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Systems-wide analysis of the ROK-family regulatory gene *rokL6* and its role in the control of glucosamine toxicity in *Streptomyces coelicolor*

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ABSTRACT Streptomycetes are saprophytic bacteria that grow on complex polysaccharides, such as cellulose, starch, chitin, and chitosan. For the monomeric building blocks glucose, maltose, and *N*-acetylglucosamine, the metabolic pathways are well-documented, but that of glucosamine (GlcN) is largely unknown. *Streptomyces nagB* mutants, which lack glucosamine-6-phosphate deaminase activity, fail to grow in the presence of high concentrations of GlcN. Here, we report that mutations in the gene for the ROK-family transcriptional regulator RokL6 relieve the toxicity of GlcN in *nagB* mutants, as a result of elevated expression of the major facilitator superfamily (MFS) exporter SCO1448. Systems-wide analysis using RNA sequencing, ChIP-Seq, EMSAs, 5'RACE, bioinformatics, and genetics revealed that RokL6 is an autoregulator that represses the transcription of *sco1448* by binding to overlapping promoters in the *rokL6-sco1448* intergenic region. RokL6-independent expression of *sco1448* fully relieved the toxicity of GlcN to *nagB* mutants. Taken together, our data show a novel system of RokL6 as a regulator that controls the expression of the MFS transporter SCO1448, which in turn protects cells against GlcN toxicity, most likely by exporting toxic metabolites out of the cell.

IMPORTANCE Central metabolism plays a key role in the control of growth and antibiotic production in streptomycetes. Specifically, aminosugars act as signaling molecules that affect development and antibiotic production, via metabolic interference with the global repressor DasR. While aminosugar metabolism directly connects to other major metabolic routes such as glycolysis and cell wall synthesis, several important aspects of their metabolism are yet unresolved. Accumulation of *N*-acetylglucosamine 6-phosphate or glucosamine 6-phosphate is lethal to many bacteria, a yet unresolved phenomenon referred to as “aminosugar sensitivity.” We made use of this concept by selecting for suppressors in genes related to glucosamine toxicity in *nagB* mutants, which showed that the gene pair of *rok*-family regulatory gene *rokL6* and major facilitator superfamily transporter gene *sco1448* forms a cryptic rescue mechanism. Inactivation of *rokL6* resulted in the expression of *sco1448*, which then prevents the toxicity of amino sugar-derived metabolites in *Streptomyces*. The systems biology of RokL6 and its transcriptional control of *sco1448* shed new light on aminosugar metabolism in streptomycetes and on the response of bacteria to aminosugar toxicity.

KEYWORDS ROK-family protein, aminosugar metabolism, systems biology, *Streptomyces* biology, control of antibiotic production

Streptomycetes are Gram-positive bacteria with a mycelial lifestyle that reproduce via sporulation. Their large GC-rich genomes encode a plethora of specialized metabolites such as antibiotics, anticancer drugs, and many other industrially and medically

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relevant compounds (1–3). The production of these molecules is closely related to the transition from vegetative to aerial growth during the development of the colonies (4, 5). Streptomycetes grow by tip extension and branching of vegetative hyphae, which are divided into multigenomic compartments. Under adverse conditions such as nutrient depletion, a complex developmental program is initiated, which results in the formation of an aerial mycelium, and eventually the aerial hyphae differentiate to form chains of unigenomic spores (3, 6). At the onset of development, the vegetative mycelium is partially degraded via programmed cell death so as to provide the building blocks necessary for aerial growth in an otherwise nutrient-depleted environment (7, 8).

During cell wall recycling, the constituents of the peptidoglycan (PG), namely *N*-acetylglucosamine (GlcNAc) and *N*-acetylmuramic acid (MurNAc) that make up the PG strands and the cross-linking amino acids, are re-imported into the cell. The lactyl ether substituent of the intracellular MurNAc 6-phosphate (MurNAc-6P) is cleaved by MurNAc-6P etherase, yielding *N*-acetylglucosamine 6-phosphate (GlcNAc-6P) (9). GlcNAc is thereby internalized by the phosphoenolpyruvate-dependent phosphotransferase system (PTS), which simultaneously phosphorylates GlcNAc to GlcNAc-6P (10, 11). The next step is GlcNAc-6P deacetylation by *N*-acetylglucosamine 6-phosphate deacetylase NagA forming glucosamine-6-phosphate (GlcN-6P), which is a central molecule at the intersection of multiple metabolic pathways, including glycolysis via conversion to fructose-6-phosphate (Fru-6P) by glucosamine-6-phosphate deaminase NagB (12). GlcNAc plays a critical role in signaling during the control of development and antibiotic production in streptomycetes (13, 14). DasR controls aminosugar transport and metabolism and is also a highly pleiotropic repressor of antibiotic production in *Streptomyces* (15–17). Metabolic control of DasR is a key step in the early activation of development and specialized metabolism under nutrient-deprived conditions.

