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# The ROK Family Regulator Rok7B7 Pleiotropically Affects Xylose Utilization, Carbon Catabolite Repression, and Antibiotic Production in *Streptomyces coelicolor*

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**Members of the ROK family of proteins are mostly transcriptional regulators and kinases that generally relate to the control of primary metabolism, whereby its member glucose kinase acts as the central control protein in carbon control in *Streptomyces*. Here, we show that deletion of SCO6008 (*rok7B7*) strongly affects carbon catabolite repression (CCR), growth, and antibiotic production in *Streptomyces coelicolor*. Deletion of SCO7543 also affected antibiotic production, while no major changes were observed after deletion of the *rok* family genes SCO0794, SCO1060, SCO2846, SCO6566, or SCO6600. Global expression profiling of the *rok7B7* mutant by proteomics and microarray analysis revealed strong upregulation of the xylose transporter operon *xylFGH*, which lies immediately downstream of *rok7B7*, consistent with the improved growth and delayed development of the mutant on xylose. The enhanced CCR, which was especially obvious on rich or xylose-containing media, correlated with elevated expression of glucose kinase and of the glucose transporter GlcP. In liquid-grown cultures, expression of the biosynthetic enzymes for production of prodigionines, siderophores, and calcium-dependent antibiotic (CDA) was enhanced in the mutant, and overproduction of prodigionines was corroborated by matrix-assisted laser desorption ionization–time-of-flight analysis. These data present Rok7B7 as a pleiotropic regulator of growth, CCR, and antibiotic production in *Streptomyces*.**

*Streptomyces* are high-G+C soil bacteria that are well known as major producers of antibiotics and natural products. Under nutrient-limited conditions they undergo complex morphological changes, which result in formation of aerial mycelium and chains of spores. The developmental program coincides with production of antibiotics (1). Sequencing of the genome of the model organism *Streptomyces coelicolor* A3(2) identified around 7,825 genes and more than 20 secondary metabolite gene clusters (2). Over 12% of the genome encodes predicted regulatory proteins, underlining the complexity of the regulatory networks that govern the transcriptional control of cell differentiation, secondary metabolite production, and signaling cascades.

Carbon source regulation is a major factor in the control of morphological differentiation and secondary metabolism (recently reviewed in references 3 and 4). We previously showed that changes in *N*-acetylglucosamine metabolism have a major effect on the production of pigmented antibiotics in *S. coelicolor* (5, 6). This is in part explained by the fact that glucosamine-6P is an allosteric effector of the global regulator DasR, which represses antibiotic production (7, 8), but it is likely that other factors play a role, including global carbon control mechanisms. Carbon catabolite repression (CCR) allows bacteria to utilize a preferred carbon source (such as glucose) when exposed to multiple carbohydrates. In Gram-negative bacteria and low-G+C Gram-positive bacteria, such as *Escherichia coli* and *Bacillus subtilis*, the phosphoenolpyruvate (PEP)-dependent phosphotransferase system (PTS) plays a major role in carbon regulation (9, 10). During PTS-mediated carbon source internalization, a phosphoryl group is transferred from the glycolytic intermediate phosphoenolpyruvate, via the PTS enzymes phosphotransferase enzyme I (EI), the histidine phosphor-carrier protein (HPr), and enzyme IIA (EIIA), onto the incoming carbon source. The PTS-mediated transport triggers

the cyclic AMP (cAMP)-receptor protein (Crp) in Gram-negative bacteria and the catabolite control protein CcpA in low-GC Gram-positive bacteria (9). In *Streptomyces*, GlcNAc and fructose transport depend on the PTS, but glucose transport is mediated via the major facilitator superfamily (MFS) transporter GlcP (11). In terms of global regulation, streptomycetes lack a CcpA orthologue, while Crp is present but does not play a role in CCR (12). In contrast, in streptomycetes CCR largely depends on the enzyme glucose kinase (Glc; SCO2126), and deletion of the *glkA* gene results in global loss of CCR (13). Interestingly, CCR is independent of the Glc enzymatic activity and the flux of glucose, and replacing Glc of *S. coelicolor* with that of *Zymomonas mobilis* restores normal glycolysis but not CCR (13).

The crystal structure of Glc of *Streptomyces griseus* was recently resolved (14). The *Streptomyces* Glc protein belongs to group III of the glucokinases, which contain a ROK signature (Pfam PF00480) that is found in a diverse family of proteins (ROK stands for repressors, open reading frames, and kinases) (15). Some 5,000 ROK-family proteins have been identified so far, mostly in prokaryotes, but family members are found in all kingdoms of life (16). ROK kinases contain a conserved N-terminal ATP binding

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motif (DxGxT), while ROK regulators contain a canonical helix-turn-helix DNA binding motif in their N-terminal domains. In *E. coli*, the ROK-family proteins NagC and Mlc are involved in the control of GlcNAc and glucose-PTS genes, respectively (17, 18). The *S. coelicolor* genome contains 25 genes encoding putative ROK-family proteins, 12 of which contain a DNA binding motif. Additionally, 16 of the ROK proteins contain a zinc binding motif, which conforms to the consensus GHX<sub>(9-11)</sub>CXCGX<sub>2</sub>G(C/H)XE and was shown to be essential for function of *E. coli* Mlc and *B. subtilis* Glk (19, 20). Interestingly, SCO6008 (Rok7B7) was identified as an activator of actinorhodin biosynthesis in a mass spectrometry (MS)-based assay for proteins that bind upstream of the regulatory gene *actII-ORF4* (21), and expression of a metagenomic library in *Streptomyces lividans* revealed that the *rok7B7* homolog *rep* accelerates antibiotic production and sporulation (22). Additionally, Rok7B7 was recently implicated in CCR in *S. coelicolor* (23).

In search of new regulatory genes that may play a role in carbon-dependent control of antibiotic production, we describe here the mutational analysis of seven of the *rok* genes of *S. coelicolor*. These seven genes all contain a DNA binding domain and a cysteine motif involved in zinc binding (24) and include the three closest relatives of *rep*. Deletion of *rok7B7* (SCO6008) resulted in global changes in antibiotic production and morphological differentiation under various growth conditions. Detailed phenotypic and genome-wide transcriptome and proteome analyses of the *rok7B7* null mutant established a relationship between Rok7B7 and CCR depending on the nutritional status of the environment.

## MATERIALS AND METHODS

**Bacterial strains and growth conditions.** All bacterial strains are listed in Table S1 in the supplemental material. *Escherichia coli* JM109 (25) and ET12567 (26) were used for routine cloning procedures and for extracting nonmethylated DNA, respectively. *Streptomyces coelicolor* A3(2) M145 was the parent for all mutants described in this work. *S. coelicolor* M512 (M145  $\Delta$ *actII-ORF4*  $\Delta$ *redD*) (27) lacks the pathway-specific activator genes for actinorhodin and undecylprodigiosin biosynthetic clusters and therefore fails to produce these antibiotics, although the strain has an intact biosynthetic machinery (28). All media and routine *Streptomyces* techniques are described in the *Streptomyces* manual (26). *E. coli* cells were grown in Luria-Bertani broth (LB) at 37°C. Phenotypic characterization of *Streptomyces* mutants was carried out on minimal medium (MM) and on R2YE<sup>-</sup> (i.e., lacking glucose) agar plates (26) with 1% carbon sources as indicated. SFM (soy flour mannitol) agar plates were used to prepare spore suspensions. For growth curves, liquid NNMP (26) was used with a low (0.1%, wt/vol) concentration of Casamino Acids. An adapted NNMP medium was used for <sup>14</sup>N/<sup>15</sup>N labeling, containing (per liter) 2 g NH<sub>4</sub>Cl, 1 g algal amino acids, 0.6 g MgSO<sub>4</sub> · 7 H<sub>2</sub>O, 50 g polyethylene glycol 6000 (NBS Biologicals, Huntingdon, United Kingdom), 0.1 mg ZnSO<sub>4</sub> · 7 H<sub>2</sub>O, 0.1 mg FeSO<sub>4</sub> · 7 H<sub>2</sub>O, 0.1 mg MnCl<sub>2</sub> · 7 H<sub>2</sub>O, 0.1 mg CaCl<sub>2</sub>, 15 mM Na/K phosphate buffer (pH 6.8), and 5 g mannitol. For <sup>15</sup>N labeling, NH<sub>4</sub>Cl (>99% <sup>15</sup>N) and algal amino acids (>98% <sup>15</sup>N) were added as <sup>15</sup>N-labeled compounds (Cambridge Stable Isotopes, Andover, MA).

