proceeds under strongly copper-limiting conditions. Evidence from genetic

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[§] This article contains “Materials and Methods,” supplemental Figs. S1–S9, and Tables S1–S6.

The atomic coordinates and structure factors (code 4adz) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (<http://www.rcsb.org/>).

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Copper Sensing in *Streptomyces lividans*

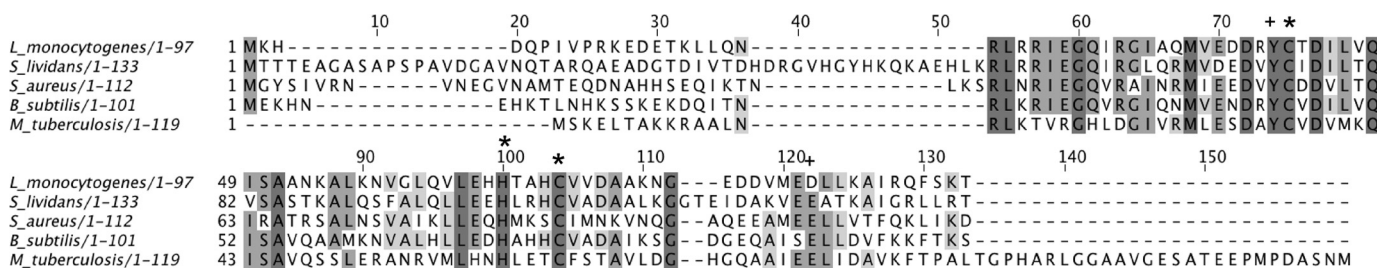


FIGURE 1. ClustalW2 multiple amino acid sequence alignment of copper-sensing CsoR proteins for which biochemical or structural data have been reported. Completely and partially conserved residues are boxed in dark and light gray, respectively. The conserved Cu(I)-binding ligands are indicated (*) along with residues reported to play a role in the Cu(I)-binding allosteric regulation of DNA (+). The UniprotKB accession numbers are Q8Y646 (*L. monocytogenes*), A6QIT1 (*S. aureus*), O32222 (*B. subtilis*), and P71543 (*M. tuberculosis*).

knock-out studies has suggested that secreted cuproproteins or cuproenzymes are needed for the development switch from vegetative to aerial mycelium to occur, but their identification and role in the development switch have not so far been elucidated (10, 11).

The copper proteome of *Streptomyces coelicolor* has been determined using bioinformatics approaches, revealing a rich assortment of putative extracellular cuproenzymes, redox proteins, and copper metallochaperone-like proteins (10, 12). An exciting yet unexpected finding was the identification of a cuproenzyme along with a dedicated copper metallochaperone in the cytosol (10, 13). A metabolic requirement for copper in the bacterial cytosol has not yet been documented; however, *in vitro* biochemical evidence indicates that this cytosolic tyrosinase-like enzyme (MelD2) is active in the oxidation of mono- and di-phenols (13). The cytosol of *Streptomyces coelicolor* also contains proteins putatively involved in copper detoxification systems (10). Two operons have been identified that contain genes encoding for a CopZ-like copper metallochaperone and a CopA-like P₁-type ATPase transporter. In certain bacteria, the *copA* and *copZ* resistance genes are transcriptionally regulated by a copper sensor protein belonging to the copper-sensitive operon regulator (CsoR)³ family (7). In *S. coelicolor*, the gene *SCO4136* has been proposed to encode for a CsoR orthologue (10). However, the genomic environment of *SCO4136* does not contain any known copper resistance genes and instead is located among genes encoding for proteins involved in phosphate transport (10). Therefore, if the gene product of *SCO4136* functions as the cytosolic copper sensor in *S. coelicolor*, it is not genetically linked to copper resistance genes such as the two putative *copZA*-like operons.

The founding member of the CsoR family was discovered in *Mycobacterium tuberculosis* (CsoR^{Mtb}) (7). Under elevated copper levels, the expression of CsoR^{Mtb} was found to be strongly induced, with the binding of Cu(I) triggering the derepression of the *rv0967–rv0970* operon in which *rv0969* encodes for a P₁-type ATPase involved in copper transport and *rv0967* for CsoR^{Mtb} (7, 14). It has subsequently been found that members of the CsoR family are widespread in bacterial genomes. Examples of CsoR proteins that transcriptionally regulate *copZA* copper resistance operons have been identified and

characterized to varying degrees in *Bacillus subtilis* (15), *Listeria monocytogenes*, and *Staphylococcus aureus* (16, 17). Furthermore, a recent report has identified a CsoR member that is not thought to be involved in copper homeostasis but instead appears to function in response to sulfur stress (17).

To understand the role copper plays in morphological differentiation and development, it is necessary to understand how cellular copper is handled under normal and elevated conditions. *S. coelicolor* and *Streptomyces lividans* both display a distinct dependence on copper for development with that of *S. lividans* being more pronounced (18). They also share a high level of sequence identity and genome organization (19, 20), and the gene numbering of the *S. coelicolor* genome database is used for *S. lividans* in this study. Sequence alignment of the *S. lividans* 4136 gene product with known CsoR orthologues reveals conservation of the amino acids that act as Cu(I)-binding ligands and two amino acids considered important for the Cu(I)-dependent allosteric regulation of DNA binding (Fig. 1). In this study, we have biochemically and structurally characterized the *S. lividans* 4136 gene product and find it to be a tetrameric Cu^I-CsoR (CsoR^{S^l}). *In vivo* studies with a genetic knockout (Δ *csoR*) and CsoR^{S^l} mutants indicate CsoR^{S^l} to be essential for the response of *S. lividans* to elevated copper levels. Furthermore, we have coupled bioinformatics and RNA-seq experiments to characterize the regulon under the direct control of CsoR^{S^l}. RNA-seq data suggest that the regulation of copper homeostasis is much more extensive than only the genes repressed by CsoR^{S^l}.

MATERIALS AND METHODS

Streptomyces Strains, Media, and Growth Conditions—The *Streptomyces* strains used are as follows: *S. lividans* 1326 (*S. lividans* 66, stock number 1326 John Innes Collection), *S. lividans* Δ *csoR* (this study), and *S. coelicolor* A3(2) strain M512 (21). The agar media soy flower mannitol, complex medium (R5), minimal agar medium (MM), the liquid complex medium tryptic soy broth with 10% sucrose, and the liquid defined medium NMMP were prepared according to Ref. 22. If required, glucose and/or mannitol were added to 0.5%. Agar plates were incubated at 30 °C, and liquid cultures were grown in 250-ml baffled flasks with 0.2- μ m vent caps (Corning Glass) with shaking at 160 rpm. Liquid cultures were inoculated with spores to a final concentration of 2×10^7 spores/ml. Spore stocks were obtained from cultures grown on soy flower mannitol plates and stored in 20% glycerol at –20 °C. Growth was

³ The abbreviations used are: CsoR, copper-sensitive operon repressor; AAS, atomic absorption spectroscopy; MM, minimal agar medium; BCS, bathocuproine disulfonate; BCA, bicinchoninic acid; AUC, analytical ultracentrifugation.

recorded by determining the dry biomass from 1.5-ml samples collected in pre-dried Eppendorf tubes, and mycelium was pelleted by centrifugation at 13,000 rpm for 10 min. After 12–16 h at 105 °C, the dry weight was determined using an analytical balance.

Generation of a CsoR Deletion Mutant (Δ csoR) of *S. lividans* 1326—The 4136 gene encoding for CsoR^{Sl} was deleted in *S. lividans* 1326 in a two-step process using the CRE-lox system (23). First, the gene (nt +4 to 399) was replaced by homologous recombination with an apramycin resistance cassette flanked by loxP sites. For this purpose, the upstream flanking region of SL4136 (–1370 to +3) and the downstream flanking region (+399 to 1680) were amplified from genomic DNA by PCR, including EcoRI, XbaI, XbaI, and HindIII sites, respectively, for cloning purposes. These two fragments and the apramycin resistance cassette flanked by loxP sites were cloned in the delivery vector pWHM3 that is unstable in *Streptomyces* (24). Following protoplast transformation, recombinants that were apramycin-resistant but had lost the vector (thiostrepton resistance) were isolated. Second, an unstable plasmid encoding the Cre recombinase was introduced (23) and allowed for the excision of the apramycin resistance cassette on the lox sites. The resulting strain, Δ csoR, has no coding sequence for CsoR^{Sl} and has only a 61-nt “scar,” including two XbaI sites left in the genome. The Δ csoR strain was analyzed by PCR to confirm the loss of 4136, the apramycin resistance cassette, and vector sequences.

RNA Isolation and Transcriptome Analysis by RNA-seq—Total RNA was isolated with Kirby mix according to standard procedures from mycelium in early log phase grown on NMMP supplemented with 0.5% glucose and mannitol (22). Cultures were induced with 400 μ M Cu(II) for 2 h followed by total RNA isolation. RNA integrity was confirmed by agarose gel electrophoresis, and the absence of genomic DNA was checked by PCR. For the removal of ribosomal RNA, an Ambion kit was used. Samples were sent to BaseClear, an independent and accredited service laboratory for DNA-based research, and transcriptome analysis by RNA-seq was carried out. The sequences obtained on an Illumina sequencer were filtered for noncoding RNAs and analyzed with CLCbio bioinformatics software packages using the annotated *S. coelicolor* genome as reference. Expression values were expressed as reads/kb of exon model/million mapped reads (25), *i.e.* dividing the total number of exon reads (in this case one exon per reference sequence) by the number of mapped reads (in Millions) times the exon length (in this case the length of the reference sequence).

Promoter Probing—The DNA sequence (–300 to +3) upstream of SL4136 was obtained from genomic DNA by PCR introducing a flanking EcoRI and BamHI site to facilitate cloning in pIJ2585 digested with the same enzymes (26). The resulting plasmid, p4136-I, was introduced in strain M512 by protoplast transformation. cultures were grown in triplicate in NMMP medium supplemented with 0.5% glucose, 0.5% mannitol, and 50 μ g/ml apramycin. Strain M512 transformed with the empty vector was grown under the same conditions to obtain background readings. Following extraction of mycelium with methanol, the concentration of undecylprodigiosin was

quantified from the absorbance at 530 nm and the extinction coefficient of 100,500 liters/mol^{–1} cm^{–1} (27).