While the metabolic pathway of GlcNAc has been well-characterized, little is known of glucosamine (GlcN) metabolism in streptomycetes. The hydrolysis of chitosan and the metabolism of chitosan-derived oligomers (GlcN)₂₋₃ are under the control of CsnR (18, 19), a repressor from the ROK [repressors, open reading frames (ORFs), and kinases] family of transcriptional regulators (20). GlcN oligomers are imported via the ABC-transporter complex CsnEFG-MsiK and are then presumably hydrolyzed and phosphorylated by CsnH, a sugar hydrolase, and CsnK, a ROK-family kinase, respectively (19).

High concentrations of either GlcN or GlcNAc are toxic to *Streptomyces coelicolor* in the absence of the enzyme glucosamine-6-phosphate deaminase (NagB) (12). Similar aminosugar sensitivity was also observed for *Escherichia coli* (21, 22). Despite extensive studies focusing on the metabolism of aminosugars, the underlying cause of aminosugar toxicity remains unresolved. Interestingly, when grown in the presence of high concentrations of either GlcN or GlcNAc, *S. coelicolor* Δ nagB strains sustain spontaneous second-site mutations that allow the colonies to survive. We previously exploited this principle to identify novel genes related to GlcN transport and metabolism (12, 23). Spontaneous mutations or deletion of *nagA*, preventing the conversion from GlcNAc-6P to GlcN-6P, relieves the toxicity of both GlcNAc and GlcN to *S. coelicolor* *nagB* mutants (23). Since NagA is not known to play a role in GlcN metabolism, this points to a caveat in our understanding of aminosugar metabolism. In addition to suppressor mutations in *nagA*, two independent suppressor mutations were found in the gene for ROK-family regulator SCO1447 (RokL6), which relieved toxicity specifically of GlcN but not of GlcNAc. ROK-family of proteins often play important roles in the (control of) sugar utilization in bacteria, and these regulators are widespread in streptomycetes (20, 24, 25).

Here, we report on the function of RokL6 by a systems-wide approach, using mutational and transcriptional analysis and *in vitro* and *in vivo* DNA binding studies. This revealed that the gene product of *rokL6* is an ROK-family regulator that specifically represses the adjacent gene *sco1448*. This gene encodes a putative major facilitator superfamily (MFS) transporter, and we propose that it acts as a pump for toxic substances created during the challenge of *nagB* mutants with higher concentrations of GlcN.

MATERIALS AND METHODS

Strains, plasmid, and growth conditions

The bacterial strains and plasmids used or constructed in this study are summarized and listed in Table 1, and all oligonucleotides are described in Table 2. *Escherichia coli* was grown and transformed according to the standard procedures (26), with *E. coli* JM109 serving as the host for routine cloning, and *E. coli* ET12567 (27) used for the isolation of non-methylated DNA for transformation into *Streptomyces*. For protein heterologous expression, *E. coli* BL21(DE3) from Novagen was used. *E. coli* was grown in Luria-Bertani media in the presence of selective antibiotics as required, with the following final concentrations: ampicillin (100 µg/mL), apramycin (50 µg/mL), kanamycin (50 µg/mL), and chloramphenicol (25 µg/mL). *Streptomyces coelicolor* M145 (28) was obtained from the John Innes Centre strain collection and was the parent of all mutants. *S. coelicolor* *nagB* mutant Δ *nagB* (23) and GlcN-derived *nagB* suppressor mutant SMG1 (29) have been described previously. All *Streptomyces* media and routine techniques are based on the *Streptomyces* manual (30). Phenotypic characterization of *Streptomyces* mutants was carried out on minimal medium (MM) agar plates with different carbon sources as indicated. Soy flour mannitol (SFM) agar plates were used for the preparation of spore suspensions and *E. coli* to *S. coelicolor* conjugations. A mixture of 1:1 yeast-extract malt extract and tryptic soy broth liquid media was used to cultivate mycelia for protoplast preparation and genome DNA isolation. Growth curves of *Streptomyces* were performed in liquid cultures containing minimal medium normal minimal media phosphate (NMMP) (30) with 1% (wt/vol) glucose or GlcN as the sole carbon source, and the dry weights of the mycelia were measured at different time points.

Gene knock-out, complementation, and overexpression

As a basis for the creation of gene deletion mutants, we used the unstable multi-copy plasmid pWHM3 (31, 35), as described previously (12). For the *rokL6* knock-out construct, a 1,254-bp 5' flanking region and a 1,366-bp 3' flanking region were amplified by PCR from the *S. coelicolor* M145 genome, using the primer pairs described in Table 2. The upstream region was cloned as an *EcoRI*-*XbaI* fragment, and the downstream region as an *XbaI*-*BamHI* fragment, and these two fragments were ligated into pWHM3 from the *EcoRI* and *BamHI* sites. The apramycin resistance cassette *aac* (3)/IV, flanked by *loxP* sites (*apra-loxP*), was subsequently cloned into the engineered *XbaI* sites between the flanks to create *rokL6* knock-out construct pKO-*rokL6*. In the same way, the flanking regions of *sco1448* were cloned into pWHM3 with the *apra-loxP* to produce the *sco1448* knock-out vector, pKO-1448. The knock-out constructs were introduced into *S. coelicolor* M145 or its *nagB* mutant. For clean gene knock-out mutants, the apramycin resistance cassette was excised by the introduction of pUWLcre, which expresses Cre recombinase (32, 36). The correct recombination event in each of the knock-out mutants was confirmed by PCR.