**Plasmids and constructs.** All constructs described in this work are listed in Table S2 in the supplemental material.

**Constructs for gene replacement and deletion mutants.** The strategy for creating knockout mutants is based on the unstable multicopy vector pWHM3 (29) as described previously (11). For each knockout construct, roughly 1.5-kb stretches of upstream and downstream regions of the respective genes were amplified by PCR from the *S. coelicolor* M145 genome. The upstream region was thereby cloned as an EcoRI-XbaI fragment and the downstream part as an XbaI-HindIII fragment, and these were ligated into EcoRI-HindIII-digested pWHM3 (for the exact location of the oli-

gonucleotides, see Table S3 in the supplemental material). In this way, an XbaI site was engineered between the flanking regions of the gene of interest. This was then used to insert the apramycin (Apr) resistance cassette *aacC4* flanked by *loxP* sites, using engineered XbaI sites. The presence of the *loxP* recognition sites allows the efficient removal of the apramycin resistance cassette following the introduction of plasmid pUWLcre (expressing the Cre recombinase) (30, 31). We constructed the knockout plasmids pGAM8 to pGAM14 for the single gene replacement of SCO0794, SCO1060, SCO2846, SCO6008, SCO6566, SCO6600, SCO7543, respectively. To analyze correctness of the mutants, PCRs were done on mycelium from liquid-grown MM cultures (oligonucleotide pairs are described in Table S3).

For complementation of the GAM33 mutant ( $\Delta$ *rok7B7*), the integrative vector pSET152 (32) harboring a recombinant *rok7B7* gene, along with a 250-bp promoter region and the stop codon replaced by the sequence 5'-ATGGACTACAAGGACCACGACGCGGACTACAAGGAC CACGACATCGACTACAAGGACGACGACGACAAGTAGAGCACGG ATGGCACCGTTGTCATCTCGTTAAGGATTTACTTCTT, was used. The recombinant *rok7B7* gene was synthesized that allows production of Rok7B7 containing a C-terminal triple FLAG-tag epitope (GenScript, La Jolla, CA).

**Agarase assays.** Spores of *S. coelicolor* M145, its *rok7B7* null mutant GAM33, and the complemented mutant were spotted onto MM and R2YE<sup>-</sup> agar plates (26) with 1% (wt/vol) different carbon sources as indicated. After 3 days of growth, the agarase production was assessed by measuring the zones of clearing of the agar around the mycelial spot and related to the biomass. Experiments were done several times, with highly similar results.

**Proteomics analysis.** *S. coelicolor* M145 and its *rok7B7* null mutant, GAM33, were grown in liquid minimal media containing either <sup>14</sup>N or <sup>15</sup>N as the sole nitrogen source, as described above, until late logarithmic phase, when antibiotic production became apparent due to color changes of the growth medium. This was repeated for a label swap experiment in which the nitrogen sources were exchanged between strains.

At the indicated time points, mycelium was harvested by centrifugation, washed, and sonicated for 5 min at 12-W output power using 5-s-on/5-s-off intervals in 100 mM Tris-HCl (pH 7.5), 10 mM MgCl<sub>2</sub>, 5 mM dithiothreitol (DTT). Debris was removed by centrifugation at 16,000 × g for 10 min at 4°C. Protein concentration of the extracts was determined with a Bradford protein assay, using bovine serum albumin (BSA) as a standard. Secreted proteins were precipitated from the growth medium in chloroform-methanol as described previously (33). Protein pellets were dissolved in 1% (wt/vol) SDS, 50 mM Tris-HCl, pH 7.5, at 70°C for 10 min, and the amount of protein was measured using a protein DC assay (Bio-Rad, Hercules, CA), using BSA as a standard.

To perform quantitative proteomics, <sup>14</sup>N- and <sup>15</sup>N-labeled media or mycelial extracts were mixed 1:1 for protein content, and proteins were precipitated using chloroform-methanol (33). Protein pellets were heated at 95°C for 5 min in SDS-PAGE sample buffer, and proteins were separated on mini-Protean TGX anyKD gels (Bio-Rad). Gels were stained with Coomassie brilliant blue, and gel lanes were cut into approximately 15 pieces. Cysteines were reduced with DTT and modified with iodoacetamide, proteins were in-gel digested overnight with trypsin, and peptides were extracted, all as described previously (34). Acidified peptides were purified and concentrated using C<sub>18</sub> stage tips (35) as described previously. Peptides were analyzed on a Surveyor nanoLC system (Thermo, Waltham, MA) connected to an LTQ-Orbitrap mass spectrometer (Thermo) as described previously (36).

Peak lists were extracted and charge states determined using DTA-SuperCharge (version 2.0b1) (37). Spectra were searched against the *Streptomyces coelicolor* genome database (Sanger Institute, Hinxton, United Kingdom) using Mascot (version 2.2.03; Matrix Science, London, United Kingdom) with a parent ion tolerance of 10 ppm and a fragment ion mass tolerance of 0.9 Da. <sup>15</sup>N-metabolic labeling was specified as the quantification method. Fixed and variable modifications were the iodo-

acetamide derivative of cysteine and oxidation of methionine, respectively. Extracted ion currents (XIC) of identified peptides and their relative ratios were derived using MSQuant (version 2.0a81) (37), and a correction algorithm for incomplete  $^{15}\text{N}$  incorporation rates was applied (38). StatQuant (version 1.2.3) (39) was used for peptide ratio outlier removal based on Z-values ( $n > 3$ ), after which relative protein ratios were calculated by averaging peptide ratios. To compensate for errors in the mixing of samples, protein ratios were normalized based on the median ratio or, in case of the time series experiment, on the elongation factor G (SCO4661; *fusA*) ratio.

StatQuant was also used to determine whether the observed difference in the amount of protein between the two samples was significant using a one-sample Student's *t* test. Protein ratios with  $P < 0.1$  were accepted and compared to protein ratios of the label swap experiment using an additional one-sample Student's *t* test. Proteins that demonstrated opposing ratios between the original and label swap experiment ( $P > 0.5$ ) or were present in only one of the experiments were eliminated.

**MALDI-TOF-MS analysis.** For matrix-assisted laser desorption ionization–time-of-flight (MALDI-TOF)-MS, the same mycelial samples as those used for proteomics were also extracted with methanol, and the extract was mixed 1:1 (vol/vol) with a saturated  $\alpha$ -cyano-4-hydroxycinnamic acid solution in 50% (vol/vol) acetonitrile–0.05% (vol/vol) trifluoroacetic acid. One  $\mu\text{l}$  was spotted on a MALDI target plate, and samples were measured on a Bruker microflex LRF mass spectrometer in the positive-ion reflection mode using delayed extraction. For each spectrum, at least 1,000 shots were acquired at 60 Hz.

**RNA isolation and DNA microarray analysis and validation.** For transcript analysis, total RNA was isolated as described previously (40) from surface-grown mycelia of *S. coelicolor* M145 and its *rok7B7* null mutant, cultivated on MM agar plates with mannitol (1%, wt/vol) as the sole carbon source (plates were covered with cellophane discs). Two independent biological replicate experiments were performed. Samples were taken at 14 h (early vegetative growth), 24 h (vegetative growth), 30 h (early aerial growth), 36 h (aerial growth), 42 h (early sporulation), and 54 h (sporulation). Total RNA was purified using the Kirby-mix protocol (26). DNase I treatment was used to fully remove any traces of DNA. Before use, the RNA samples were checked for integrity on an Agilent 2100 Bioanalyzer (Agilent Technologies), and lack of DNA was checked by reverse transcription-PCR (RT-PCR) without reverse transcriptase. cDNA preparation, labeling, and hybridization was performed as previously described (41, 42). For the microarray analysis, 4 $\times$ 44K whole-genome *Streptomyces* arrays (Agilent Technologies) were used (42). The filtered data sets were analyzed using rank product analysis (43) via the Rank-Prod package in R (version 2.5.0) (44). Differentially expressed genes were identified as demonstrating a >2-fold change in expression between the parental strain M145 (wild type) and the *rok7B7* null mutant. To verify the output from the microarray analysis, semiquantitative RT-PCR analysis was carried out as described previously (45). For each RT-PCR, 200 ng of RNA was used (concentration was assessed using a Nanodrop spectrophotometer), and amplification without reverse transcriptase was used as the control. Oligonucleotides used for RT-PCR are presented in Table S3 in the supplemental material.

**Computer analysis.** Alignments of ROK-family proteins were made using ClustalW (46). This alignment was then used as the input for the Genebee website (<http://www.genebee.msu.ru/clustal/basic.html>) to produce a phylogram. Shaded protein alignments were visualized using the BoxShade program at [http://www.ch.embnet.org/software/BOX\\_form.html](http://www.ch.embnet.org/software/BOX_form.html).