Determination of Tyrosinase and Cytochrome *c* Oxidase Activity—The wild type, *S. lividans* 1326, and the mutant strain Δ csoR were each transformed with pIJ703 (plasmid harboring the *Streptomyces antibioticus melC* operon under control of its own promoter). Four independent transformants of each strain were grown in liquid tryptic soy broth with 10% sucrose medium supplemented with 25 μ M Cu(II). Tyrosinase activity was determined in spent medium with 10 mM 3,4-dihydroxy-L-phenylalanine in 100 mM phosphate buffer, pH 6.8, as substrate (28). Cytochrome *c* oxidase activity was visualized with *N,N,N',N'*-tetramethyl-*p*-phenylenediamine as substrate, essentially according to Refs. 29, 30.

Cloning, Overexpression, and Purification of CsoR^{Sl} from *Escherichia coli*—The 4136 gene was cloned from *S. lividans* 1326 genomic DNA (supplemental “Materials and Methods”) and overexpressed in *Escherichia coli* using a pET28a (Kan^r) vector (Novagen). This construct, designated pET4136, was transformed to *E. coli* BL21(DE3) cells, and single colonies were transferred to 2 \times YT medium (Melford) with kanamycin (50 μ g/liter) (Melford) at 37 °C. Overexpression of the N-terminal His-tagged CsoR^{Sl} was induced by 1 M isopropyl β -D-1-thiogalactopyranoside (Melford) to a final concentration of 1 mM, and the temperature was decreased to 25 °C for overnight incubation. Cultures were harvested by centrifugation at 4000 rpm for 20 min at 4 °C, and the cell pellet was resuspended in 50 mM Tris/HCl, 500 mM NaCl (Fisher) and 20 mM imidazole (Sigma) at pH 7.5 (Buffer A). The resuspended cell suspension was lysed using an EmulsiFlex-C5 cell disrupter (Avestin) followed by centrifugation at 18,000 rpm for 20 min at 4 °C. The clarified supernatant was loaded onto a 5-ml nickel-nitrilotriacetic acid-Sepharose column (GE Healthcare) equilibrated with Buffer A and eluted by a linear imidazole gradient using Buffer B (Buffer A with 500 mM imidazole). A single peak at ~30% Buffer B was eluted from the column, and fractions were pooled and dialyzed overnight at 4 °C against 10 mM MES, pH 6.5, 150 mM NaCl, 2 mM dithiothreitol (DTT) (Melford), and 4 mM EDTA (Sigma) (Buffer C). Following dialysis, the N-terminal His tag was removed by incubating the protein at room temperature overnight with 125 units of thrombin (Sigma). The protein/thrombin mixture was reappplied to the nickel-nitrilotriacetic acid-Sepharose column (GE Healthcare), and the flow-through was collected and concentrated using a Centricon (VivaSpin) with a 5-kDa cutoff at 4 °C for application to a G-75 Sephadex column (GE Healthcare) equilibrated with Buffer C. Fractions eluting from the major peak of the G-75 column were analyzed by SDS-PAGE, and those deemed of good purity were concentrated and stored at –20 °C until required.

Site-directed Mutagenesis of CsoR^{Sl} and Cloning for *in Vivo* Studies—The QuikChange site-directed mutagenesis method (Stratagene) was used to create the C75A and H100A mutants of CsoR^{Sl}. Forward and reverse primers were designed with the respective nucleotide change(s) to create the desired mutation (supplemental “Materials and Methods”), and the pET4136 plasmid was used as template. The respective mutations were confirmed by DNA sequencing. For expression in *S. lividans*, the wild type and mutant CsoR^{Sl} open reading frames (ORF)

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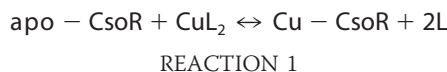
were cloned under control of the 4136 promoter and the constitutive *ermE* promoter. For this purpose, the 4136 promoter region was obtained by PCR from *S. lividans* 1326 genomic DNA introducing an NdeI site on the ATG start codon of the ORF, and the *ermE* promoter was obtained from plasmid pHM10a (31). The wild type, C75A, and the H100A ORFs were cloned downstream of the promoters in the low copy vector pHJL401 (32).

UV-visible and Circular Dichroism Spectroscopies—A Varian Cary 50 UV-visible spectrophotometer and an Applied Photophysics Chirascan circular dichroism (CD) spectrophotometer (Leatherhead, UK) both equipped with a thermostatic cell holder controlled with a Peltier system were routinely used. An extinction coefficient (ϵ) at 280 nm of $3105 \text{ M}^{-1} \text{ cm}^{-1}$ was calculated for the CsoR^{SI} monomer. This value was used throughout to determine the concentration of apo-CsoR^{SI} samples. Far-UV CD spectra at 20 °C with 20 μM CsoR^{SI} in 10 mM potassium phosphate, 50 mM potassium fluoride, pH 7.0, were acquired in the range 260 to 190 nm.

Electrospray Ionization Mass Spectrometry (ESI-MS) and Atomic Absorption Spectroscopy (AAS)—CsoR^{SI} samples for mass spectrometry were diluted 1:20 with a 50% methanol and 1% formic acid solution. Spectra were acquired on a Micromass Quattro Ultima triple quadrupole instrument using the following experimental parameters: capillary voltage 1.7 kV, cone voltage 80–120 V, and cone gas 100 liter/h. Acquisition and processing were carried out using MassLynx software (Waters, Manchester, UK). Metal content was determined with a Unicam 939/959 atomic absorption graphite furnace spectrometer.

Cu(I) Titrations and Determination of Binding Affinity—Apo-CsoR^{SI} samples for experiments with Cu(I) were prepared in an anaerobic chamber (DW Scientific [O₂] <2 ppm) by first incubating for 2–3 h with 2 mM DTT followed by desalting using a PD-10 column (GE Healthcare) equilibrated with either 10 mM MOPS, pH 7.5, 150 mM NaCl, or 10 mM MES, pH 6.5, 150 mM NaCl. Free thiol content was determined by the reduction of 5,5'-dithiobis(2-nitrobenzoic acid) monitored at 412 nm ($\epsilon = 13,500 \text{ M}^{-1} \text{ cm}^{-1}$) (33). Cu(I)Cl (Sigma) was dissolved under anaerobic conditions in 10 mM HCl and 500 mM NaCl and diluted with either MES or MOPS buffer. Cu(I) concentration was determined spectrophotometrically by stepwise addition using a gastight syringe (Hamilton) to a known concentration of the Cu(I)-specific bidentate chelator bicinchoninic acid (BCA) using an extinction coefficient at 562 nm of $\epsilon = 7900 \text{ M}^{-1} \text{ cm}^{-1}$ for [Cu^I(BCA)₂]³⁻ (34). Apo-CsoR^{SI} proteins (30–50 μM) were sealed in an anaerobic quartz cuvette (Hellma), and the absorbance change at 240 nm was monitored upon titrating in the Cu(I) solution. Competition assays were set up anaerobically with either BCA or bathocuproine disulfonate (BCS) (Sigma). Increasing protein concentrations (0–90 μM) were added to solutions of [Cu^IL₂]³⁻ of defined molar ratio L:Cu(I) ≥ 3 (to ensure the presence of the 1:2 complex [Cu^IL₂]³⁻ with negligible contribution from the 1:1 complex [Cu^IL₂]⁻) creating a series of individual solutions with constant [Cu^I] and [L] and varying [apo-CsoR^{SI}]. Samples were left for between 1 and 4 h, and the transfer of Cu(I) from the [Cu^IL₂]³⁻ complex to apo-CsoR^{SI} was determined by measuring the

absorbance of the [Cu^IL₂]³⁻ complex spectrophotometrically for L = BCA at 562 nm ($\epsilon = 7900 \text{ M}^{-1} \text{ cm}^{-1}$) and L = BCS at 483 nm ($13,000 \text{ M}^{-1} \text{ cm}^{-1}$) (34, 35). By interchanging L, assays favoring competitive or noncompetitive Cu(I) binding could be set up, which for the latter led to an estimate of the binding stoichiometry. The dissociation constant for Cu(I) ($K_D(\text{Cu}^{\text{I}})$) was determined from competitive assays by assuming Reaction 1 (1),



and by using Equation 1,

$$K_D\beta_2 = \frac{([\text{apo-CsoR}]_{\text{tot}}/[\text{M-CsoR}]) - 1}{\{([\text{L}]_2/[\text{ML}_2]) - 2\}[\text{ML}_2]} \quad (\text{Eq. 1})$$

where [L] is the total ligand concentration (BCA or BCS), and the overall formation constant (β_2) is $10^{17.2} \text{ M}^{-2}$ for [copper (BCA)₂]³⁻ and $10^{19.8} \text{ M}^{-2}$ for [copper (BCS)₂]³⁻ (34, 36). Assays were performed in duplicate, and the $K_D(\text{Cu}^{\text{I}})$ value for a series was initially calculated for each individual solution and then averaged. By using the average $K_D(\text{Cu}^{\text{I}})$ value, a simulated curve was plotted using Equation 1.

Bioinformatic Identification of CsoR^{SI} Operator Targets—The computational prediction of CsoR^{SI} *cis*-acting elements was performed as described previously (37). Experimentally validated binding sites of CsoR orthologues in *B. subtilis* (TAATACCCTACGGGGGTATGG) (15, 38), *S. aureus* (ATATACCTATAGGGGGTACAT) (17), *Geobacillus thermodenitrificans* (TTATACCCGAAGGGGGTATAT) (17), and *M. tuberculosis* (RicR.1, ATATACCACCCGGGGGTATAG; RicR.2, ATATACCCTATAGGGGGTAGG; RicR.3, ATATACCCTATACGGGGTATCT; RicR.4, TTGTACCCAGCGGGGTATCG) (39) were used to generate via the PREDetector program (40) a first weight matrix to identify similar putative *cis*-acting elements within *S. coelicolor* (SCO), *Streptomyces griseus* (SGR), *Streptomyces scabies* (SCAB), and *Streptomyces avermitilis* (SAV) genomes. This first round of prediction allowed the identification of highly reliable putative *cis*-acting sequences in terms of (i) scores, (ii) interspecies conservation, and (iii) physiological meaning (hits related to copper sensitivity). Hits conserved among the selected four streptomycetes and identified upstream of copper utilization-related genes, *i.e.* upstream of *copZ* (copper chaperone) and upstream of the CsoR^{STREP} orthologues, were used to generate a new weight matrix (named “CsoR streptomycetes”), more specific for predictions in *Streptomyces* species. Sequences used to generate the CsoR streptomycetes weight matrix were *copZ*^{SCO}, *copZ*^{SGR}, *copZ*^{SCAB}, and *copZ*^{SAV}, and *csoR*^{SCO}, *csoR*^{SGR}, *csoR*^{SCAB}, and *csoR*^{SAV} and are presented in supplemental Fig. S6, and reliable CsoR-binding sites identified in these four *Streptomyces* species are presented in supplemental Tables S2–S5. Putative CsoR-like sequences predicted in SCO were used to identify *cis*-acting elements in the closely related strain *S. lividans*.