For *rokL6* complementation, the entire coding region of *rokL6* with its own promoter was amplified from the *S. coelicolor* chromosome. The PCR product was digested with *XbaI*/*EcoRV* and then inserted into pSET152 (33) to obtain *rokL6*-complemented vector pCOM-*rokL6*. This construct was introduced into *nagB-rokL6* double mutant Δ *nagB* Δ *rokL6*, and the successfully complemented strains Δ *nagB* Δ *rokL6*-C were selected by apramycin. The empty pSET152 was conjugated in Δ *nagB* Δ *rokL6* to produce Δ *nagB* Δ *rokL6*-E, which was used as a control strain.

For SCO1448 overexpression, a 1,212-bp DNA fragment containing *sco1448* was amplified and ligated into pSET152 with the 63-bp highly efficient promoter P30, namely the promoter SP30 with the 20-bp ribosomal binding site RBS15 (37), to produce SCO1448-overexpressing vector pOE-1448. The pOE-1448 construct was then introduced into Δ *nagB* to obtain the SCO1448 overexpression strain Δ *nagB*SCO1448-O.

TABLE 1 Bacterial strains and plasmids used in this study

Bacterial strains	Description ^{a,b}	Reference or source
<i>S. coelicolor</i>		
M145	<i>S. coelicolor</i> M145, SCP1 [−] SCP2 [−] prototroph	(30)
$\Delta nagB$	M145 <i>nagB</i> ^d	(12)
SMG1	$\Delta nagB$ suppressor mutant	(29)
$\Delta rokL6$	M145 <i>rokL6</i> ^d	This study
$\Delta nagB\Delta rokL6$	M145 <i>nagB</i> ^d <i>rokL6</i> ^d	This study
$\Delta nagB\Delta rokL6$ -C	$\Delta nagB\Delta rokL6$ complemented with <i>rokL6</i>	This study
$\Delta nagB\Delta rokL6$ -E	$\Delta nagB\Delta rokL6$ containing empty pSET152	This study
$\Delta nagB$ -E	$\Delta nagB$ containing empty pSET152	This study
$\Delta nagB$ SCO1448-O	$\Delta nagB$ overexpressing SCO1448	This study
M145- <i>rokL6</i> -FLAG	M145 containing <i>rokL6</i> fused to 3× FLAG tag sequence	This study
$\Delta nagB\Delta rokL6$ -FLAG	$\Delta nagB\Delta rokL6$ complemented with <i>rokL6</i> containing a 3× FLAG tag sequence	This study
$\Delta sco1448$	M145 <i>sco1448</i> ^d	This study
$\Delta nagB\Delta sco1448$	$\Delta nagB$ <i>sco1448</i> ^d	This study
SMG1 $\Delta sco1448$	SMG1 <i>sco1448::aac(3)/IV</i>	This study
$\Delta nagB\Delta rokL6\Delta sco1448$	$\Delta nagB$ <i>rokL6</i> ^d <i>sco1448::aac(3)/IV</i>	This study
M145- <i>ProkL6</i> -eGFP	M145 with eGFP transcribed from <i>ProkL6</i>	This study
M145-P1448-eGFP	M145 with eGFP transcribed from P1448	This study
M145- <i>PnagKA</i> -eGFP	M145 with eGFP transcribed from <i>PnagKA</i>	This study
M145-P30-eGFP	M145 with eGFP transcribed from P30	This study
$\Delta rokL6$ - <i>ProkL6</i> -eGFP	$\Delta rokL6$ with eGFP transcribed from <i>ProkL6</i>	This study
$\Delta rokL6$ -P1448-eGFP	$\Delta rokL6$ with eGFP transcribed from P1448	This study
$\Delta rokL6$ - <i>PnagKA</i> -eGFP	$\Delta rokL6$ with eGFP transcribed from <i>PnagKA</i>	This study
$\Delta rokL6$ -P30-eGFP	$\Delta rokL6$ with eGFP transcribed from P30	This study
M145 $\Delta sco0136$ -0137	M145 <i>sco0136</i> -0137:: <i>aac(3)/IV</i>	This study
$\Delta nagB\Delta sco0136$ -0137	$\Delta nagB$ <i>sco0136</i> -0137:: <i>aac(3)/IV</i>	This study
<i>E. coli</i>		
<i>E. coli</i> JM109	<i>E. coli</i> strain for routine cloning	(26)
<i>E. coli</i> ET12567/pUZ8002	Strain used for conjugation between <i>E. coli</i> and <i>Streptomyces</i>	(27)
<i>E. coli</i> BL21(DE3)	Strain used for protein expression	Novagen
Plasmids	Description	Reference or source
pWHM3	<i>E. coli</i> / <i>Streptomyces</i> shuttle vector, high copy number and unstable in <i>Streptomyces</i>	(31)
pUWLcre	<i>E. coli</i> / <i>Streptomyces</i> shuttle vector expressing the Cre recombinase in <i>Streptomyces</i>	(32)
pSET152	Integrative <i>E. coli</i> / <i>Streptomyces</i> shuttle vector	(33)
pET28a (+)	Vector for His ₆ -tagged protein expression	Novagen
pCRISPomyces-2	<i>E. coli</i> / <i>Streptomyces</i> shuttle vector, harboring codon optimized cas9, designed for easy inserted spacer of specific genes	(34)
pKO- <i>rokL6</i>	Construct for the deletion of <i>rokL6</i>	This study
pKO-1448	Construct for the deletion of <i>sco1448</i>	This study
pCOM- <i>rokL6</i>	<i>rokL6</i> complementation vector based on pSET152	This study
pOE-1448	<i>sco1448</i> expression vector based on pSET152	This study
pKI-FLAG ₃	3× FLAG knock-in vector based on pCRISPomyces-2	This study
pCOM-FLAG ₃	<i>RokL6</i> -3 × FLAG complementation vector	This study
pEX- <i>RokL6</i>	<i>RokL6</i> -His ₆ expression vector based on pET28a (+)	This study
pEGFP- <i>rokL6</i>	Construct expressing eGFP from <i>ProkL6</i>	This study
pEGFP-1448	Construct expressing eGFP from <i>P</i> <i>sco1448</i>	This study
pEGFP- <i>nagKA</i>	Construct expressing eGFP from <i>P</i> <i>nagKA</i>	This study
pEGFP-P30	Construct expressing eGFP from P30	This study