## RESULTS

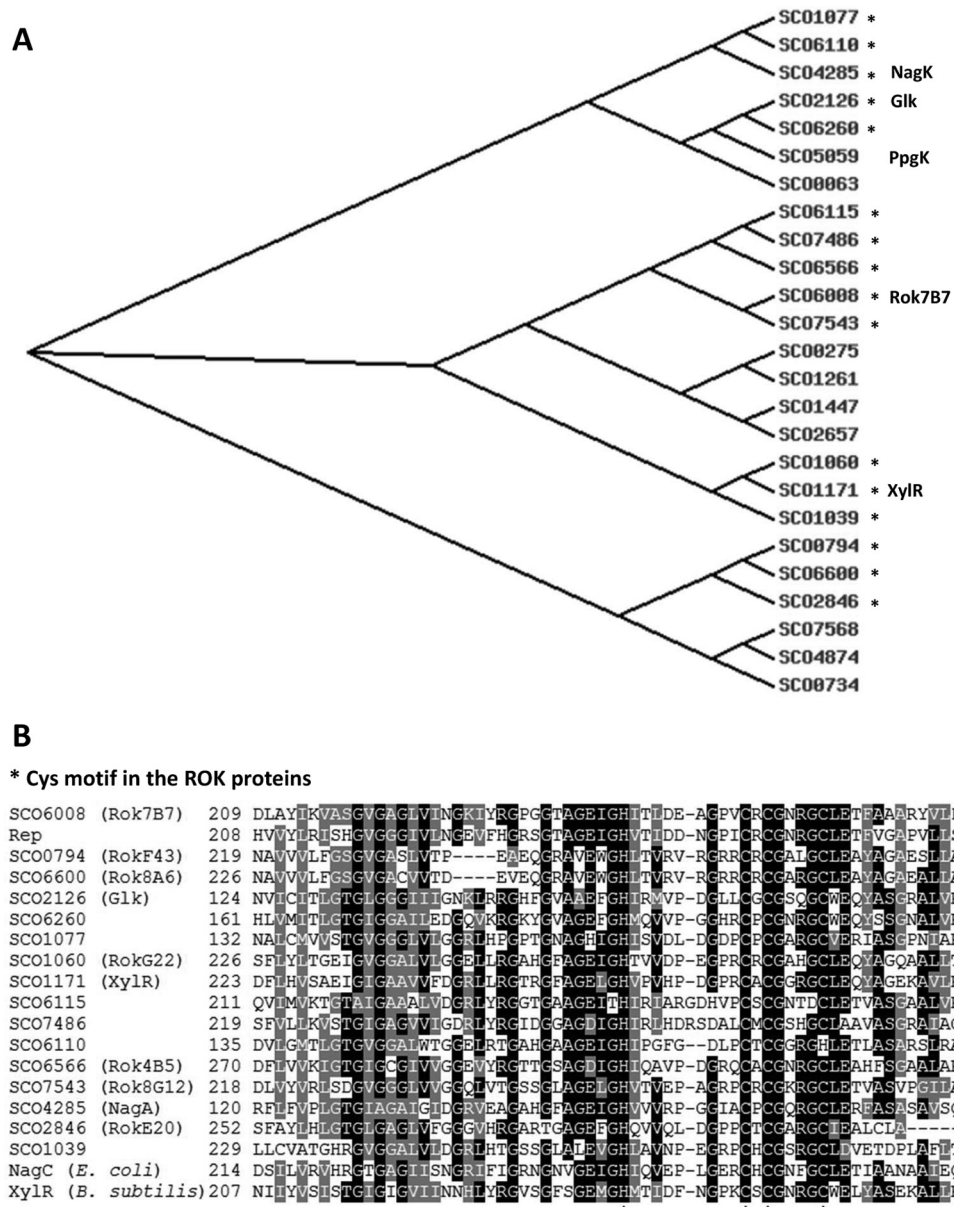
**The *rok*-family genes in *S. coelicolor*.** In total, 25 genes encoding proteins with a ROK-family signature are found on the *S. coelicolor* genome (2). A phylogram is shown in Fig. 1A. The family includes glucose kinase (Glk; SCO2126), which catalyzes the phosphorylation of glucose and plays a key role in carbon catab-

olite repression (47), and *N*-acetylglucosamine kinase NagK (SCO4285) (5). Sixteen of the ROK-family proteins contain a conserved zinc binding cysteine motif (Fig. 1B). Introduction of *rep*, an ROK-family regulatory gene obtained from a metagenomic library, accelerated sporulation and enhanced antibiotic production in *S. lividans* (22). BLASTP analysis of the ROK-family proteins of *S. coelicolor* proteins with the Rep protein sequence as the input revealed SCO6008 as the nearest orthologue in *S. coelicolor*, with 48% amino acid identity, while the next strongest hits show 30 to 35% amino acid identity (Table 1). However, the gene synteny of *rep* does not correspond to that of SCO6008 or to that of any of the other ROK genes. To analyze the possible role of the ROK-family proteins of *S. coelicolor*, we performed mutational analysis of seven *rok* genes (SCO0794, SCO1060, SCO2846, SCO6008, SCO6566, SCO6600, and SCO7543), whereby the predicted gene products, similar to the Rep protein, contain both a DNA binding and a zinc binding motif and have not been studied previously.

**Gene synteny for the *rok* genes.** The genetic context of the *rok* genes studied in this work is presented in Fig. S1 in the supplemental material. Suggestively, five out of seven genes are located in direct proximity of one or more ABC transporter systems. SCO6008, previously named *rok7B7* (23, 24, 48), is highly conserved in streptomycetes and is flanked by two sugar transport operons, namely, SCO6005 to SCO6007 (upstream) and the xylose transporter operon *xyIFGH* (SCO6009 to SCO6011; downstream), with its gene synteny highly conserved in streptomycetes. *Rok7B7* exhibits 27% amino acid identity to the xylose repressor XylR of *Bacillus subtilis*. *Rok7B7* itself shares 92% amino acid identity with NgcR of *Streptomyces olivaceoviridis*, which controls the upstream-located *N*-acetylglucosamine and chitobiose ABC transporter operon *ngcEFG* (49). However, in *S. coelicolor* *N*-acetylglucosamine is transported via the PTS, with NagE2 and NagF as the membrane components EIIC and EIIB, respectively (50), while chitobiose is transported via DasABC (51). Since SCO6005 to SCO6007 shows limited sequence similarity to NgcEFG (around 35% amino acid identity), it likely transports a molecule other than *N*-acetylglucosamine (5, 6).

Several of the other *rok* genes are also connected to ABC transporter operons. Like *rok7B7*, SCO6566 is connected to the utilization of C<sub>5</sub> sugars, as it is located upstream of a likely ribose transporter operon, with conserved gene synteny in many streptomycetes. In the direct environment of SCO1060 are two sugar transport operons, with SCO1062 to SCO1065 possibly encoding a galactose transporter based on the phylogenetic linkage to galactose utilization genes (*galK*, *galT*, and genes for  $\beta$ -galactosidases) and SCO1056 to SCO1055 having similarity to the maltose operon *malEFG* (SCO2231–2234) (52). SCO6600 is divergently transcribed from sugar transport operons SCO6601 to SCO6604, which are involved in glucoside transport (48). SCO7543 lies adjacent to the operons SCO7539 to SCO7542 and SCO7544 to SCO7548. SCO7547 to SCO7544 encode a likely sulfonate ABC transporter with similarity (37.5, 43.4, and 23.3% protein identity, respectively) to TauABC of *E. coli* (53).

**Phenotypic characterization of the *rok* mutants.** As a first step toward the characterization of the *rok* genes, null mutants were constructed in *S. coelicolor* M145. For this, we first replaced the entire coding region of each gene of interest with the apramycin cassette (*aacC4*) flanked by *loxP* sequences. For each experiment, five transformants exhibiting the desired double-



**FIG 1** Phylogeny of the predicted ROK family proteins of *S. coelicolor* M145 and alignment of Cys motifs. (A) Phylogram of the ROK proteins of *S. coelicolor* based on ClustalW analysis (46). ROK-family proteins containing a conserved zinc binding motif are marked by asterisks. The names of proteins along with known or suspected functions are given. (B) Alignment of the partial amino acid sequences of the ROK-family proteins that contain the zinc binding motif, with *B. subtilis* XylR and *E. coli* NagC included for reference purposes. Asterisks refer to the conserved Cys and His residues that coordinate the zinc atom. For shading, at least 50% of the aligned proteins should share the same or similar amino acid residues. Identical residues are shaded black, similar residues are shaded gray.

crossover phenotype (apramycin resistant and thiostrepton sensitive) were selected and verified by PCR as described in Materials and Methods. Following removal of the *aacC4* cassette by Cre recombinase, deletion mutants were obtained and designated GAM30 (M145ΔSCO0794), GAM31 (M145ΔSCO1060), GAM32 (M145ΔSCO2846), GAM33 (M145ΔSCO6008), GAM34 (M145ΔSCO6566), GAM35 (M145ΔSCO6600), and GAM36 (M145ΔSCO7543). For all experiments, both the apramycin-resistant and the deletion mutants were analyzed. Considering that there were no phenotypic differences between them in terms of antibiotic production and development, from this point

on we will only describe the results obtained for the deletion mutants unless stated otherwise.