Electrophoretic Mobility Shift Assays of CsoR^{SI} Targets—Intergenic DNA fragments (208, 232, and 240 bp) containing the

predicted target sequence for CsoR^{Sl} identified by PREDetector were amplified from genomic *S. lividans* 1326 DNA as described in the supplemental “Materials and Methods”. DNA oligomers (Sigma) for use in EMSA studies were between 35 and 36 bp in length. Complementary pairs were annealed by heating at 96 °C in a water bath for 5 min and left to cool to room temperature overnight. 0.5 μM of a DNA oligomer target was incubated with concentrations of apo-CsoR^{Sl} monomer ranging between 4 and 30 μM in 10 mM HEPES, pH 7.5, 150 mM NaCl, 1 mM DTT. Cu(I)-CsoR^{Sl} samples were either prepared by pre-loading apo-CsoR^{Sl} with a stoichiometric amount of Cu(I) in an anaerobic chamber before mixing with DNA samples or added directly to the DNA-protein complex under anaerobic conditions. All samples were incubated at room temperature for 30 min and then loaded (20 μl) to a pre-run 6% Tris/borate EDTA (TBE) polyacrylamide gel. Gels were stained for 30 min in an ethidium bromide solution followed by imaging.

Crystallization and Structure Determination of Apo-CsoR^{Sl}—Crystals of apo-CsoR^{Sl} were grown using the hanging drop vapor diffusion method at 20 °C. 1 μl of protein solution at a concentration of 15 mg/ml was mixed with an equal volume of reservoir solution containing 1.26 M ammonium sulfate, 0.1 M sodium citrate, pH 4. Crystals of dimensions ~0.2 × 0.2 × 0.2 mm grew within 1 week. A single crystal was transferred to a cryoprotectant solution containing 1.3 M ammonium sulfate, 0.1 M sodium citrate, pH 4, and 15% glycerol, prior to flash-cooling to 100 K by plunging into liquid nitrogen. Crystallographic data were measured to 1.7 Å resolution at Diamond Light Source beamline I04 using an ADSC Q315r CCD detector and an x-ray wavelength of 0.9795 Å. Data were indexed using iMosflm (41) and scaled and merged using Scala (42) in the CCP4i suite. The structure was solved by molecular replacement in BALBES (43). The initial model was built into the electron density map using Buccaneer (44) and refined using Refmac5 (45). Riding hydrogen atoms were added when refinement of the protein atoms had converged. Models were rebuilt between refinement cycles in Coot (46), and the final structure was validated using the MolProbity server (47) and Coot. Coordinates and structure factors were deposited in the RCSB Protein Data Bank with accession number 4adz. A summary of data and refinement statistics and the quality indicators for the structure are given in Table 1.

RESULTS

Mass, Metal Content, Secondary Structure, and Assembly State of Purified CsoR^{Sl}—Purified CsoR^{Sl} ran as a single band on an SDS-polyacrylamide gel (supplemental Fig. S1) and denaturing ESI-MS gave a single component spectrum with a species of molecular mass 14,739 Da, in excellent agreement with the predicted mass (supplemental Table S1). AAS revealed negligible copper or nickel ion content, and thus an apo-form of CsoR^{Sl} was purified. The isolated protein reduced 5,5'-dithiobis(2-nitrobenzoic acid) to give a protein/thiol ratio of 1:1.8, suggesting that the thiol groups of Cys-75 and Cys-104, putatively involved in Cu(I) binding, remain reduced under the purification conditions (see below). The CD spectrum of apo-CsoR^{Sl} is shown in Fig. 2A with the two negative minima at 222 and 208 nm being

TABLE 1
Crystallographic data collection and processing statistics for apo-CsoR^{Sl}

Values in parentheses refer to the outermost resolution shell (1.79 to 1.70 Å).

Wavelength	0.9795 Å
Resolution	45.9 to 1.70 Å
Space group	P2 ₁ 2 ₁
Unit cell	<i>a</i> = 41.63 Å, <i>b</i> = 54.55 Å, <i>c</i> = 91.75 Å
Unique reflections	22,220 (2403)
Completeness	93.8% (71.7%)
<i>R</i> _{merge}	0.058% (0.283%)
<i>I</i> / <i>σ</i> (<i>I</i>)	12.1 (2.1)
<i>R</i> _{cryst}	0.185
<i>R</i> _{free}	0.229
ESU based on ML	0.071 Å
Root mean square deviation bond lengths	0.017 Å
Root mean square deviation bond angles	1.66°
Ramachandran favored	100%
Wilson <i>B</i> -factor	22.4 Å ²
Protein Data Bank accession code	4adz

typical signatures for α-helical secondary structure. Using this CD spectrum, a secondary structure content of 51% α-helix, 8% β-strand, 13% loops, and 27% unordered was predicted using Dichroweb (48, 49). Size-exclusion chromatography using a calibrated G-75 column revealed a major peak eluting at a mass of ~74 kDa, suggesting that under native conditions apo-CsoR^{Sl} exists as a higher order assembly (Fig. 2B). This was confirmed from analytical ultracentrifugation (AUC) experiments (supplemental “Materials and Methods”) where sedimentation equilibrium scans were fitted to a single component model to give an average single species molecular mass of 59.3 kDa (supplemental Fig. S2A). Based on the mass determined using ESI-MS, the mass obtained from AUC is consistent with apo-CsoR^{Sl} existing as a tetrameric assembly (expected mass 58,955.6 Da).

Apo-CsoR^{Sl} Binds Cu(I) with Attomolar Affinity—An anaerobic titration of Cu(I) into apo-CsoR^{Sl} gave rise to absorbance changes in the UV region of the spectrum (Fig. 2C). The absorbance increase at 240 nm is indicative of the formation of Cys-thiolate copper coordination bonds, whereas the 320 nm increase has not previously been reported for other CsoR orthologues (7, 38). The absorbance change at 240 nm saturates at ~1 mol eq of Cu(I) per monomer of CsoR^{Sl} (Fig. 2C, inset). Beyond this point, a shallow increase in absorbance occurs upon further addition of Cu(I). The end point sample from the titration was passed through a desalting column to remove excess copper, and AAS gave a 1:1 copper :CsoR^{Sl} monomer stoichiometry. Cu(I)-CsoR^{Sl} samples were characterized by CD spectroscopy, gel filtration (Fig. 2, A and B), and AUC (supplemental Fig. S2B). The secondary structure content remains essentially unchanged compared with apo-CsoR^{Sl} and gel filtration (Fig. 2B), and AUC results are consistent with Cu(I)-CsoR^{Sl} maintaining a tetramer assembly.

Accurate determination of the *K*_D(Cu^I) for CsoR^{Sl} necessitates the use of competition assays employing the specific Cu(I) bidentate chelators BCA and BCS (36, 50). At pH 7.5, addition of apo-CsoR^{Sl} into [Cu^I(BCA)₂]³⁻ leads to the extraction of 1 eq of Cu(I) from the [Cu^I(BCA)₂]³⁻ complex (supplemental Fig. S3). This indicates that under the concentrations employed, BCA cannot compete with CsoR^{Sl} for Cu(I), and that 1 eq of copper is bound per CsoR^{Sl} monomer, corroborating the