^a"d" indicates the gene before "d" is in-frame deleted.^b"::*aac(3)/IV*" indicates the gene before "::" is replaced by *aac(3)/IV* cassette.

TABLE 2 Oligonucleotides used in this study

Name	5'-3' sequence	Function or descriptions
Cloning used		
rokl6-KO-LF	GTCAGAATTACCCCTCGGAACACCACCAGGCA	rokl6 knock out
rokl6-KO-LR	GAAGTTATCCATCACCTCTAGACATCGCGGATCCTTCCAGAT	rokl6 knock out
rokl6-KO-RF	GAAGTTATCGGCATCTCTAGATTGCGACCGCGGAGCGGTAG	rokl6 knock out
rokl6-KO-RR	GTCAAAGCTTGCATGCGCAGGCGGTCAAGC	rokl6 knock out
rokl6-COM-F	CTAGGATATCCATCGCACGTCCGACA	RokL6 complementation
rokl6-COM-R	CATGCTAGAGGTGAGGCCCTTCCGGG	RokL6 complementation
rokl6-KO-checkF	TGCTGCTGCCGACGGTACTC	rokl6 mutant check
rokl6-KO-checkR	CTATCAGGAGCCTGCGCTGATAG	rokl6 mutant check
sco1448-KO-LF	GTCAAGATTCTGCTGCCGACGGTACTCGGGTGG	sco1448 knock out
sco1448-KO-LR	GAAGTTATCCATCACCTCTAGATGTGTTTCATGTCCACCCCTC	sco1448 knock out
sco1448-KO-RF	GAAGTTATCGGCATCTTAGACCGGCAGTCTGAACGCTCGC	sco1448 knock out
sco1448-KO-RR	GTCAAAGCTTCTCCGCGATCAGGGCGATGACG	sco1448 knock out
sco1448-KO-checkF	CGGGCATGCCGGATCCTTC	sco1448 mutant check
sco1448-KO-checkR	GGGTGCTGGTCCGGCTGGAC	sco1448 mutant check
sco1448-OE-F	CTAGGATATCTGTTACATTGCAACCGTCTCTCTTTGACATGCTGTGGCGCTTGGGTGTAAAGTCGTGGCCATCTAAGTAAGGAGTGTCCATATGAACACAGGG	sco1448 overexpression
	ACCTACGGG	
sco1448-OE-R	GTGCACTCTAGAGTGGACTCAC	SCO1448 overexpression
rokl6-spacer-F	ACGCGCGCGCGGGCTACCGCTC	rokl6 spacer assembly
rokl6-spacer-R	AAACGAGCGGTAGCCCGCGGGCC	rokl6 spacer assembly
rokl6-flag-LF	CAGCTATGACCATGATTACGCTCGCCCTTTCATCTGCTCCAGC	rokl6 with 3 × FLAG PCR
rokl6-flag-LR	GGCGCGGGGC TACTTGTGTGTCGTCTTGTAGTCGATGTCGTGTCCTTGTAGTCGCGCTCGTGGTCTTGTAGTCCCGTCCGGCGGTGCGAAG	rokl6 with 3 × FLAG PCR
rokl6-flag-RF	GACAAGTAGCCGCGCGCGCCGGAAGGGCCTCACCCGG	rokl6 CRISPR template assembly
rokl6-flag-RR	GTTGTAAACGACGCGCCAGTGCCACGCGCGCGGGGAC	rokl6 CRISPR template assembly
RokL6-flag-checkF	CGGGCTACTTGTCTCGT	RokL6-FLAG check
RokL6-flag-checkR	TCGACCTGCTGCTGGTG	RokL6-FLAG check
RokL6-EXP-F	GAATAATTTTGTTTAACTTTAAGAAGGAGATACATGCCGATCACCGAGCAC	RokL6-His ₆ expression
RokL6-EXP-R	GACAGCAATGGGTGCGGATCATGCCCCGCATCACCGAGCAC	RokL6-His ₆ expression
egfp-F	ATGGTGAGCAAGGGCGAGGAG	eGFP gene PCR
egfp-R	CTTGGGCTGCAGGTGACTTTACTTTGTACAGTCGTCTCATGC	eGFP gene PCR
P1447-egfp-F	GCTATGACATGATTACGAATTCGATGAGCGCCAGCCCACTGGC	rokl6 promoter PCR
P1447-egfp-R	TCGCCCTTGC TCACCATGCGGGATCTTCAGATCGG	rokl6 promoter PCR
P1448-egfp-F	GCTATGACATGATTACGAATTCGATGCGGTGAGGCGTTTCGAGAG	sco1448 promoter PCR
P1448-egfp-R	TCGCCCTTGC TCACCATGTTCCACCCCTCCGTGTCG	sco1448 promoter PCR
Phag/K4-egfp-F	TATGACATGATTACGAATTCGATCGCGAGACCGCGTCATGC	nagK4 promoter PCR
Phag/K4-egfp-R	CTCGCCCTTGTCTACCATCCCGTGCGCCACATCGAG	nagK4 promoter PCR
SP30R15-egfp-F	GCTATGACATGATTACGAATTCGATGTTACATTCGAACCGTCTC	SP30R15 PCR
SP30R15-egfp-R	TCGCCCTTGC TCACCATGAGACCTCTTACTTAGATGG	SP30R15 PCR