The phenotype of all mutants was tested on MM (minimal medium) and R2YE (rich medium) agar plates supplemented with glycerol, mannitol, arabinose, ribose, xylose, fructose, glucose, galactose, mannose, cellobiose, lactose, maltose, sucrose, trehalose, glucosamine (GlcN), or *N*-acetylglucosamine (GlcNAc) as the sole carbon source, while taurine was tested as a sulfur source considering the possible relationship between SCO7543 and sulfur utilization. The mutants were phenotypically similar to the parental strain M145 in terms of development when grown on R2YE<sup>-</sup> or MM agar plates

TABLE 1 ROK regulators and their knockout strains studied in this work<sup>a</sup>

Gene	Name of the null mutant	Length of gene product (aa)	% Amino acid identity to:	
			NagC	Rep
SCO0794	GAM30	429	28	28
SCO1060	GAM31	413	27	35
SCO2846	GAM32	406	29	29
SCO6008 ( <i>rok7B7</i> )	GAM33	403	30	48
SCO6566	GAM34	584	28	33
SCO6600	GAM35	441	26	30
SCO7543	GAM36	424	27	34

<sup>a</sup> The amino acid identities of *S. coelicolor* A3(2) ROK-family proteins to NagC of *E. coli* and Rep, which was expressed from a metagenomic library, are presented. Protein lengths and amino acid identities are indicated.

supplemented with fructose, sucrose, trehalose, or taurine. The same was true for MM agar plates supplemented with mannitol or maltose and on R2YE<sup>-</sup> plates supplemented with GlcNAc.

GlcNAc activated antibiotic production of all strains under poor nutritional conditions (MM), while under rich growth conditions this process and aerial mycelium formation were blocked in all strains (Fig. 2A and B). This is in accordance with previous reports (8). Furthermore, cellobiose also inhibited development and antibiotic production on both MM- and R2YE-grown cultures, which may be due to the repressing effect of its monomer, glucose. Interestingly, actinorhodin production (blue pigment) and development was blocked for all strains grown on R2YE<sup>-</sup> agar plates with the C<sub>5</sub> carbon source ribose and also for several of the strains when grown on MM with ribose (Fig. 2A and B). Development of all strains was also inhibited on MM supplemented with arabinose, which is a C'-2 carbon epimer of ribose. Carbon source dependence of antibiotic production (such as repression by glucose) is a rather well-known phenomenon (3, 54), but the effect of C<sub>5</sub> sugars is far less well established.

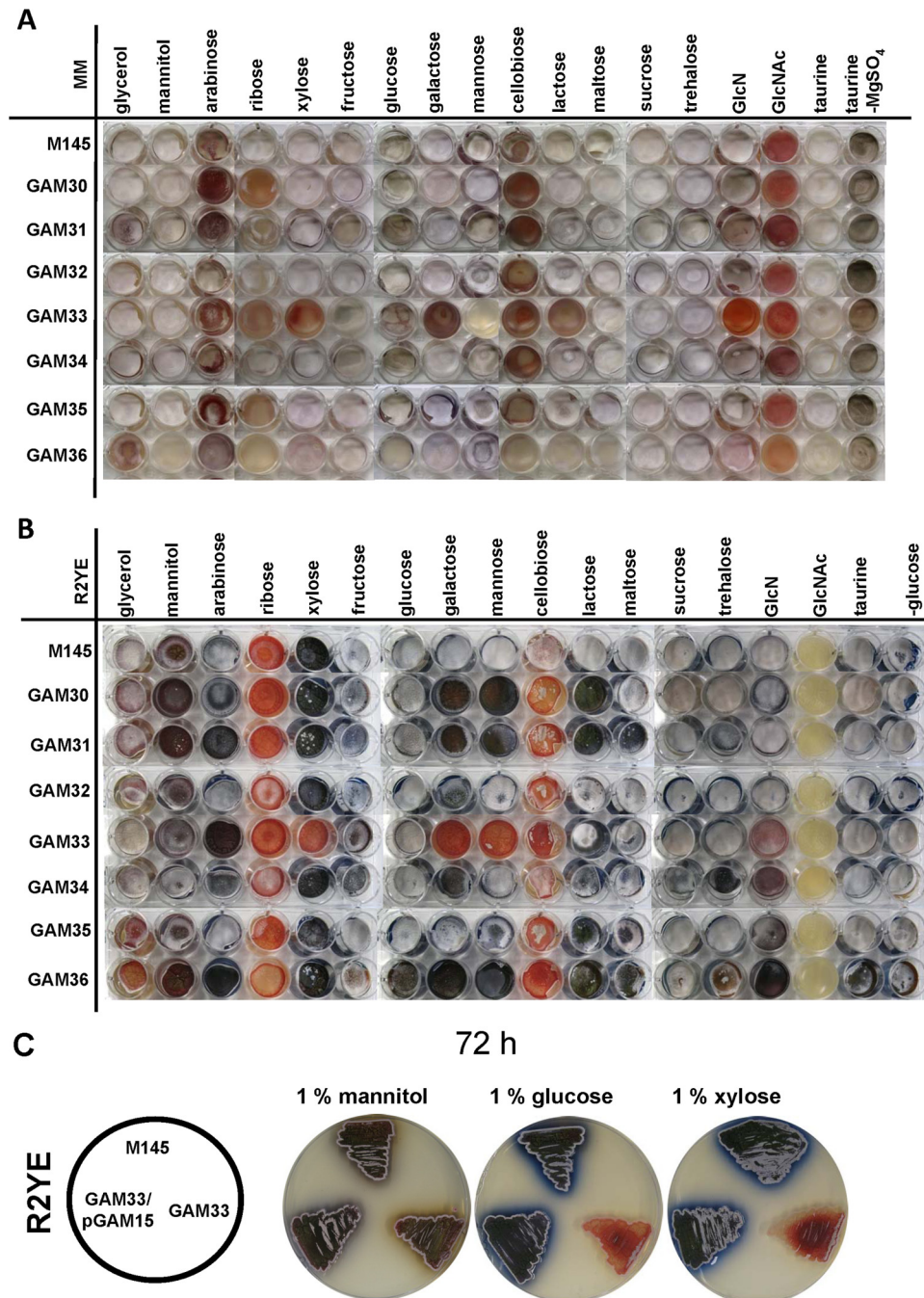
The most pleiotropic changes were observed in the *rok7B7* null mutant and, to a lesser extent, the ΔSCO7543 (GAM36) mutant (Fig. 2A and B). Growth on MM and R2YE agar medium supplemented with GlcNAc did not show significant differences between M145 and the *rok7B7* deletion mutant, suggesting that GlcNAc transport still takes place. The *rok7B7* null mutant showed delayed development and actinorhodin production on R2YE<sup>-</sup> with xylose, glucose, galactose, mannose, cellobiose, or GlcN (Fig. 2B and C). Furthermore, development of the *rok7B7* mutant was delayed on MM with arabinose and blocked on MM with ribose, xylose, galactose, mannose, cellobiose, lactose, or GlcN (Fig. 2A). Development of the ΔSCO7543 null mutant GAM36 was delayed under most growth conditions (Fig. 2A and B). The other five *rok* gene null mutants were similar to the parent under most growth conditions (Fig. 2A and B). Considering the pleiotropic effect of the deletion of *rok7B7*, we subjected its null mutant, GAM33, to more extensive analysis. In a control experiment, GAM33 was complemented with pGAM15 harboring *rok7B7* and its native promoter, which restored the wild-type phenotype (Fig. 2C).

**Rok7B7 controls xylose utilization and carbon catabolite repression.** Considering the *in silico* predictions and the possible involvement of Rok7B7 in control of xylose utilization, we monitored growth of the *rok7B7* mutant GAM33 in defined liquid MM with mannitol (1%, wt/vol), xylose (1%, wt/vol), or glucose (1%,

wt/vol) as the sole carbon source. The *rok7B7* mutant grew with a doubling time (~4 h) similar to that of the parent strain M145 in the presence of mannitol (Fig. 3A). Interestingly, in MM with glucose or xylose, GAM33 had a slightly higher growth rate than the parent M145 and reached significantly higher biomass levels due to postponed entry into the stationary phase (around 6 or 20 h later for glucose and xylose, respectively), suggesting enhanced glucose and xylose utilization (Fig. 3B and C). Indeed, complementation of GAM33 with plasmid pGAM15 harboring *rok7B7* and its native promoter restored the wild-type phenotype with reduced growth on glucose and xylose (Fig. 3B and C). This strongly suggests that the improved sugar utilization by GAM33 was due solely to the deletion of *rok7B7*. The prolonged vegetative growth phase corresponds well to the observed delay in development and antibiotic production on MM and R2YE agar plates supplemented with either glucose or xylose. The previously reported upregulation in glucose-grown cultures of the Rok7B7 protein in the *glkA* mutant, which is pleiotropically defective in CCR, prompted analysis of the role of Rok7B7 in carbon control (23). The secreted agarase produced by *S. coelicolor* is a classical CCR target and provides a simple assay (13). Therefore, agarase production by M145, its *rok7B7* null mutant GAM33, and the complemented mutant (harboring pGAM15) was compared on solid media containing mannitol, glucose, cellobiose, or xylose as the sole carbon source. The same amount of agarase was produced by all three strains on MM agar plates supplemented with mannitol. On MM agar plates with glucose or its dimer, cellobiose, neither strain produced significant amounts of agarase due to efficient glucose repression (Fig. 4 and Table 2). Interestingly, while the parental strain M145 produced normal levels of agarase on MM with xylose, agarase production was completely repressed in the *rok7B7* mutant (Fig. 4 and Table 2). Agarase production was restored following complementation of GAM33 with pGAM15 (Fig. 4), confirming that repression resulted solely from the inactivation of *rok7B7*. The *rok7B7* mutant also showed enhanced CCR on MM agar plates with maltose or glucosamine as the sole carbon source, although not as strong as that with xylose (Table 2). CCR was strongly enhanced in the *rok7B7* mutant grown on R2YE agar plates with various carbon sources (except glucose and cellobiose), with the complete block of agarase production on R2YE with GlcNAc. These data suggest that Rok7B7 acts as a global regulator of CCR.