Copper Sensing in *Streptomyces lividans*

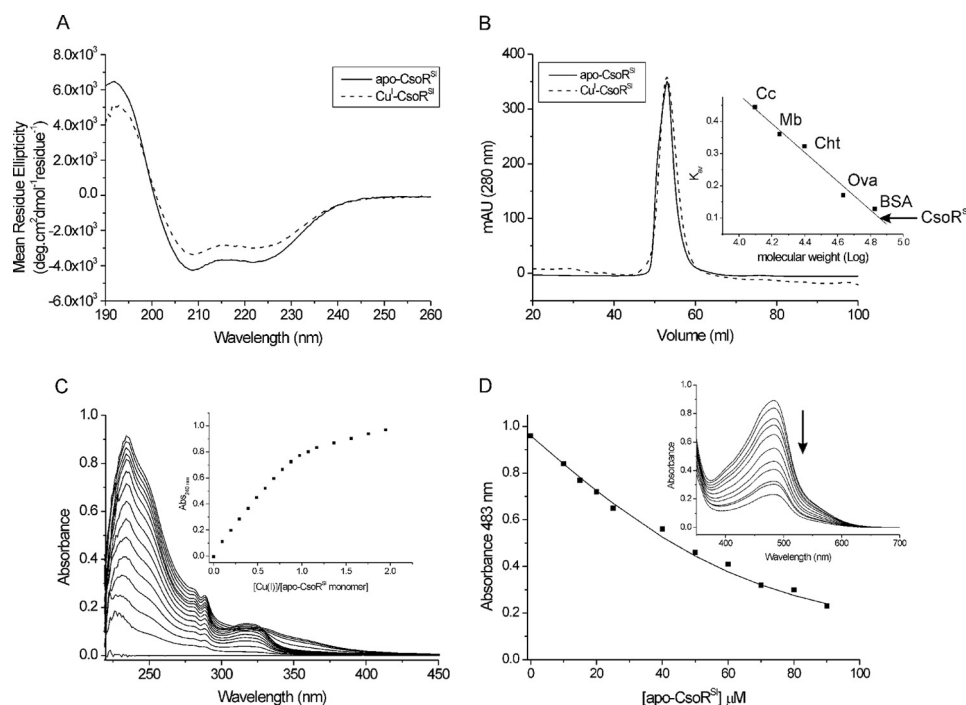


FIGURE 2. **CsoR^{SI} exists in solution as a tetramer and binds Cu(II).** *A*, far-UV CD spectra of CsoR^{SI} at pH 7 and 20 °C. *B*, G-75 size-exclusion chromatography profile of CsoR^{SI}. The gel phase distribution coefficients (K_{av}) versus log molecular weight of the following standards run individually on the column; cytochrome *c* (*cc*; 12.5 kDa), myoglobin (*Mb*; 17 kDa), chymotrypsin (*Cht*; 25 kDa), ovalbumin (*ova*; 43 kDa), and bovine serum albumin (*BSA*; 67 kDa) are plotted (*inset*) to estimate the molecular weight of the eluted CsoR^{SI}. *C*, changes in the UV region of the apo-CsoR^{SI} base-lined spectrum upon addition of Cu(II)Cl at pH 7.5, 20 °C. The *inset* shows the increase in absorbance at 240 nm plotted as a function of [Cu(II)]/[monomer-CsoR^{SI}]. *D*, determination of the K_D (Cu^I) for apo-CsoR^{SI} at pH 7.5 under copper-limiting conditions imposed by [Cu^I(BCS)₂]³⁻ as a competitive probe. The absorbance at 483 nm in the visible spectrum of [Cu^I(BCS)₂]³⁻ decreases upon addition of apo-CsoR^{SI} (*inset*) and can be used to determine the K_D (Cu^I) using Equation 1. The *line* represents a best fit to the data using an average K_D (Cu^I) 2.6×10^{-18} M. Conditions used are as follows: 10–90 μ M [apo-CsoR^{SI}], 74 μ M [Cu^I]_{total}, and 300 μ M [BCS].

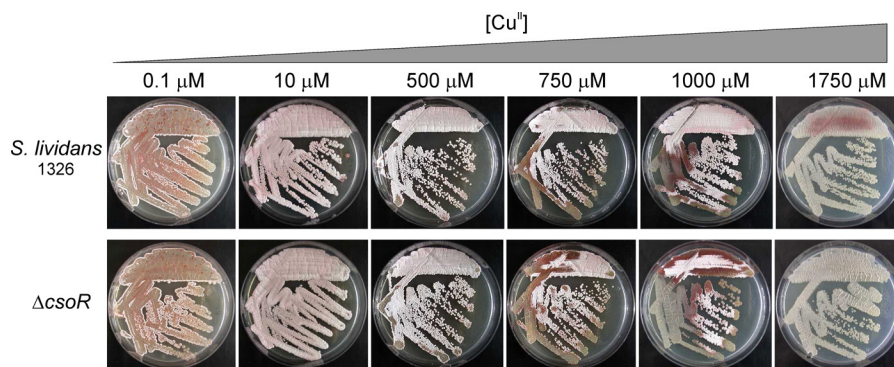


FIGURE 3. **Effect of copper on morphological development of *S. lividans* 1326.** Spores of wild type *S. lividans* 1326 and the Δ *csoR* mutant were streaked on R5 agar plates in the presence of increasing concentrations of exogenous Cu(II) and incubated at 30 °C. Photographs were taken after 90 h.

results from the UV titration and AAS experiments. Competition for Cu(I) between apo-CsoR^{SI} and [Cu^I(BCS)₂]³⁻ was observed (Fig. 2D), and analysis of the data set shown in Fig. 2D using Equation 1 and a β_2 of $10^{19.8} \text{ M}^{-2}$ for [Cu^I(BCS)₂]³⁻ gave a K_D (Cu^I) of 2.6×10^{-18} M. This K_D (Cu^I) was used to simulate a fit of the data, as shown by the *solid line* in Fig. 2D. Duplicate data sets were obtained with varying [Cu^I]_{total} or [BCS], and the average K_D (Cu^I) for CsoR^{SI} at pH 7.5 was 6.7×10^{-18} M. Competition between CsoR^{SI} and [Cu^I(BCS)₂]³⁻ was also observed at pH 6.5, and an average K_D (Cu^I) of 3.2×10^{-17} M was determined.

Growth and Development of *S. lividans* 1326 Versus That of Δ *csoR* Strain—The attomolar affinity of CsoR^{SI} for Cu(I) was consistent with a role in sensing, buffering, and handling cop-

per in the cytosol under all growth conditions, including stress elicited by elevated copper concentrations. To assess the effect CsoR^{SI} has on growth and morphological development of *S. lividans* 1326 under normal conditions and when copper stress is applied, a deletion strain (Δ *csoR*) was constructed. Fig. 3 shows an example of growth on R5 agar media of the wild type *S. lividans* 1326 and the Δ *csoR* strain at increasing copper concentrations. At low concentrations of copper, no significant differences between the two strains were observed regarding vegetative, aerial growth, and sporulation (Fig. 3). This was also the case on various other solid media and in the presence of copper chelators (supplemental Fig. S4A). However, above 750 μ M Cu(II), the Δ *csoR* strain appears to be slightly more affected in morphological development and in the production of the red

TABLE 2

Growth rates (doubling/h) of the wild type *S. lividans* 1326 and the $\Delta csoR$ mutant in NMMP medium supplemented with the indicated [Cu(II)]

Standard deviation of the growth rates was 0.01.

Strain	Concentration of Cu(II)				
	0	100	400	750	1000
<i>S. lividans</i> 1326	0.27	0.28	0.28	0.27	0.28
$\Delta csoR$	0.28	0.29	0.27	0.26	0.26

pigment (undecylprodigiosin) than the wild type (Fig. 3). Copper therefore has a triphasic effect on development. Under conditions where all copper in the medium is bound by the chelator bathocuproine disulfonic acid, development is completely blocked (supplemental Fig. S4A). Stimulation of aerial hyphae production and spores occurs when the bioavailability of copper concentrations increase to 2–5 μM , and development is inhibited again at concentrations above 500 μM . This effect is not observed on MM or soy flower mannitol solid media, but on MM development is also severely retarded above 500 μM Cu(II) and inhibited completely at 1000 μM (supplemental Fig. S4B). Growth rate determinations in liquid defined media (NMMP) supplemented with Cu(II) in the range from 0 to 1000 μM corroborated the observations on the different solid media. Essentially no differences were seen between the growth rates of the wild type and the $\Delta csoR$ strain under all conditions (Table 2 and supplemental Fig. S5).

Challenging the wild type and the $\Delta csoR$ strain with stresses that could affect copper homeostasis, such as diamide (redox stress) and hydrogen peroxide, showed a similar response in both strains (data not shown), as is the case for the addition of the iron chelator bathophenanthroline disulfonic acid to the medium (supplemental Fig. S4A). However, when the *in vivo* maturation of two cuproenzymes was measured, a small but consistent difference between wild type and the $\Delta csoR$ strain was observed. Both the activity of cytochrome *c* oxidase and the secreted heterologous tyrosinase (MelC2) are significantly higher in the $\Delta csoR$ strain (Fig. 4). Because deletion of the *csoR* gene is not likely to affect the expression of the endogenous *cox* genes (as confirmed by the RNA-seq data, see supplemental Table S6A) or the heterologous *melC* operon, these data would suggest that maturation/incorporation of the copper cofactor is more efficient in the $\Delta csoR$ strain.

X-ray Crystal Structure of Apo-CsoR^{SI}—The crystal structure of apo-CsoR^{SI} was determined to 1.7 Å resolution. Two protomers (chains A and B) were found in the crystallographic asymmetric unit of a crystal of CsoR^{SI} with well defined electron density visible for residues 44–133 in each protomer. The overall protomer fold (residues 44–133) consists of three α -helices of varying lengths. No electron density was visible for residues 1–43 suggesting that these residues are disordered. By applying crystallographic symmetry, a tetramer assembly, consistent with AUC, was generated with chain A packing against chain C and chain B packing against chain D (Fig. 5A). A surface representation of the tetramer is shown in Fig. 5B indicating that CsoR^{SI} has a “closed” tetrameric assembly as opposed to a donut-like structure reported for a CsoR-like protein from *Thermus thermophilus* and CsoR^{M^{tb}} (7, 51). The overall fold of

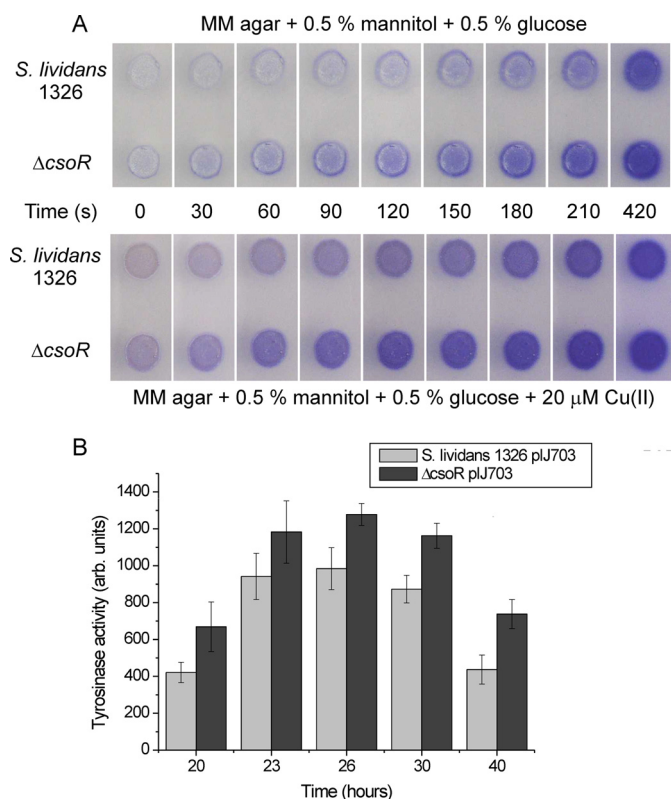


FIGURE 4. Effect of $\Delta csoR$ on cytochrome *c* oxidase and tyrosinase activity in *S. lividans* 1326. A, 1000 spores were spotted on MM agar plates and incubated for 50 h at 30 °C. The *in vivo* activity of cytochrome *c* oxidase was determined using *N,N,N',N'*-tetramethyl-*p*-phenylenediamine (TMPD) by monitoring the production of the blue compound than the wild type both on plates with and without added Cu(II). B, tyrosinase activity in the spent medium was determined with 3,4-dihydroxy-L-phenylalanine as substrate at the indicated time points and normalized for the biomass. The average tyrosinase activity of four transformants is plotted.