(Continued on next page)

TABLE 2 Oligonucleotides used in this study (Continued)

Name	5'-3' sequence	Function or descriptions
sco0136-0137-KO-LF	CAGCTATGACCATGATTACTCGTGTGACGGTGAAGTTTCAC	sco0136-0137 knock out
sco0136-0137-KO-LR	ACGAAGTATTCGCGCATCTTTCGCTCATACAGTGAAC	sco0136-0137 knock out
sco0136-0137-KO-RF	GCTATACGAAGTTATCATCATCACTCTCTCCCTGAGCGCGCTTGAC	sco0136-0137 knock out
sco0136-0137-KO-RR	GTGTGAAAAACGACGGCCAGTGTGACGGACCGATCATGGACC	sco0136-0137 knock out
sco0136-37-KO-checkF	GCTGGAACCGACCGCTTAC	sco0136-0137 mutant check
sco0136-37-KO-checkR	CGGCTGTCCGCCGAGATTCC	sco0136-0137 mutant check
qPCR used		
<i>hrdB</i> -qPCR-F	GCACATGGTTCGAGGTCATCA	<i>hrdB</i> qPCR
<i>hrdB</i> -qPCR-R	GGTCATGTCGAGCTCTTGG	<i>hrdB</i> qPCR
<i>egfp</i> -qPCR-F	CCTGTGACCACTCTGACCTAC	eGFP gene qPCR
<i>egfp</i> -qPCR-R	TTGCCGTGCTCTTGAAGAAGATG	eGFP gene qPCR
sco1448-qPCR-F	CCAACCTGCTCACCTGTG	sco1448 qPCR
sco1448-qPCR-R	TCCAGACGGTCTCGATCTC	sco1448 qPCR
5'RACE used		
Oligo(dT) anchor primer	GACCAACGGTATCGATGTCGACTTTTTTTTTTTTTTTT	5'RACE anchor primer
Anchor primer	GACCAACGGTATCGATGTCGAC	5'RACE anchor primer
<i>rokl6</i> -GSP	GTGATGGGACGGGACG	<i>rokl6</i> specific primer
<i>rokl6</i> -NP1	ACGCTTTCGGTGGCGACGTC	<i>rokl6</i> nested primer1
<i>rokl6</i> -NP2	GCTTCAACTGCCCTGCCGTC	<i>rokl6</i> nested primer2
sco1448-GSP	GCGAGGCACAGGTGAGCAG	sco1448 specific primer
sco1448-NP1	GCGGCATGGCGAGGGACG	sco1448 nested primer1
sco1448-NP2	CTCACGGCGGTGCTCTG	sco1448 nested primer2
EMSA used		
EMSA- <i>rokl6</i> -P-F	CGCACCGGTAATCCACTATCTATCAGGAGGCTCCCTGATAGTTTACGCCGGGATTGAC	RokL6 BS probe annealing
EMSA- <i>rokl6</i> -P-R	GTCAATCCGGCGGTAATACTATCAGGAGCGCTGCCTGATAGATGGATTACCGGTGCG	RokL6 BS probe annealing
EMSA- <i>hrdB</i> -P-F	CGGCCGCCCGCACCGTTCGGCCCATTCACAGCGGTGGTCCCTGTCCGCCGTGGA	RokL6 EMSA <i>hrdB</i> -P annealing
EMSA- <i>hrdB</i> -P-R	TCCACGGCGGACAGGGCGCACCGGCTTGGGAATGGCCGACGGTGGCGGGGGCCG	RokL6 EMSA <i>hrdB</i> -P annealing
EMSA- <i>rokl6</i> -MP1-F	CGCACCGGTAATCCACTATCTATCAGGAGCGCTCCCTGATAGTTTACGCCGGGATTGAC	RokL6 EMSA M1 probe annealing
EMSA- <i>rokl6</i> -MP1-R	GTCAATCCGGCGGTAATACTATCAGGAGCGCTGCCTGATAGTGGATTACCGGTGCG	RokL6 EMSA M1 probe annealing
EMSA- <i>rokl6</i> -MP2-F	CGCACCGGTAATCCACTATCTATCAGGAGCGCTCCGAAATTCGTTACGCCGGGATTGAC	RokL6 EMSA M2 probe annealing
EMSA- <i>rokl6</i> -MP2-R	GTCAATCCGGCGGTAATACTATCAGGAGCGCTGCCTGATAGTGGATTACCGGTGCG	RokL6 EMSA M2 probe annealing
EMSA- <i>rokl6</i> -MP3-F	CGCACCGGTAATCCACTATCTATCAGGAGCGCTCCGATCCGTTTACGCCGGGATTGAC	RokL6 EMSA M3 probe annealing
EMSA- <i>rokl6</i> -MP3-R	GTCAATCCGGCGGTAATACTATCAGGAGCGCTGCCTGATAGTGGATTACCGGTGCG	RokL6 EMSA M3 probe annealing