**Global transcriptional profiling of the *rok7B7* null mutant by microarray analysis.** To obtain insight into the global transcriptional changes as a result of the deletion of *rok7B7* DNA microarray analysis was performed on RNA isolated from solid-grown MM mannitol cultures of *S. coelicolor* M145 and its *rok7B7* null mutant, GAM33. For this, plates were overlaid with cellophane discs and RNA was harvested after 14 h (early vegetative growth), 24 h (corresponding to vegetative growth), 30 h (vegetative/aerial growth), 36 h (aerial growth), 42 h (aerial growth/sporulation), and 54 h (sporulation). Under these conditions, which have been employed for a variety of microarray experiments with *S. coelicolor*, the mutant showed a timing of development similar to that of its parent.

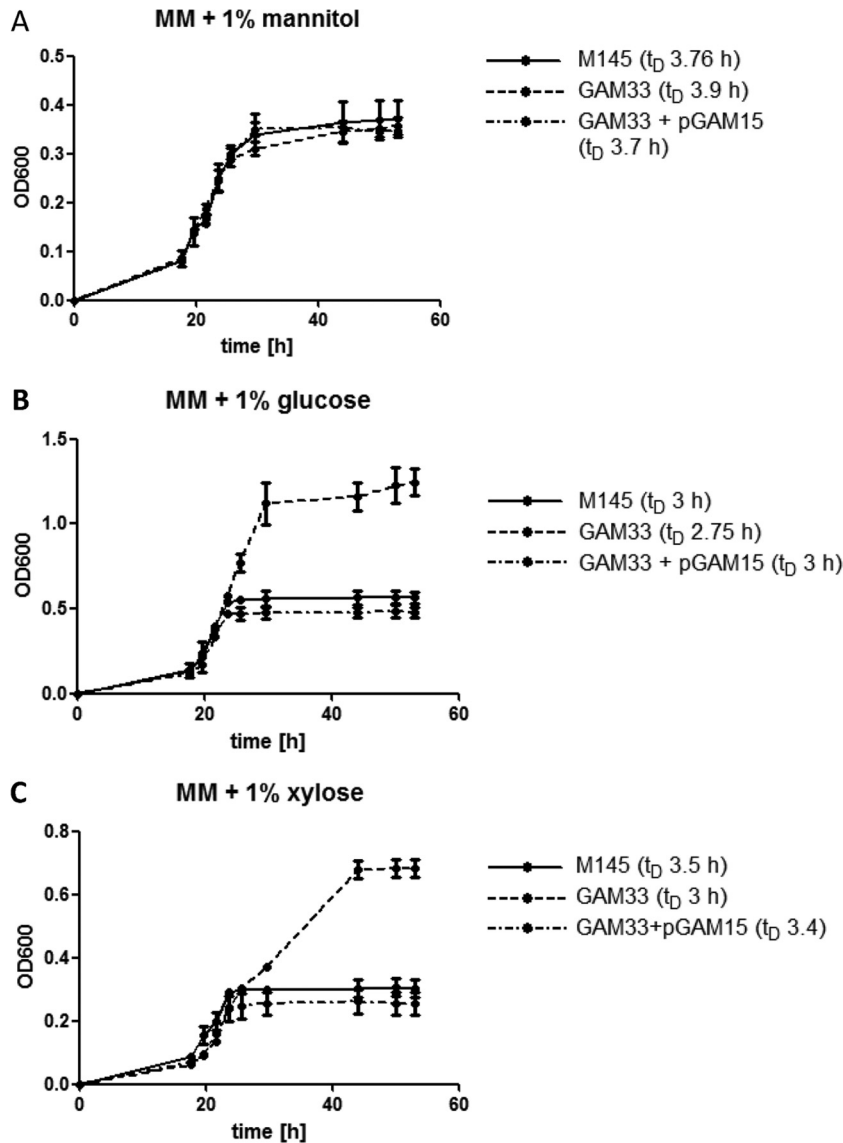
The transcription of over 160 genes was more than 2-fold up- or downregulated at one or more time points in the *rok7B7* mutant. These genes are listed in Table S4 in the supplemental material. Only a few targets showed over 2-fold differential expression between GAM33 and the wild-type strain at several time points,



**FIG 2** Phenotypic analysis of the *rok* mutants. Phenotypes of *S. coelicolor* M145 and its  $\Delta$ SCO0794,  $\Delta$ SCO1060,  $\Delta$ SCO2846,  $\Delta$ SCO6008 ( $\Delta$ *rok7B7*),  $\Delta$ SCO6566,  $\Delta$ SCO6600, and  $\Delta$ SCO7543 mutants (designated GAM30 to GAM36, respectively) were tested on MM (A) and R2YE (B) microtiter plates supplemented with 1% (wt/vol) carbon sources as indicated. Cultures were grown for 120 h. (C) Complementation of the *rok7B7* mutant GAM33 with pGAM15 harboring the *rok7B7* gene and its native promoter. Phenotypes were analyzed on R2YE agar plates supplemented with 1% (wt/vol) mannitol, glucose, or xylose. Cultures were grown for 72 h. Note that development of the *rok7B7* mutant is delayed on R2YE glucose and not completely blocked, and after 120 h normal sporulation occurred (see panel B).

including genes of the *xylFGH* operon (*xylF* and *xylG*) (Fig. 5A). Interestingly, the target that was most strongly downregulated in the *rok7B7* mutant was SCO2494, encoding pyruvate phosphate dikinase (PPDK; SCO2494), which generates PEP from pyruvate. This enzyme was recently implicated to play a key role in glucose kinase-independent carbon control in streptomycetes (23). In

support of the enhanced expression of *xylFGH* observed in the microarray data, quantitative proteomics experiments also showed that the components of the XylFGH transporter are about 10-fold increased at the protein level in the *rok7B7* mutant compared to the parental strain M145 (see below). Interestingly, transcription of *glcP* for the glucose permease GlcP (which occurs in



**FIG 3** Growth curves in minimal media with various carbon sources. The wild-type strain M145 (solid line), GAM33 (lacking *rok7B7*) (dashed line), and the complemented mutant GAM33/pGAM15 (dashed/dotted line) were grown in MM supplemented with mannitol (A), glucose (B), or xylose (C).  $t_D$ , doubling time. All growth analyses were performed in duplicate. Note that the mutant grows significantly better on glucose or xylose than the parental strain or the complemented mutant. The standard deviations for each data point are indicated. OD600, optical density at 600 nm.

two identical copies, *glcP1* and *glcP2*) was strongly upregulated at several time points (Fig. 5A). Most of the signal observed in the microarray experiments probably stems from *glcP1*, as *glcP2* is hardly expressed under any of the conditions tested, including during growth on glucose (11). The genes for glucose kinase and for the glycolytic enzymes were not changed significantly. Besides the primary metabolic genes discussed above, another class of genes that was deregulated in the *rok7B7* null mutant was that of the *chpABCDEFGHI*, *rdlAB*, and *ramS* genes, which encode the surfactant proteins chaplins, rodlins, and the spore-associated protein SapB, respectively (55, 56). All of these genes showed the same trend, but transcription of *chpDEH* and *rdlAB* was most strongly downregulated, namely, over 2-fold at 42 h compared to the parental strain M145 (see Table S4). Conversely, transcription was upregulated at 54 h.

In a control experiment, the microarray data were further validated by semiquantitative RT-PCR performed on the same RNA, using oligonucleotide pairs for *xylG* and *glcP1*, while *rpsI* was the control (Fig. 5B). The relative transcript levels of *xylG* and *glcP1* corresponded well to the microarray analysis, confirming their strong upregulation in the *rok7B7* null mutant.