CsoR^{SI} and the packing of the three α -helices of each protomer within the tetramer assembly are similar to that in the homodimer structure of Cu^I-CsoR^{M^{tb}} (Fig. 5C) (7). Superposition of the two structures by secondary structure matching gave a root mean square deviation in C α positions of 1.98 Å. An extended α -helix 3 in CsoR^{SI} appears to be the reason for the closed assembly such that in apo-CsoR^{SI} the C termini of each pair of protomers pack together. Further differences between these structures are observed in the hairpin loop connecting α -helices 2 and 3, and at the N terminus of α -helix 1 (Fig. 5C). The largest differences are in the vicinity of the copper-binding sites, with residues 104–106 having deviations of >4 Å. The electrostatic potential of the CsoR^{SI} tetramer reveals a central region of strong positive potential at the interface of two homodimers that extends out toward helix 1 (Fig. 5B). Negative potential is located at the start and end of helix-2 and flanks the central positive potential in the tetramer assembly (Fig. 5B).

From the sequence alignment in Fig. 1, Cys-75, His-100, and Cys-104 are predicted to be the copper ligands in CsoR^{SI}. In the structure of the tetrameric assembly, His-100 and Cys-104 are located toward the end of helix 2 in each protomer, and Cys-75' is located on a loop connecting α -helices 1 and 2 of an opposite protomer creating a putative intersubunit binding site, as found in CsoR^{M^{tb}}. The side chain of Cys-75 has been modeled in two

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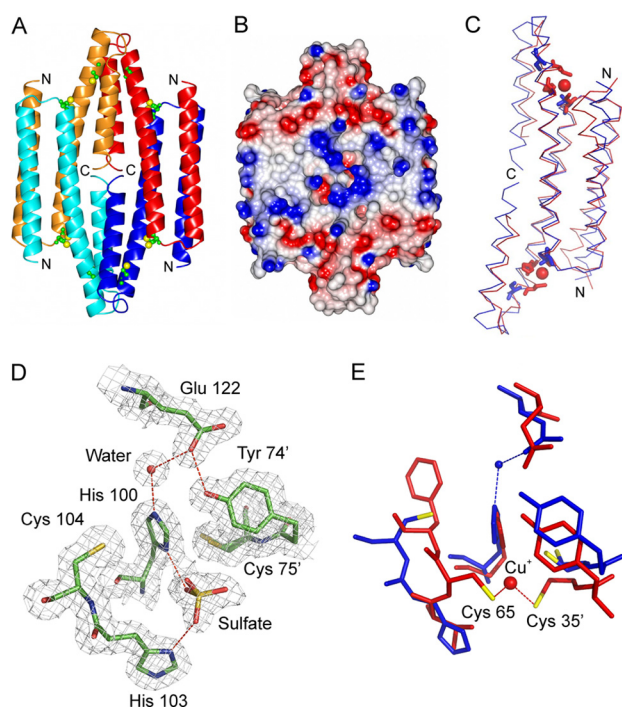


FIGURE 5. Structure of apo-CsoR^{SI}. *A*, physiologically relevant tetramer assembly, with protomers colored *red* (*A*), *orange* (*B*), *blue* (*C*), and *cyan* (*D*). The close packing of the C termini of all chains is apparent. The Cys residues predicted to be involved in copper binding are shown as *sticks* with the *S_γ* atom colored *yellow*. *B*, electrostatic surface representation of the CsoR^{SI} tetramer. Positive charges are indicated in *blue* and negative charges in *red*. *C*, superposition (calculated in the program Superpose, part of the CCP4 suite), based on secondary structure matching, of protomers *A* and *C* of the CsoR^{SI} structure (*blue*) with the dimer of CsoR^{Mtb} (*red*). The copper-binding Cys residues from CsoR^{Mtb} and the corresponding residues in CsoR^{SI} are shown as *sticks*. *D*, $2F_o - F_c$ electron density map, contoured at 1σ for the putative copper-binding region in the structure of CsoR^{SI}. Hydrogen bonds are shown as *dashed red lines*. The proposed copper binding residues Cys-75 and Cys-104 lie some 7.6 Å apart and are separated by the imidazole side chain of His-100, which forms a hydrogen bond to a well ordered water molecule. A sulfate anion is present at the protein surface, forming bonds to His-100 and His-103. *E*, comparison of the copper-binding region in CsoR^{SI} and CsoR^{Mtb}, based on the superposition shown in *C*. *A* and *B* prepared using CCP4MG (56) and *C–E* prepared using PyMOL.

slightly different conformations with occupancies of 0.7:0.3 (Fig. 5*D*), with the *S_γ* atom pointing either toward or away from the His-100 side chain. It is apparent that, in the absence of a copper ion, the His-100 imidazolate in apo-CsoR^{SI} intersects the two Cys ligands, which are some 7.6 (monomer *A*)/7.3 (monomer *B*) Å apart such that no disulfide bond between the two Cys residues is formed (Fig. 5*D*). The Ne2 atom of His-100 forms an H-bond (2.7/2.6 Å) with a well ordered water molecule (Wat1), which further hydrogen bonds (2.7/2.8 Å) to the side chain Oe1 atom of Glu-122 in α -helix 3 (Fig. 5*D*). A further H-bond interaction (2.6/2.7 Å) involving the Oe1 atom of Glu-122 and the OH atom of Tyr-74' of an adjacent protomer is also observed (Fig. 5*D*). A number of sulfate ion-binding sites are present in the structure, consistent with the ammonium sulfate component of the crystallization solution. One of these sites is located in the vicinity of each of the putative copper-binding sites. This sulfate anion is clearly positioned to be within H-bond distance of the N δ 1 atom of His-100 (2.6/2.6 Å) and 2.9/3.1 Å from N δ 1 of His-103 (Fig. 5*D*). It is interesting to note that the position of the sulfate anion is close to that of the

modeled copper ion in the 2.55 Å resolution CsoR^{Mtb} structure (7).

In Silico Identification of Putative CsoR^{SI} Operator Targets—The structural similarity of the 4136 gene product to Cu^I-CsoR^{Mtb} adds further credence to a role as repressor in *S. lividans*. Based on experimentally validated *cis*-acting sequences bound by *B. subtilis*, *S. aureus*, and *M. tuberculosis* CsoR orthologues (7, 15, 17, 38), we bioinformatically predicted putative CsoR^{SI} operator sequences in *S. lividans*. Regulons tend to be highly conserved in distantly related species, and therefore to ensure the identification of highly reliable CsoR^{SI} putative target genes, the computational prediction was performed in four *Streptomyces* species, *S. coelicolor*, *S. scabies*, *S. avermitilis*, and *S. griseus* (supplemental Tables S2–S5), so as to increase the reliability of *cis*-acting element predictions. Predictions from *S. coelicolor* were used to identify putative CsoR operator sequences in *S. lividans*. Three target sequences were selected from a stringent prediction procedure for further *in vitro* analyses. The highest score was 21.69, and the target sequence (AAATACCCCTGGTGGGTATAT) was located –42 nucleotides upstream from the start codon of *csoR* and –183 nucleotides upstream from the start codon of 4137 encoding a putative phosphate transport regulator (Fig. 6*A*). The identification of the most highly reliable operator sequence upstream of *csoR* itself strongly supports the validity of the prediction procedure and suggests autoregulation of CsoR^{SI} expression as reported for CsoR orthologues in other bacteria (7, 16, 17). The other two targets had scores of 19.37 (TTATACCCCTAGGGGTAAGG) and 14.15 (GGGTACCCCTAGGGGTATAC) and were found to be located –25 nucleotides upstream of the gene 2730 and –83 nucleotides upstream of the gene 1045 (Fig. 6*A*). Both of these genes are part of a *copZA*-like operon, predicted to encode a CopZ-like copper metallochaperone protein and a CopA P₁-type ATPase, which in tandem operate as a specific Cu(I) efflux system in many bacterial systems. Based on our computational predictions, the deduced consensus binding sequence of streptomycetes CsoR orthologues corresponds to the 21-nt palindromic sequence ATATACCCCT-NAGGGGTATAT, where positions 3–8 and 14–19 (underlined) appear to be the most conserved and thus probably the more crucial for CsoR^{SI} recognition (supplemental Figs. S6 and S7).

EMSA Analysis of CsoR^{SI} Operator Targets and Assembly State of DNA-CsoR^{SI} Complex—To test whether the three operator sequences identified with PREDetector would bind CsoR^{SI} *in vitro*, EMSAs were carried out. These were initially performed with intergenic regions consisting of 240 bp (1044/1045), 232 bp (2729/2730), and 208 bp (4136/4137) and with smaller 35/36-bp DNA oligomers, containing the CsoR^{SI} operator sequence flanked by 10 or 11 bp (Fig. 6*A*). Incubation of the intergenic regions or oligomers with apo-CsoR^{SI} resulted in the formation of a low mobility CsoR^{SI}-DNA complex visualized by the retardation of the DNA in the EMSA (Fig. 6, *B* and *C*). No shift in mobility of a random DNA sequence was observed suggesting that binding of the targets to CsoR^{SI} is specific (Fig. 6*C*). Incubation of the DNA oligomers with Cu^I-CsoR^{SI} samples or anaerobic addition of Cu(I) to preincubated apo-CsoR^{SI}-DNA complexes resulted in the absence of a band shift, and only the