FLAG₃ tag knock-in by CRISPR

To express FLAG-tagged RokL6 in *S. coelicolor*, the 3× FLAG sequence (FLAG₃) was fused to the end of the original copy of *rokL6* on the genome using codon-optimized CRISPR-Cas9 system (34) as described previously (38). Briefly, the spacer sequence (5'- GGCG CCGCGGGCTACCGCTC-3') specific to *rokL6*, located at the end of the coding region, was inserted into the pCRISPOmyces-2 plasmid from *BbsI* sites. Next, the 2,263-bp template containing FLAG₃ for homology-directed repair was made and inserted into the spacer-containing plasmid, resulting in a *rokL6*- FLAG₃ knock-in construct designated as pKI-FLAG₃. Mutagenesis was done according to the previous study (38). After conjugation of pKI-FLAG₃ to *S. coelicolor* A3(2) M145, ex-conjugants were patched on SFM agar plates with 20 µg/mL apramycin. Then, positive ex-conjugants were patched on antibiotic-free SFM agar plates and grown at 37°C. Spores were collected and checked for loss of construct. Apramycin-sensitive strains were selected for spore collection, and their genomes were checked for desired recombination events. The successful *in situ* FLAG₃ knock-in strain was identified as M145-*rokL6*-FLAG. Additionally, *rokL6* with FLAG₃ was ligated into pSET152 to generate pCOM-FLAG₃ and introduced into the *nagB-rokL6* double mutant for evaluating RokL6-FLAG₃ function *in vivo*.

Heterologous expression and purification of His₆-tagged RokL6 protein

For the heterologous expression of *S. coelicolor* RokL6 in *E. coli*, the 1,197-bp *rokL6* coding region was amplified from *S. coelicolor* genomic DNA using primer pair RokL6-exp-F/RokL6-exp-R. The PCR fragment was ligated into pET-28a (+) from *XhoI* and *NcoI* sites, generating expression vector pEX-RokL6. The RokL6 expression vector was transformed into *E. coli* BL21(DE3), and the expression of C-terminal His₆-tagged RokL6 recombinant protein, RokL6-His₆, was induced by the addition of isopropyl β-D-1-thiogalactopyranoside at a final concentration of 1 mM when the cell density was reached around an optical density at 600 nm of 0.6, followed by overnight incubation at 16°C. Cells were harvested, washed, and disrupted in lysis buffer (39) by sonication. Soluble RokL6-His₆ was purified from the supernatant using HisPur Cobalt Resin (Thermo Fisher Scientific, USA) and dialyzed against electrophoretic mobility shift assay (EMSA binding buffer).