**Global protein expression profiling of the *rok7B7* mutant by proteomics.** To further study the involvement of Rok7B7 in primary metabolism, carbon catabolite repression, and secondary metabolites biosynthesis, proteomics analysis was performed. For this, *S. coelicolor* M145 and its *rok7B7* null mutant, GAM33, were grown in liquid minimal media containing either  $^{14}\text{N}$  or  $^{15}\text{N}$  as the sole nitrogen source and mannitol as the sole carbon source until late logarithmic phase, when antibiotic production became apparent due to color changes of the growth medium.  $^{14}\text{N}$ - and  $^{15}\text{N}$ -

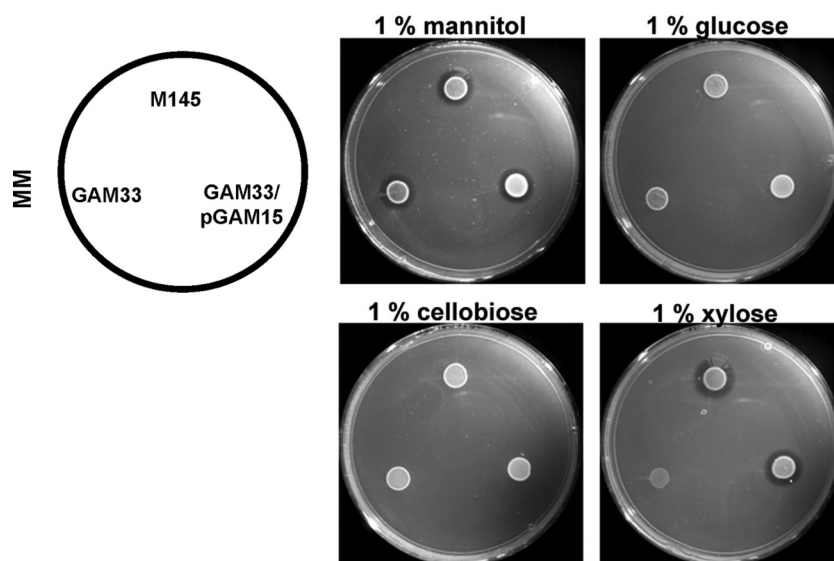


FIG 4 Effects of mannitol, glucose, cellobiose, and xylose on agarase production. Strains were M145 (parental strain), its *rok7B7* null mutant GAM33, and GAM33 complemented with pGAM15 (harboring the *rok7B7* gene). Five  $\mu$ l of spores of either strain ( $10^9$  CFU/ml) was spotted onto MM plates. After 3 days of growth, the agarase production was detected as a zone of clearing of the agar around the mycelial spot and was related to the biomass (for details, see Table 2 and reference 63).

TABLE 2 *Rok7B7* and carbon catabolite repression of agarase production

Agar plate and supplement	Value for strain <sup>a</sup> :						Fold change
	M145			$\Delta$ <i>rok7B7</i> (GAM33)			
	Agarase (mm)	Total (mm)	Ratio	Agarase (mm)	Total (mm)	Ratio	
<b>MM</b>							
Agar	6	26	0.23	6	26	0.23	1
Mannitol	5	25	0.2	5	25	0.2	1
GlcN	5	23	0.22	3	19	0.16	0.73
Fructose	6	26	0.23	6	25	0.24	1.04
Maltose	6	26	0.23	5	26	0.19	0.83
GlcNAc	5	26	0.19	5	26	0.19	1
Glucose	1	17	0.06	1	17	0.06	1
Xylose	5	25	0.2	0	15	0	0
Chitin	6	26	0.23	6	27	0.22	0.96
Cellobiose	1	17	0.06	1	17	0.06	1
<b>R2YE</b>							
Agar	5	22	0.23	1	15	0.07	0.29
Mannitol	5	22	0.23	1.5	16	0.09	0.41
GlcN	5	22	0.23	1.5	16	0.09	0.41
Fructose	5	23	0.22	2	17	0.12	0.54
Maltose	5	22	0.23	3	20	0.15	0.66
GlcNAc	2	17	0.12	0	14	0	0
Glucose	1	18	0.06	1	18	0.06	1
Xylose	5	25	0.2	2	17	0.12	0.59
Chitin	5	22	0.23	2	17	0.12	0.52
Cellobiose	1	18	0.06	1	18	0.06	1

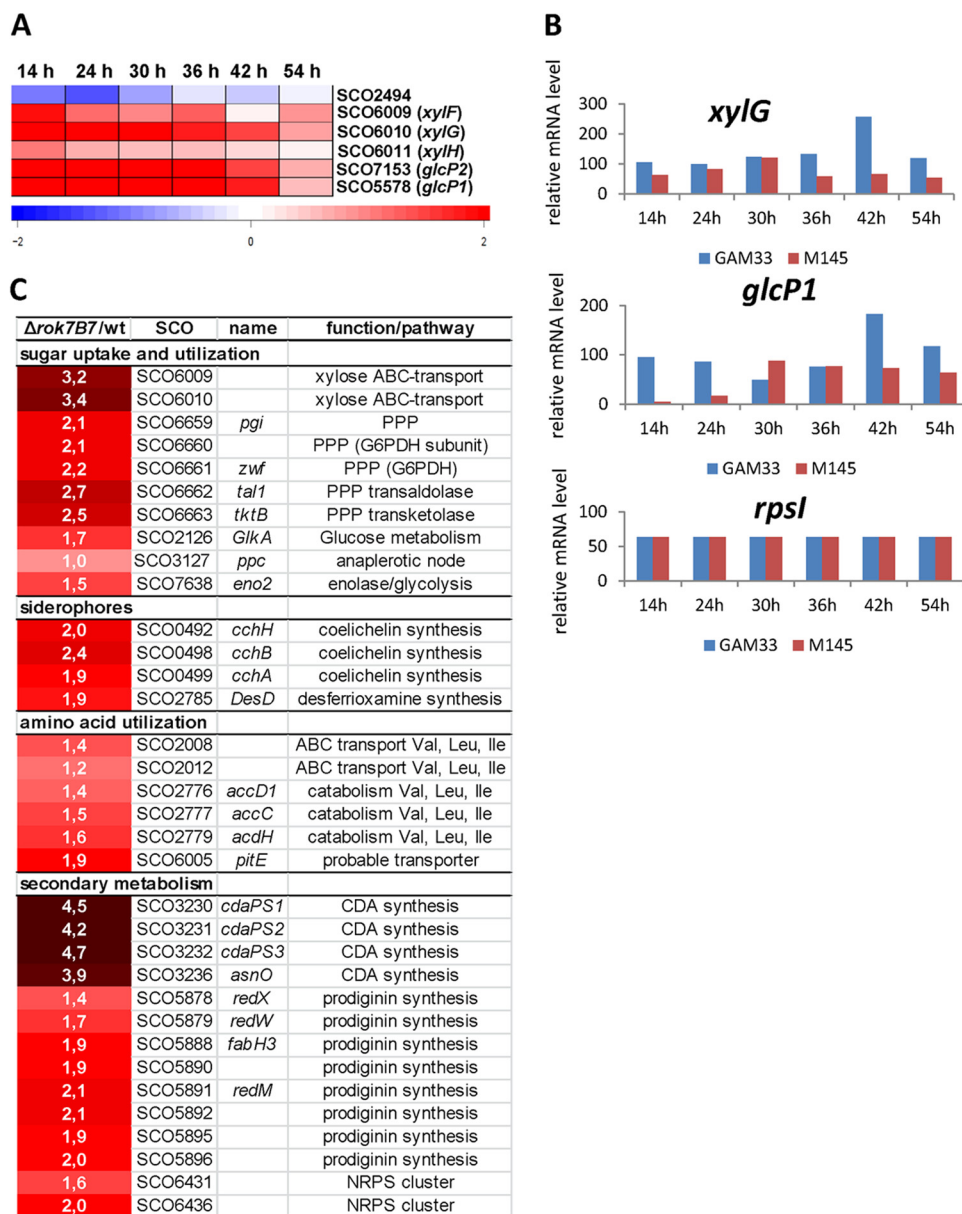
<sup>a</sup> Agarase indicates agarase zone, measured from the outside of the colony to the edge of the zone of clearing. Total indicates the combined length of the agarase zone and the colony. Ratio indicates the length of the agarase zone to the combined length of the agarase zone and the colony. Fold change indicates the ratio of the expression level of  $\Delta$ *rok7B7* to that of M145.

labeled protein extracts were obtained from both the mycelium itself and the spent media. These growth conditions are very similar to those used in the study of the *glkA* mutant (23), allowing direct comparison of the outcome.

Quantitative proteomics identified 59 proteins (53 proteins in the mycelial fraction and 7 in the secretome, with the xylose binding protein SCO6009 in both lists) whose expression changed with statistical significance in the *rok7B7* null mutant (Fig. 5C; also see Table S5 in the supplemental material). Classes of proteins that were overrepresented were those related to (i) primary metabolism (sugar and amino acid transport, metabolic enzymes), (ii) siderophore biosynthesis, and (iii) secondary metabolism (antibiotic biosynthesis) (Fig. 5C).