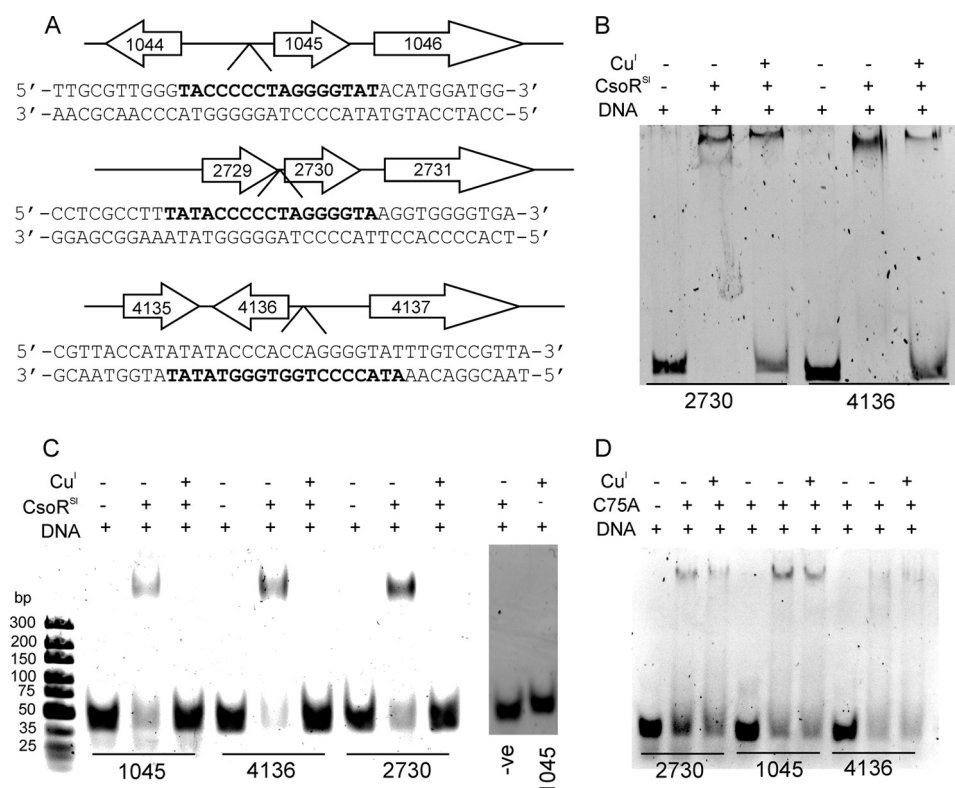


FIGURE 6. CsoR^{SI} operator DNA targets and EMSA studies. *A*, three-locus regulon and the DNA sequences identified by PREDetector (**bold**) to be target operator binding sites for CsoR^{SI}. The position of the DNA target sequence in the oligomers used in EMSA is highlighted in **bold**. *B*, EMSA of CsoR^{SI} in the presence of the intergenic DNA fragments containing the target sequences for 2730 (232 bp) and 4136 (208 bp). *C*, EMSA of CsoR^{SI} in the presence of the individual DNA oligomers containing the target sequences shown in *A*. The gel image to the *right* is a control EMSA showing the effect on binding CsoR^{SI} to a randomly generated double-stranded DNA oligomer with the sequence 5'-GACGAGGACGTCTACGCCATCGACATACTG-3' and the mobility of the 1045 DNA oligomer in the presence of only Cu(I). *D*, EMSA of the C75A mutant of CsoR^{SI} in the presence of the individual DNA oligomers containing the target sequences shown in *A*. The components present in each lane of the gel image are indicated with 0.5 μ M [DNA] and 5 μ M [apo-CsoR^{SI} monomer] and [Cu^I-CsoR^{SI} monomer] used.

high mobility band corresponding to the free DNA target was observed (Fig. 6C). This strongly suggests that once Cu(I) is bound to CsoR^{SI}, the affinity for these small DNA fragments is significantly reduced such that a complex is not detected by EMSAs. For the intergenic regions, similar behavior is observed, but it is noted that some low mobility complex remains (Fig. 6B). DNA incubated with only Cu(I) was not affected in mobility (Fig. 6C). To determine the DNA:CsoR^{SI} binding stoichiometry, size-exclusion chromatography was used (supplemental "Materials and Methods"). This revealed a 1:8 ratio (DNA:CsoR^{SI} monomer) or two tetramers of CsoR^{SI} are required to bind the DNA operator (supplemental Fig. S8).

Effect of Mutating Residues Cys-75 and His-100 in CsoR^{SI}—Size-exclusion chromatography and AUC measurements verified that the mutants do not disrupt the tetramer assembly state, and AAS indicated the absence of stoichiometric copper in the purified proteins. Both mutants showed an absorbance increase at 240 nm upon addition of Cu(I) under anaerobic conditions, saturating at ~ 1 mol eq of Cu(I) (Fig. 7A and supplemental Fig. S9), and for the H100A mutant, the absorbance increase at ~ 320 nm that was observed in the wild type protein upon addition of Cu(I) was absent (Fig. 7A). The K_D (Cu^I) values for the mutants were determined using BCA as a competitive probe (Fig. 7B). Duplicate data sets at pH 7.5 were acquired for both mutants with varying [Cu^I]_{total} or [BCA], and

the data averaged to give a K_D (Cu^I) of 1.3×10^{-14} M and 9.8×10^{-16} M for the C75A and H100A variants, respectively. Both mutants therefore display an increased K_D (Cu^I) compared with wild type CsoR^{SI}.

The mutants were each able to bind to the three DNA operators as indicated by EMSA (Fig. 6D). Addition of Cu(I) to the mutant-DNA operator complexes did not result in the disappearance of the low mobility band or a substantial increase in the intensity of the high mobility (free) DNA band (Fig. 6D). This suggests that Cu(I) binding in the mutants does not alter the affinity for the DNA operator to the extent observed for the wild type and suggests that the allosteric mechanism of operator release or exposure is affected. This apparent inability of the CsoR mutants C75A and H100A to disengage *in vitro* from the DNA operator following Cu(I) binding raises the question of what the effect on growth is when these mutant proteins are expressed in the mycelium. The wild type and mutated CsoR^{SI} genes were cloned on a low copy plasmid under control of their own promoter. These constructs did not show any effect on growth. Therefore, the CsoR^{SI} wild type gene and the mutant genes encoding the C75A and H100A variants were cloned under control of a strong constitutive promoter, and the wild type and Δ csoR strain were transformed with these constructs. Fig. 8 clearly shows that the mutant forms of CsoR^{SI} have a strong inhibitory growth effect in both strains starting from 100 μ M Cu(II). The effect of mutant CsoR expression is slightly

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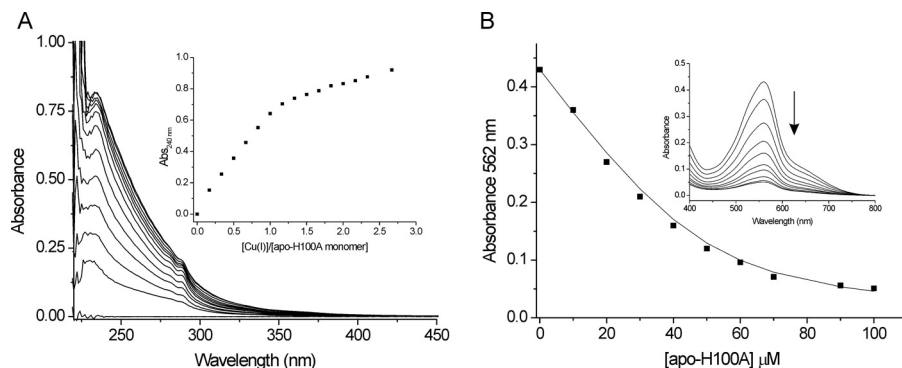


FIGURE 7. **H100A mutant binds Cu(II) with a reduced affinity.** *A*, changes in the UV region of the apo-H100A mutant base-lined spectrum upon addition of Cu(II)Cl at pH 7.5 and 20 °C. The *inset* shows the increase in absorbance at 240 nm plotted as a function of [Cu(II)]/[apo-H100A monomer]. *B*, determination of the $K_D(\text{Cu}^I)$ for apo-H100A under copper-limiting conditions imposed by $[\text{Cu}^I(\text{BCA})_2]^{3-}$ as a competitive probe. The *inset* shows the decrease in absorbance at 562 nm in the visible region of the spectrum upon increasing [apo-H100A], and Equation 1 was used to determine the $K_D(\text{Cu}^I)$. The *line* represents a best fit to the data using $K_D(\text{Cu}^I)$ 6.3×10^{-16} M. Conditions used are as follows: 10–100 μM [apo-H100A], 54 μM $[\text{Cu}^I]_{\text{total}}$, and 260 μM [BCA].

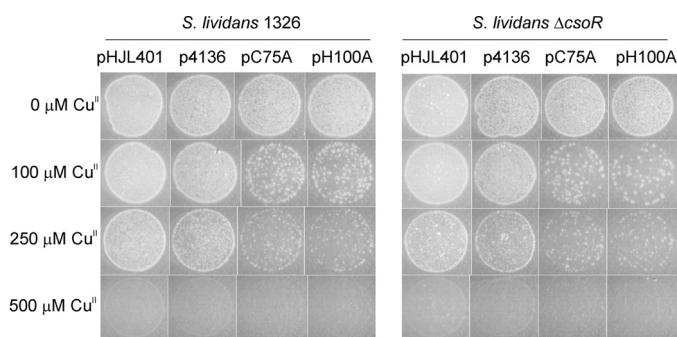


FIGURE 8. **Effect of C75A and H100A CsoR^{SI} mutants on growth.** *S. lividans* 1326 and the ΔcsoR mutant were transformed with pHJL401 (empty vector) or the same vector harboring the *csoR* gene (p4136) or the mutants C75A (pC75A) or H100A (pH100A) under control of the constitutive *ermE* promoter. Spores (1000) were spotted on MM agar plates containing varying [Cu(II)] and incubated at 30 °C. After 50–60 h of incubation, a confluent lawn was produced. Images were taken with a Leica M80 stereomicroscope equipped with a DFC295 camera. Note the higher toxicity of exogenous copper supply due to mutations preventing the copper-dependent modulation of CsoR^{SI} DNA binding ability.

more pronounced in the ΔcsoR mutant than in the wild type because the latter also produces CsoR from the genomic gene copy resulting in a mix of plasmid and genome encoded proteins. Upon prolonged incubation, the control transformants harboring the empty vector or the wild type CsoR^{SI} gene produce confluent growth with 250 and 500 μM Cu(II), although those expressing the mutant CsoRs do not. These observations support the *in silico* data indicating that CsoR^{SI} has control over the expression of genes that encode proteins involved in exporting excess copper ions and its own expression under copper stress conditions.