Chromatin immunoprecipitation sequencing of RokL6

Chromatin immunoprecipitation sequencing (ChIP-Seq) experiments were carried out essentially as described previously (38). In brief, *S. coelicolor* M145-*rokL6*-FLAG was grown on MM agar covered with cellophane disks, using mannitol with and without 50 mM GlcN as the carbon source. Mycelia were collected after 24 h (vegetative growth) and 48 h (sporulation). Mycelia were treated with phosphate-buffered saline (PBS) buffer containing 1% formaldehyde for 20 min to cross-link the DNA and protein. After thorough washing in PBS, mycelia were resuspended in lysis buffer [10 mM Tris-HCl pH 8.0, 50 mM NaCl, 15 mg/mL lysozyme, 1× protease inhibitor (Roche, Bavaria, Germany)] and incubated at 37°C for 20 min. After incubation, 0.5 mL IP buffer (100 mM Tris-HCl pH 8.0, 250 mM NaCl, 0.8%, vol/vol Triton-X-100) was added to the mycelia samples, and chromosomal DNA was sheared to 100–500 bp fragments using the Bioruptor Plus water bath sonication system (Diagenode, Liège, Belgium). The lysates were incubated with 40 µL Anti-FLAG M2 affinity gel (cat A2220, Sigma-Aldrich, St. Louis, USA) and incubated at 4°C overnight. After centrifugation, the pellet and untreated total extracts (control) were incubated in 100 µL of IP elution buffer (50 mM Tris-HCl pH 7.5, 10 mM EDTA, 1%, [wt/vol] SDS) at 65°C overnight to reverse the cross-link. The beads were removed by centrifugation before DNA isolation by phenol-chloroform. The extracted DNA samples were then further purified with the DNA Clean & Concentrator kit (Zymo Research, CA, USA). The enriched DNA samples were sent to Novogene Europe (Cambridge, UK) for library construction and next-generation sequencing. ChIP-Seq data analysis was performed as described previously (38).

RNA isolation, RNA sequencing, and quantitative PCR

Spores (10^7 CFU) of *S. coelicolor* M145 and the *rokL6* deletion mutant were grown on MM agar plates overlaid with cellophane discs, with either 1% (wt/vol) mannitol or 1% mannitol + 50 mM GlcN as the carbon sources. Biomass from two time points (24 h on mannitol or 26 h on mannitol with GlcN for vegetative growth phase, VEG; and 42 h on mannitol or 44 h on mannitol with GlcN for sporulation phase, SPO) was collected and snap-frozen in liquid N_2 . All samples were assessed as biological triplicates. After breaking the mycelia using a TissueLyser II (Qiagen, Venlo, The Netherlands), the RNA was extracted using a modified Kirby mix (30). The transcriptome sequencing library preparation and sequencing were outsourced to Novogen Europe (Cambridge, UK). Removal of rRNA from the samples was carried out using NEBNext Ultra directional RNA Library Prep Kit (NEB, MA, USA). Sequencing libraries were generated using the NEBNext Ultra RNA Library Prep Kit for Illumina (NEB), and sequencing was performed on an Illumina NovaSeq 6000 platform. Raw data were cleaned using fastp v0.12.2 (40), and then mapped to the *S. coelicolor* M145 genome sequence (GenBank accession [AL645882.2](#)) using bowtie2 v2.4.4 (41). Read counts for each gene were generated by featureCounts v2.0.1 (42). Values for transcripts per million were generated using a custom Python script. Differentially expressed genes and log2Foldchange were determined using DESeq2 v1.32.0 (43) with the data shrinkage function “apeglm” (44).

For quantitative PCR (qPCR), cDNA was synthesized using the iScript cDNA Synthesis Kit (Biorad, CA, USA). Briefly, qPCR was performed using the iTaq Universal SYBR green qPCR Kit (Biorad, CA, USA), and the program used was set as follows: 95°C for 30 s; 40 cycles of 95°C for 10 s, 60°C for 30 s, plate reading; melting curve from 65°C to 95°C with 5 s per 0.5°C increment. The principal RNA polymerase σ factor encoding gene, *hrdB* (sco5820), was used as the internal control, and the qPCR data were analyzed by CFX Manager software (version 3.1, Biorad, CA, USA), using $\Delta\Delta C_q$ standard, which is an implementation of the method described in reference (45).

Electrophoretic mobility shift assays

Electrophoretic mobility shift assays (EMSAs) were performed as described previously (38). Double-stranded DNA probes (60 bp) were generated by gradually cooling reverse complemented single-strand oligonucleotides in 30 mM HEPES, pH 7.8, heating to 95°C for 5 min, then ramping to 4°C at a rate of 0.1°C/s. The *in vitro* DNA-protein binding assays were performed in EMSA binding buffer (20 mM HEPES pH 7.6, 30 mM KCl, 10 mM $(NH_4)_2SO_4$, 1 mM EDTA, 1 mM DTT, 0.2% Tween20). The binding reactions (10 μ L), including 1 picomole DNA probe, 20 ng/ μ L bovine serum albumin (BSA), and various concentrations of RokL6-His₆, were incubated at 30°C for 20 min. The reactions were then loaded on 5% non-denatured polyacrylamide gels and separated by electrophoresis. The gels were briefly stained with ethidium bromide and imaged using the Gel Doc imaging system (BioRad, CA, USA).

Determination of transcriptional start sites

Transcriptional start sites (TSS) of *rokL6* and *sco1448* were determined by 5' RACE using a 5'/3' RACE Kit 2nd generation (Roche, CA, USA). Total RNA (2 μ g) extracted from 48-h cultures of *S. coelicolor* grown on solid minimal medium containing 1% mannitol was used for reverse transcription with gene-specific primers. The obtained cDNA was purified, and an oligo(dA) tail was added to the 3' end by terminal transferase, followed by PCR amplification of the tailed cDNA with oligo(dT) anchor primer and gene-specific nested primers. Using the resulting PCR product (diluted 1,000-fold) as a template, an additional round of PCR was performed with a more inner nested primer and an anchor primer provided in the kit to produce a single specific DNA band. The final PCR product was purified and sent for sequencing, and the TSS was determined as the first nucleotide following oligo(dA).