**Control of primary metabolism and CCR.** In line with the notion that *Rok7B7* controls the downstream-located xylose operon SCO6009 to SCO6011 (*xylFGH*), SCO6009 and SCO6010 were strongly upregulated (9- to 11-fold) in the *rok7B7* null mutant. Interestingly, glucose kinase (*Glk*; SCO2126) was 3-fold upregulated in the *rok7B7* null mutant, while phosphoenolpyruvate carboxylase (*PPC*; SCO3127) was 2-fold upregulated. These results are remarkable, since no glucose is present in the growth medium, and this again links *Rok7B7* to glucose utilization and carbon catabolite repression.

Enzymes of the pentose phosphate pathway (*PPP*; SCO6659 to SCO6663) were also significantly more abundant (4- to 6-fold) in the *rok7B7* mutant. Considering the chromosomal location, SCO6658 to SCO6663 is considered the secondary *PPP* cluster and may be involved in providing NADPH for secondary metabolite production (57). Abundance of proteins for glutamate import (SCO5775 to SCO5777) was 2- to 3-fold lower in the mutant, while those for import and catabolism of branched-chain amino acids (SCO2008 to SCO2012 and SCO2776 to SCO2779) and an enolase (SCO7638) were 2- to 3-fold enhanced in the mutant. Furthermore, the solute binding protein SCO6005, which is a dis-



**FIG 5** Transcriptomic and proteomic analysis of the *S. coelicolor* *rok7B7* null mutant. (A) Genes whose transcription was significantly differentially expressed between the *rok7B7* null mutant (GAM33) and its parent, *S. coelicolor* M145. RNA was isolated from *S. coelicolor* M145 or its *rok7B7* null mutant, GAM33, after 14 h (early vegetative growth), 24 h (vegetative growth), 30 h (vegetative/aerial growth), 36 h (aerial growth), 42 h (aerial growth/sporulation), and 54 h (sporulation) of growth on MM agar plates supplemented with 1% mannitol. The heat map indicates the fold changes. (B) Validation of microarray data by semiquantitative RT-PCR. RT-PCR was performed on RNA isolated from the parental strain M145 and the *rok7B7* null mutant, using oligonucleotide pairs for a selection of targets (*xylG* and *glcP1*). *rpsI* (for ribosomal protein S9) was used as a control. (C) Differentially expressed proteins between the *rok7B7* null mutant and the wild type. Quantitative proteomic comparison of the *rok7B7* deletion strain and parent strain (M145) is shown. Strains were grown in liquid MM supplemented with mannitol. The comparison between strains is indicated as ratios of the  $\Delta$ rok7B7 strain to M145. Results are given as  $\log_2$  ratios, therefore positive numbers indicate upregulation.

tant homologue of the NgcE protein that is involved in *N*-acetylglucosamine transport in *Streptomyces olivaceoviridis* (49), was 4-fold more abundant in the *rok7B7* mutant.

**Control of siderophore and antibiotic biosynthesis.** The *rok7B7* mutant shows strongly increased expression of biosynthetic proteins for the siderophores coelichelin (SCO0492, SCO0494, SCO0498, and SCO0499; 3- to 5-fold enhanced) and desferrioxamine (SCO2782 and SCO2785; 2- to 4-fold enhanced). These compounds bind extracellular iron, after which they can be

reimported through ABC transporters (58–60). A significant number of proteins that showed differential expression in the *rok7B7* mutant compared to M145 related to antibiotic production, namely, calcium-dependent antibiotic (CDA; SCO3230 to SCO3232 and SCO3236), prodiginine (SCO5878 to SCO5896, eight proteins detected), and an uncharacterized nonribosomal peptide (SCO6431 and SCO6436) all demonstrated increased expression levels in the *rok7B7* mutant. In particular, the biosynthetic proteins for CDA production were massively upregulated (Fig. 5C).

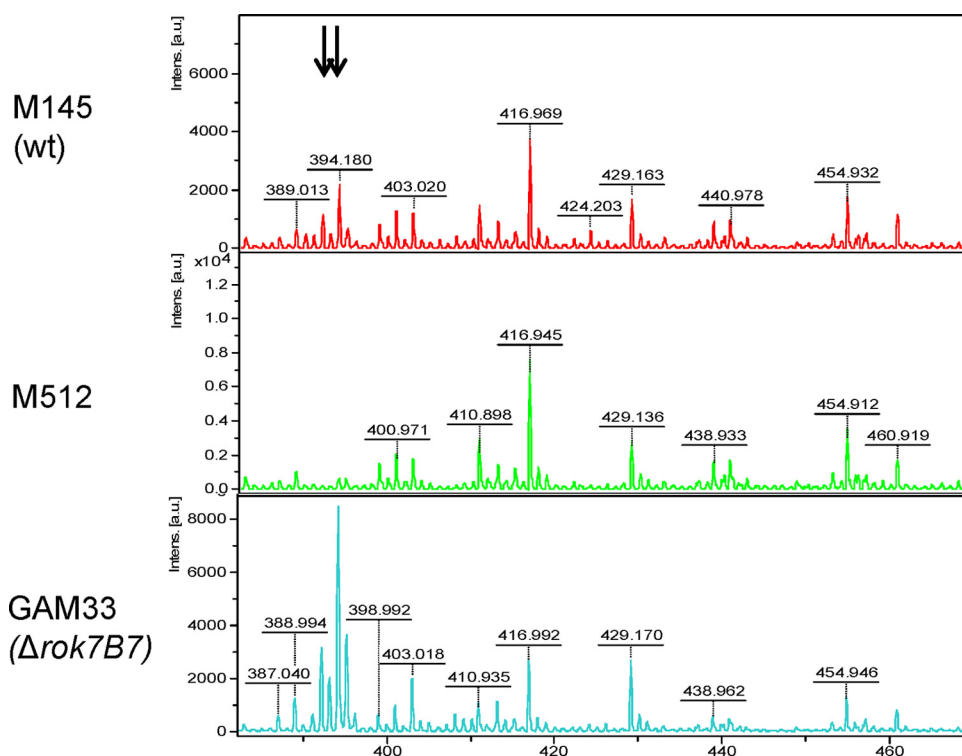


FIG 6 MALDI-TOF analysis of prodiginine production. Production of prodiginines was analyzed in mycelia of strains M145, GAM33, and M512. Arrows correspond to the 392- and 394-Da peaks for prodiginines. Horizontal axis,  $m/z$  (Da); vertical axis, intensity (arbitrary units [a.u.]).

To validate these data and assess the effect of the enhanced expression of antibiotic-related proteins to secondary metabolite production, we subjected mycelial samples corresponding to the samples used for proteomics to MALDI-TOF-MS analysis. *S. coelicolor* M512, which fails to produce prodiginines, was used as a negative control. Interestingly, prodiginines ( $m/z$  392 and 394) were readily detected in the *rok7B7* mutant and approximately 4-fold enhanced compared to the level in the parent M145 (Fig. 6). This validates the proteomics data, demonstrating that the deletion of *rok7B7* does result in upregulation of antibiotic production in minimal medium cultures.

## DISCUSSION

ROK-family proteins are widespread in actinomycetes, which are generally associated with the control of carbon metabolism. Except for glucose kinase (13, 23, 47, 61), study of this promising family of regulators and (sugar) kinases is relatively scarce. Analysis of a series of novel ROK null mutants identified *rok7B7* as a likely new pleiotropic regulatory gene in *S. coelicolor*. *rok7B7* null mutants showed a delay in aerial mycelium formation under most growth conditions as well as deregulation of antibiotic production, and they showed enhanced carbon catabolite repression. Deletion of the other *rok* family genes, SCO0794, SCO1060, SCO2846, SCO6566, SCO6600, and SCO7543, had less obvious phenotypic consequences, although the  $\Delta$ SCO7543 null mutant was somewhat affected in antibiotic production and aerial mycelium formation.

The *rok7B7* mutant showed strongly enhanced growth in liquid-grown cultures with glucose or xylose as the sole carbon source, while at the same time development and antibiotic pro-

duction were delayed in solid-grown cultures with the same carbon sources. This suggests that in the absence of *rok7B7*, the switch to development is postponed, as the onset of development and antibiotic production roughly correspond to the transition phase in liquid-grown cultures (62). This corresponds well to previous data showing that Rok7B7 likely activates actinorhodin biosynthesis (21). Furthermore, *rok7B7* is the nearest orthologue of *rep* isolated from a metagenomic library, the introduction of which into *S. lividans* accelerated both development and antibiotic production (22). The control of development by Rok7B7 is likely mediated at least in part via the control of carbon utilization, and it is upregulated during CCR in *S. coelicolor* (23). A model for the Rok7B7 regulon is presented in Fig. 7.