Mapping the Global Response of Wild Type S. lividans to Elevated Copper Levels and the ΔcsoR Gene by RNA-seq—RNA-seq is a second generation high resolution sequencing technique that enables gene expression profiling in an organism to be analyzed in response to a mutation or an external stimulus (52–54). We have applied RNA-seq to further our insight into the CsoR regulon in *S. lividans* by analysis of the transcriptome of liquid-grown cultures. The transcriptomes of wild type *S. lividans* strain 1326 grown without and with a 2-h exposure to 400 μM Cu(II) were compared with that of the ΔcsoR strain. From analysis of the RNA-seq data, both CopZ genes (gene numbers 2730 and 1045) and the 1044 gene encoding for an

uncharacterized secreted protein are all induced by exogenous copper and in the ΔcsoR strain (Table 3). These data corroborate the *in silico* approach and identify these genes as *bona fide* CsoR^{SI} targets. The CopA-like ATPases (2731 and 1046) are also up-regulated under elevated copper conditions (Table 3) but less so in the ΔcsoR strain. Nevertheless, this suggests regulation by the same promoter as the cognate CopZ. It is noticeable that the RNA-seq data do not provide clear support for the copper induction of gene 4137 or the *csoR* gene, which was predicted from PREDetector analysis (Table 3 and supplemental Table S2). Therefore, the promoter of *csoR* was analyzed in a promoter-probing experiment. The data clearly show that transcription originating from the *csoR* promoter is both copper-inducible and CsoR-dependent (Fig. 9A). The low induction by a 2-h exposure to 400 μM Cu(II) seen in the RNA-seq analysis (1.3-fold, Table 3) is confirmed by the promoter probing experiment (Fig. 9A). It also shows that upon longer exposure to Cu(II) the transcription is induced to around 2-fold, a similar level as observed in the ΔcsoR strain.

Finally, the transcriptome response to either copper or deletion of the *csoR* gene is not limited to the genes reported in Table 3. The global response can be appreciated visually by the use of Venn diagrams (Fig. 9B). A large number of genes are up- and down-regulated in response to copper and deletion of the *csoR*, with a significantly greater proportion of genes being down-regulated in the ΔcsoR strain. A considerable overlap is present between copper induction and *csoR* deletion (Fig. 9B and supplemental Table S6, B–G). Together, these data indicate that the response to changes in copper homeostasis in *S. lividans* is much more extensive than only the direct CsoR^{SI} regulon reported in Table 3. Aside from the genes up-regulated in Table 3, there is no obvious enrichment of genes that encode proteins known to be directly related to copper homeostasis. However, a gene for a putative copper transporter (3964), part of an operon encoding for two putative copper chaperones, is down-regulated in copper-induced *S. lividans* but not in the *csoR* deletion (supplemental Table S6, E–G).

DISCUSSION

At the molecular level, our understanding of proteins involved in copper detoxification, storage, and trafficking has

TABLE 3

Expression levels obtained by RNA-seq of the genes predicted by PREDetector to be under control of CsoR^{SI} along with other putative copper proteins up-regulated

Annotation	Gene	PREDetector score	<i>S. lividans</i> 1326 RPKM ^a	<i>S. lividans</i> 1326 copper-induced (RPKM) ^a	-Fold increase	Δ csoR (RPKM) ^a	-Fold increase
CsoR	4136	21.7	169.8	217.5	1.3		
Pit accessory protein	4137	21.7	418.2	470.5	1.1	496.3	1.2
Membrane protein	3280	21.3	21.7	112.8	5.2	20.0	0.9
CopZ	2730	19.4	86.7	436.0	5.0	220.9	2.5
Secreted protein	1044	14.2	4.0	17.1	4.2	7.2	1.8
CopZ	1045	14.2	0.0	9.8	>>	5.8	>>
CopA-like P ₁ -type ATPase ^b	2731		32.4	65.5	2.0	25.5	0.8
CopA-like P ₁ -type ATPase ^b	1046		3.4	9.2	2.7	4.8	1.4
Uncharacterized protein with CusF domain ^c	7265		3.2	23.0	7.3	18.0	5.7
Uncharacterized protein with a putative type 1 copper site	6710		0.0	6.3	>>	2.1	>>

^a The selected expression measure is the RPKM. It is defined as the reads/kb of exon model/Million mapped reads (25).

^b Genes 2731 and 1046 are in the same operon as 2730 and 1045, respectively.

^c Gene 7265 is the only putative copper-trafficking linked gene that is induced both by copper stress and by *csoR* deletion (see Table S6A for all putative copper proteins).

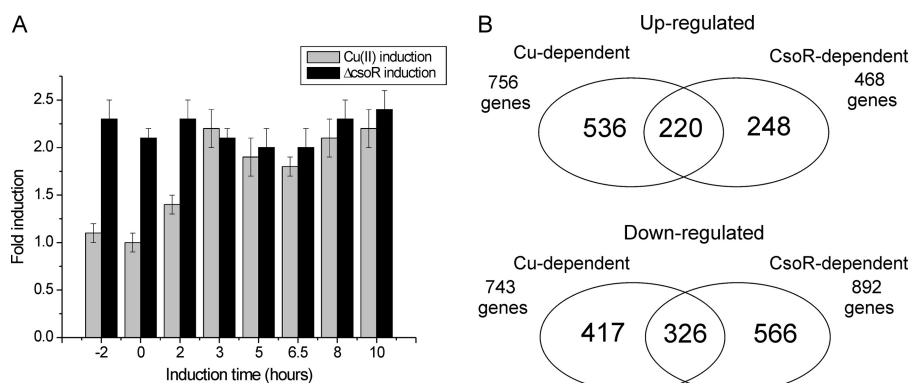


FIGURE 9. Induction of *csoR* by promoter probing and global differential gene expression. *A*, promoter activity in response to the addition of 400 μ M Cu(II) in the wild type and the Δ csoR strain are expressed as fold change relative to the wild type strain without addition of Cu(II). Note the copper-induced Red production in the *S. lividans* wild type background and the constant and copper-insensitive Red production in the *S. lividans* Δ csoR background. *B*, Venn diagrams depicting a global overview of the up- and down-regulated genes (>2-fold reads/kb of exon model/million) upon Cu(II) induction of wild type *S. lividans* and in the Δ csoR strain. The supplemental Table S6, A–G, reports the lists of genes affected.

advanced considerably in recent times (1, 3, 4). The next challenge is to interface our molecular and mechanistic understanding of copper homeostasis with the global response of an organism to copper overload. In this study, we have extensively characterized a Cu(I)-CsoR metalloregulator from *S. lividans* and uncovered the direct and extended CsoR regulon.

Cu(I) Affinity for CsoR Is Unified but How DNA Is Engaged Remains Unresolved—Recent reports have highlighted the need for a unifying approach to accurately determine Cu(I) binding affinities in homeostatic cuproproteins (36, 50). The use of the Cu(I)-specific chelators BCA and BCS has been strongly advocated, and these have been used in this work. Accurate determination of metal affinities of metal sensors is particularly important as these values are considered the thresholds for homeostasis (2). By using BCS, an upper limit K_D (Cu^I) of 1×10^{-20} M was initially reported for CsoR^{Mtb} (7) and later revised to 1×10^{-19} M (55). For CsoR^{Bs}, an upper limit of 1×10^{-21} M has been reported using BCS (38) and most recently a K_D (Cu^I) of 7.9×10^{-19} M for CsoR^{Sa} (17). The K_D (Cu^I) value determined for CsoR^{SI} at pH 7.5 is 6.7×10^{-18} M, an order of magnitude higher than CsoR^{Sa} and CsoR^{Mtb}, and thus a comparatively lower affinity for Cu(I). These studies clearly highlight the consistency in using BCS as a probe to determine Cu(I) binding affinity in the attomolar range, enabling for cross-species comparisons. The K_D (Cu^I) values determined using BCS are all consistent with the CsoR sensors being

triggered by [copper] \sim 9 orders of magnitude higher than the minimal intracellular copper concentration, consistent with the bacterial cytosol having no free atoms of copper (5). Furthermore, this clearly indicates the lengths to which cells go to avoid the potentially deleterious effects of uncomplexed copper ions.

At 1.7 Å resolution, the crystal structure of apo-CsoR^{SI} is the highest resolution structure to date for a member of the CsoR family. The primary structure of CsoR^{SI} is unique compared with other characterized CsoR proteins in that it possesses an extended (43 amino acids) N-terminal tail (Fig. 1). Structural organization of this region was not revealed from the crystal structure as electron density was only observed from residue 44 onward. Likewise, the C-terminal extension in CsoR^{Mtb} (7) was not structurally observed, and in the absence of proteolytic cleavage, dynamics outside the “core” structure are likely to be relevant. The core structure of CsoR^{SI} is very similar to that of Cu^I-CsoR^{Mtb} (7) and has no recognizable DNA structural binding motif, such as a helix-turn-helix motif. Despite this, apo-CsoR^{SI} has high specificity and affinity for operator DNA targets (Fig. 6). The absence of a recognizable DNA-binding motif leaves the question of how CsoR orthologues engage with their operator target unanswered. From our structure of CsoR^{SI}, we observe that the tetramer assembly creates a large continuous surface area with strong electropositive charge centered in the middle of each face of the tetramer, which may be of signifi-

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cance for DNA binding (Fig. 5B). An alternative possibility is that the absence of a DNA-binding motif is the reason why two tetramers associate with the DNA, as also reported for CsoR^{Bs} (38), so as to enhance binding, specificity, or both.

Apo-CsoR^{SI} Structure Provides Insights into the Mechanism of Allosteric Regulation—An hypothesis for the mechanism of allosteric Cu(I) regulation in the CsoR family has been put forward based on the crystal structure of Cu^I-CsoR^{Mtb} (7) and tested experimentally using unnatural amino acid substitutions of the copper -coordinating His residue (55). These unnatural amino acids were designed to abolish the H-bonding capability of the Ne2 atom of the imidazole ring so that, on binding Cu(I) to the Nδ1 atom, a second coordination sphere H-bond network centered on the Ne2 atom could not form and trigger release of the DNA (Fig. 5E). Cu(I) binding to the non-native His-substituted CsoR^{Mtb} did not significantly affect the DNA binding affinity, but the allosteric coupling free energy (ΔG_c) was determined to be close to zero (55). This observation was taken to indicate that allosteric switching is initiated upon binding Cu(I) to the Nδ1 atom of the His ligand, triggering the formation of a H-bond network to the Ne2 atom that results in dissociation from the operator DNA sequence. However, this H-bond network is also present in the apo-CsoR^{SI} structure and suggests that copper binding is not an essential requirement for its formation (Fig. 5E). One difference in the network found in CsoR^{SI} is the presence of a bridging water molecule between His-100 and Glu-122 (Fig. 5D). In the Cu^I-CsoR^{Mtb} structure, no such water molecule is observed, and the H-bond length between the Cu(I) coordinating His and the Glu is >4 Å (7). This is indicative of a very weak interaction, which may lead to a destabilization of the DNA-bound structure. Whether or not this water molecule plays a role in the mechanism of allosteric Cu(I) regulation or in DNA binding is presently unclear. From Fig. 5E, it is clear that His-100 and Cys-75' in the apo-CsoR^{SI} structure are closely aligned to the corresponding residues in the Cu^I-CsoR^{Mtb} structure. However, Cys-104 in apo-CsoR^{SI} must undergo a significant movement to complete the Cu(I) coordination sphere (Fig. 5E). It is conceivable that the movement associated with the positioning of the Cys-104 side chain to enable Cu(I) coordination may be of significance in allosterically regulating the dissociation from the DNA operator.

CsoR Regulon in Streptomycetes—A CsoR-responsive element prediction in four different *Streptomyces* species (supplemental Tables S2–S5) confirmed a very limited occurrence of highly conserved sequences that have been identified only upstream of one or two sets of orthologues of the copper chaperone *copZ* and the efflux ATPase *copA*, as well as upstream of the copper -sensing repressor *csoR* itself. EMSA studies (Fig. 6) revealed that CsoR^{SI} recognizes specifically these sequences, and the effect of Cu(I) on DNA binding supports a mechanism of allosteric regulation most likely similar to other CsoR orthologues (7, 17, 38). RNA-seq data corroborate the *in silico* predictions, revealing clear induction of transcripts for two operons encoding copper efflux systems (2730/2731 and 1045/1046) (Table 3) and the divergent expression of 1044 (Table 3), a secreted protein of unknown function but not predicted to bind copper. RNA-seq data also indicates that CsoR^{SI} transcript levels, under normal conditions, are relatively high compared

with the two *copZ* genes and suggest that a significant proportion of CsoR^{SI} is constitutively present (Table 3). The observation that expression of the mutant *csoR* genes (C75A and H100A) under control of their own promoter does not lead to growth inhibition at higher [copper] illustrates that CsoR^{SI} controls its own transcription. Failure of the mutants to disengage from the DNA operator has been reported for other copper ligand variants of CsoR orthologues (7, 16, 17) and appears to lead to permanent transcription repression, resulting in too low protein expression levels to block transcription of the other genes with CsoR^{SI}-binding sites. The induction of *csoR* was not observed from the RNA-seq data (Table 3) but was confirmed after >2 h of incubation with exogenous copper (Fig. 9A). This suggests the possibility that a modular system of CsoR repression may be in operation. Interestingly, the *in silico* analysis in *S. griseus* predicted four other CsoR-responsive elements suggesting a somewhat wider regulon in this strain (supplemental Table S5). The *SGR3189* (*cutC*) gene encoding a putative copper homeostasis protein (*cutC*) is located upstream of a *copA*-like gene but is divergently transcribed. This offers the possibility that the CsoR-responsive elements could act for both *copA* and *cutC* in *S. griseus*. Two other CsoR-binding sites are found at position –30 nt from the *SGR5260* gene, which could encode for the first member of an operon, including a putative multicopper oxidase (*SGR5259*) and at position –36 nt from *SGR1262* (*cstR*) encoding for a putative non-copper sensing member of the CsoR family (17).

A Model of the Directly Regulated CsoR Regulon in *S. lividans*—Our findings clearly show that CsoR^{SI} directly acts to regulate a three-locus regulon. Under homeostasis conditions, all operator sequences are occupied by two tetramers of CsoR^{SI} resulting in repression of transcription with some free apo-CsoR and some copper -bound CsoR and CopZ (2730) acting as buffers (Fig. 10). Upon elevated cytosolic copper levels, we suggest, based on our data, a three-step response of the system. Low [copper] can be buffered by free apo-CsoR and CopZ (2730), and no further action is required. Medium [copper] will require a stronger response, achieved by derepressing the 2730 and 1044/1045 operons resulting in expression of CopZ and the CopA P₁-type ATPases (Table 3). A further increase in [copper] will result in copper binding by the apo-CsoR^{SI} still occupying the *csoR* operator. As a consequence, CsoR will be produced at a higher level and will assist in buffering copper in the cytoplasm. We assume that CsoR can also mediate the trafficking of copper by donating its copper to CopZ, which in turn will have the specificity to deliver the metal to its cognate CopA P₁-type ATPase for export. As soon as copper levels are restored, apo-CsoR^{SI} begins to occupy the operator sequences, and the system returns to its “ground state.” This model assumes that the up-regulation of CsoR^{SI} occurs later. Quite how this may be possible is not known, but promoter probing and RNA-seq data are consistent with this phenomenon.

What Can Be Learned About Copper Homeostasis in *S. lividans* from RNA-seq Analysis?—Although RNA-seq analysis clearly provides insight into the response of transcripts under the direct control of CsoR^{SI} (Table 3), it is very much evident that response to exogenous copper or the deletion of the *csoR* gene is a complex process (Fig. 9B and supplemental Table S6,

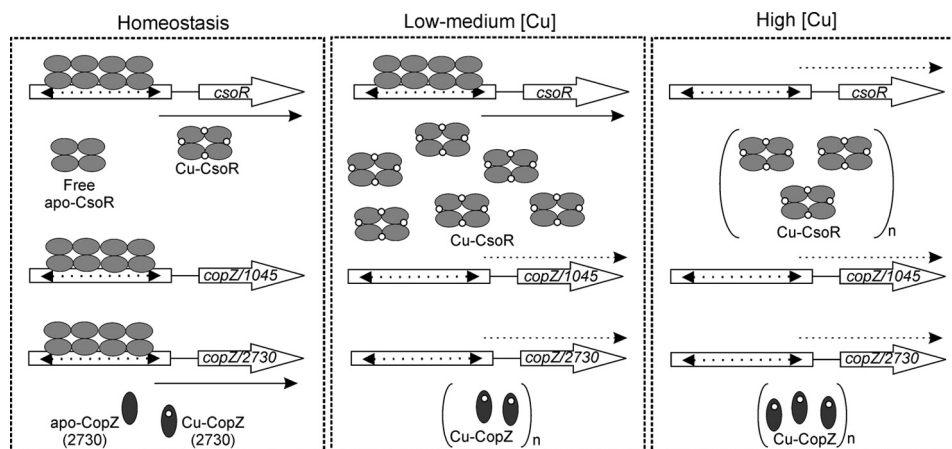


FIGURE 10. **A model of CsoR regulon and its response to copper stress in *S. lividans*.** Under normal conditions, both *csoR* and 2730 (CopZ) are expressed at a low CsoR-independent level to ensure an immediate response to an increase in [copper] is present (left panel). Upon higher copper levels, the apo-CsoR and apo-CopZ can buffer copper at first, but as soon as copper levels increase, the free apoproteins will all become copper-bound, and the repression of 2730 and 1045 will be relieved because of Cu(I) binding to the DNA-bound CsoR resulting in production of sufficient export capacity to maintain homeostasis (middle panel). Upon further increase of [copper] the repression of *csoR* will be removed, and the full response becomes available (right panel).

A–G). Over 400 genes show significantly higher expression in the $\Delta csoR$ strain without presenting putative CsoR^{SI} binding sequences in their adjacent genetic environments. Therefore, the expression of these genes must be responding in an indirect way to CsoR^{SI} and may possibly be due to the copper-binding ability of CsoR^{SI} rather than its DNA-binding ability. Indeed, the inactivation of *csoR* would somehow mimic copper overload due to the absence of free cytosolic apo-CsoR^{SI} involved in buffering excess copper.

Of the genes annotated to encode cuproenzymes and putative copper transport proteins, the expression of some could be CsoR^{SI}-dependent (supplemental Table S6A). For example, from the initial PREDetector analysis of the streptomycetes CsoR regulon, a fourth type of gene that could potentially be controlled by CsoR^{SI} was identified (supplemental Table S2). Gene 3280 encodes for a membrane protein, but experimental validation of the predicted site was not included in our present EMSA studies due to the unusual location of the predicted *cis*-acting element at position +24 nt within the predicted 3280 coding sequence. Similar sequences identified at position +4 and –186 nt relative to the translational start of the 3280 orthologue in *S. avermitilis* and *S. scabies* are more appropriately located for controlling gene expression by a transcriptional repressor (supplemental Tables S3 and S4). The RNA-seq data highlighted a 5.2-fold higher expression of 3280 in RNA samples collected from copper-induced cultures, whereas the expression level in the $\Delta csoR$ remains unchanged (Table 3). This suggests that 3280 is involved in copper homeostasis, and an as yet unidentified transcription control system plays a role when CsoR^{SI} is no longer present due to the gene deletion. A closer inspection of the 3280 product predicts a protein (~200 amino acids) with four transmembrane helices and a putative Cu(I) Cys-Xaa-Xaa-Xaa-Cys-binding motif at the start of the predicted first transmembrane helix.

The 220 genes that are induced by both exogenous copper and $\Delta csoR$ (Table S6D) indicate that the CsoR^{SI} and copper overload regulons only partially overlap with each other. One of these overlap genes is 7265, annotated as a membrane protein with a CusF domain and therefore implemented in copper

binding. RNA-seq data reveal this gene to have the greatest fold increase of all known and predicted genes that encode for copper-binding proteins/transporters (Table 3) and may therefore have an as yet uncharacterized role in copper homeostasis. Gene 6710 encoding a protein containing a putative type I copper site is not expressed in the control culture but does show a low level of expression in the *csoR* mutant and upon copper induction. However, it is not clear at present how this protein could be involved in copper homeostasis. The strongest transcription increase both in response to copper overload and in the $\Delta csoR$ mutant, by 29- and 15-fold, respectively, is a putative [2Fe-2S] thioredoxin-like protein (SCO5830). A functional reason for this is unclear as is the case for the hundreds of other genes that are induced by either copper or by the $\Delta csoR$. It is reasonable to assume from these data that besides CsoR^{SI} control, one or most likely several other control systems operate in copper homeostasis, e.g. redox homeostasis. One possibility may be that the way the copper excess regulon is stimulated requires more sophisticated and efficient mechanisms than the slow and energy-consuming means involving transcriptional control by a DNA-binding protein.

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