Combined transcriptomic and proteomics analyses suggested that Rok7B7 acts as a repressor of the adjacent xylose operon *xyIFGH* (SCO6009 to SCO6011), since expression of the operon was strongly enhanced in the mutant. Furthermore, in a model in which Rok7B7 acts as a repressor, one would expect that expression of the components of the xylose transporter are downregulated following enhanced expression of Rok7B7, but recent proteomics data revealed the opposite, namely, that Rok7B7 and XylFGH are both strongly induced in *glkA* deletion mutants (23). Such a direct correlation of expression is not consistent with a simple model of Rok7B7 acting as a specific repressor of *xyIFGH*. In fact, quantitative proteomics experiments identified XylFGH as the only sugar transport system that was strongly activated by glucose in a Glk-independent manner, along with its putative ROK-family regulator, Rok7B7. We propose that the upregulation of the xylose transporter operon is an attempt to compensate for the absence of Rok7B7, which would be consistent with a role

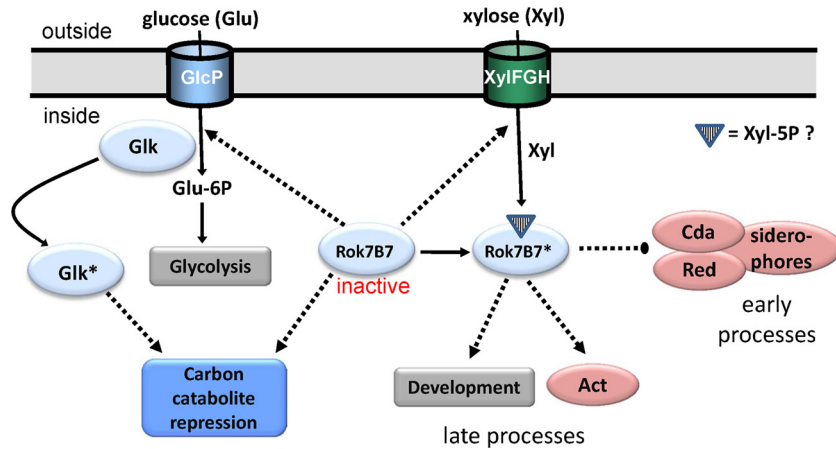


FIG 7 Model for the Rok7B7 regulatory network. Our data show that active Rok7B7 (indicated with an asterisk) activates actinorhodin (Act) production and development (as shown previously for its orthologue, Rep [22]), which explains the delayed development and Act production in the *rok7B7* null mutant. Conversely, the production of the secondary metabolites Red, CDA, and siderophores is upregulated. Deletion of *rok7B7* (presented in the model as “inactive Rok7B7”) results in enhanced expression of GlcP, Glk, and XylFGH as well as induction of CCR (including Glk-independent CCR). During glucose utilization, Glk is likely posttranslationally activated (shown as Glk\*), resulting in CCR (11). Solid lines, conversion or pathway; dotted lines, regulation (arrows represent activation, large dots indicate repression). We speculate that a metabolic derivative of xylose, such as xylose-5P, acts as a ligand for Rok7B7, thus activating the protein.

of xylose-5P or a related metabolite as an inducer molecule of Rok7B7.

In terms of the control of xylose utilization, the xylose regulator XylR of *S. coelicolor* is likely encoded by SCO1171, with the gene located adjacent to, and its gene product likely controlling the transcription of, *xylA* (SCO1169; for xylose isomerase) and *xylB* (SCO1170; for xylose kinase). XylA and XylB together catalyze the conversion from xylose to xylulose-5P. Since there is no phylogenetic linkage between the *xylABR* genes and *rok7B7* or *xylFGH* and deletion of *rok7B7* does not affect any of the genes among SCO1169 to SCO1171, it is less likely that these genes belong to the *rok7B7* regulon. However, to obtain full insight into the relationship between the regulons controlled by SCO1171-XylR and Rok7B7, the control of xylose and xylulose metabolism needs to be investigated in more detail. It should be noted, however, that deletion of *rok7B7* alone is sufficient to obtain strong CCR of agarase on xylose-containing MM agar plates, suggesting that xylose transport and utilization was completely derepressed in the *rok7B7* mutant. A similar observation was made for the control of maltose utilization by MalR in *S. coelicolor*, where *malR* deletion mutants showed complete catabolite control of agarase when maltose was used as the sole carbon source, while the wild-type strain hardly grows on maltose (63).

Interestingly, the combined data suggest that deletion of *rok7B7* induces glucose kinase-independent CCR. In other words, Rok7B7 may repress this poorly studied CCR pathway (23). The glucose transporter gene *glcP* was strongly upregulated in the *rok7B7* null mutant, suggesting direct control of glucose metabolism by Rok7B7. The expression of GlcP is strongly induced by glucose in a Glk-independent manner (23). The transmembrane GlcP binds to Glk, and its release during sugar transport may be a key mechanism in the activation of CCR (61). The target that was most strongly downregulated at the transcriptional level in the *rok7B7* null mutant was SCO2494, encoding pyruvate phosphate dikinase (PPDK; SCO2494), which generates PEP from pyruvate, the opposite of the reaction catalyzed by pyruvate kinase. PPDK also belongs to those proteins that are most strongly repressed by

glucose in a Glk-independent manner (23). In fact, PPDK is some 10- to 30-fold downregulated by glucose in a Glk-independent manner in both mannitol- and fructose-grown cultures, and it may play a key role in Glk-independent carbon control in streptomycetes. The change in expression of GlcP (upregulated) and SCO2494 (downregulated) by glucose in a Glk-independent manner is consistent with cells exerting glucose repression, as was indeed observed for the *rok7B7* mutant. This is further corroborated by the enhanced expression of Glk in *rok7B7* null mutants, as shown by the proteomics experiments. The fact that no significant change was found in transcription of the gene for glucose kinase (*glkA*) is consistent with the constitutive transcription of *glkA* and confirms the earlier concept that glucose kinase expression is controlled at the posttranscriptional level (61). All in all, our data suggest that Rok7B7 negatively controls CCR. However, despite our efforts using purified Rok7B7, we failed to detect direct binding of Rok7B7 to the promoter regions of any of the genes whose expression was most significantly affected by the deletion of *rok7B7* (data not illustrated). We therefore performed global *in vivo* DNA binding studies using chromatin immunoprecipitation (ChIP)-chip technology on cultures grown under standard conditions of minimal media agar plates with mannitol as the sole carbon source, but this also failed to identify significant binding to specific sites. Therefore, we expect that a ligand or cofactor is required to facilitate binding of Rok7B7 to the DNA, with derivatives of xylose, cellobiose, or glucose as candidate molecules. ChIP-chip experiments under different growth conditions should provide more insight into the direct Rok7B7 regulon and its binding site.

In terms of antibiotic production, both the red-pigmented prodiginines and their biosynthetic proteins were enhanced in the *rok7B7* mutant in liquid-grown MM cultures supplemented with mannitol, as shown by proteomics and subsequent MALDI-TOF-MS analysis of secondary metabolites. Conversely, in solid-grown cultures on rich media (R2YE), the production of the blue-pigmented antibiotic actinorhodin was strongly delayed in the *rok7B7* mutant, which was independent of the carbon source and

consistent with earlier reports that Rok7B7 activates actinorhodin production (21, 22). Following a DNA-affinity capture assay (DACA), Rok7B7 was isolated as one of the proteins that was bound to the promoter regions of *actII-ORF4* and *redD*, the pathway-specific transcriptional regulators of the *act* and *red* gene clusters, respectively (21). All data point to control of antibiotic production by Rok7B7, with activation of Act and repression of Red and CDA. The proteomics experiments also showed upregulation of the degradation pathway for branched-chain amino acids (leucine, isoleucine, and valine), which are precursors for biosynthesis of polyketides such as actinorhodin (64), suggesting that besides antibiotic production, Rok7B7 also controls precursor supply.

While the C<sub>5</sub> sugar xylose had a stimulatory effect on actinorhodin production in wild-type cells, perhaps acting via Rok7B7, the C<sub>5</sub> sugars arabinose and ribose, which are closely related compounds, suppress actinorhodin production. To the best of our knowledge, the stimulatory effect of xylose on antibiotic production was not reported previously, but the inhibitory effect of the C<sub>5</sub> sugars ribose and arabinose on antibiotic production was established in *Streptomyces hygroscopicus* (65) and *Streptomyces olivocinereus* (66). Actinorhodin production is strongly delayed in the *rok7B7* mutant when grown on media containing xylose. As argued above, xylose or a derivative thereof (e.g., xylose-5P) may act as an effector molecule of Rok7B7, leading to its activation and linking nutrient sensing to antibiotic production and development. Antibiotic production was also repressed by cellobiose. Carbon source dependence of antibiotic production is widely reported in the literature (reviewed in references 3 and 54). Since cellobiose is a dimer of glucose (β-1,4), its repressing activity regarding antibiotic production may be explained by the accumulation of glucose produced from its extracellular hydrolysis, thus effecting glucose repression.

In conclusion, our work underlines the complexity of the regulatory networks involved in the control of antibiotic production and highlights Rok7B7 as a novel pleiotropic transcriptional regulator of CCR that links the control of primary and secondary metabolism in *S. coelicolor* (Fig. 7). Rok7B7 therefore qualifies as a possible target for genetic engineering approaches toward the improvement of antibiotic production.

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