Promoter activity test

The promoter regions of *rokl6*, *sco1448*, *nagKA* (12), and P30 (37) were ligated with a gene expressing enhanced green fluorescent protein (eGFP) and integrated into pSET152 via the *XbaI*/*EcoRV* sites, generating four recombinant vectors: pEGFP-*rokl6*, pEGFP-1448, pEGFP-*nagKA*, and pEGFP-P30. The recombinant constructs were then introduced into *S. coelicolor* and *rokl6* deletion mutant $\Delta rokl6$ to generate the following strains: M145-*Prokl6*-eGFP, M145-P1448-eGFP, M145-P*nagKA*-eGFP, M145-P30-eGFP, and $\Delta rokl6$ -*Prokl6*-eGFP, $\Delta rokl6$ -P1448-eGFP, $\Delta rokl6$ -P*nagKA*-eGFP, and $\Delta rokl6$ -P30-eGFP. The *in vivo* activities of all the promoters were evaluated by comparing the transcription level of the gene for eGFP via confocal microscopy and confirmed by qPCR.

Confocal imaging

Sterile coverslips were inserted into MM with 1% mannitol agar plates at an angle of 45°, and spores of eGFP-harboring strains were inoculated at the intersection angle and incubated at 30°C for 48 h. The mycelium was scraped off from coverslips into a drop of water and then imaged with a TCS SP8 confocal microscope (Leica) using a 63× oil immersion objective (NA: 1.40) (46). Image processing and the analysis of fluorescent intensity were performed using ImageJ (version 1.54d).

Bioinformatics analysis

DNA and protein database searches were performed using the BLAST server of the National Centre for Biotechnology Information (<http://www.ncbi.nlm.nih.gov>) and the *S. coelicolor* genome page services (<http://strepdb.streptomyces.org.uk>). Polypeptide sequences were aligned using Clustalw available at <http://www.ebi.ac.uk/clustalw>. Gene synteny analysis was performed using SynTax (47). To visualize the consensus sequence for the predicted binding site of RokL6, WebLogo (48) was used, and regulon predictions were done using PREDetector (49). The comparative analysis of the intergenic regions of *rokl6*/*sco1448* and their orthologous pairs was performed by MEME (50).

RESULTS

RokL6 is involved in GlcN metabolism

We previously showed that null mutants of *S. coelicolor nagB* are sensitive to high concentrations of GlcN, whereby spontaneous second-site mutations arise that allow the colonies to survive. Exploitation of this principle led to the identification of suppressor mutations in *nagA* and in a novel gene related to GlcN transport and metabolism (12, 23). Two independent suppressor mutants were obtained in the gene *sco1447* when *S. coelicolor* $\Delta nagB$ was grown on GlcN, namely SMG1 and SMG38, and these mutations specifically alleviated the toxicity exclusively on GlcN, but not GlcNAc (23, 51). Suppressor mutant SMG1 had sustained a single nucleotide insertion at nucleotide (nt) position 26 within the coding region of *sco1447*, while SMG38 had a single nucleotide deletion at nt position 120 in *sco1447*. In both cases, the mutations resulted in a frameshift at the beginning of the gene, thereby preventing the expression of the active protein. *sco1447* encodes a ROK-family transcriptional regulator, which we designated RokL6, based on the terminology that ROK-family proteins are named after the specific cosmid they are located on (24) within the ordered cosmid library that was used for the *S. coelicolor* genome sequencing project (52).

The observation that the mutations specifically alleviate the toxicity of GlcN to *nagB* null mutants, but not that of GlcNAc, suggests that RokL6 plays a specific role in the control of GlcN metabolism. To test this, the *rokl6* single mutant $\Delta rokl6$ and *nagB-rokl6* double mutant $\Delta nagB\Delta rokl6$ were generated by deleting *rokl6* in *S. coelicolor* M145 and in the previously published *nagB* null mutant (29), respectively. To create the mutants, the entire coding region of *rokl6* (nt positions + 3 to +1,200) was replaced with the apramycin-resistance cassette *aac* (3)/IV, which was flanked by *loxP* sites. The *aac* (3)/IV

gene was subsequently excised from the genome using the Cre recombinase expressed from plasmid pUWLcre. The resulting *rokL6* single mutant and *nagB-rokL6* double mutant, Δ *nagB* Δ *rokL6*, were confirmed by PCR. In order to ascertain that the phenotypes were specifically caused by the deletion of *rokL6*, we genetically complemented the Δ *nagB* Δ *rokL6* mutant by expressing *rokL6*. For this, the $-478/+1,226$ region of *rokL6*, harboring the entire gene and its promoter region, was amplified from the *S. coelicolor* chromosome and cloned into integrative vector pSET152 to obtain pCOM-*rokL6* (see Materials and Methods for details).

As expected, *S. coelicolor* M145 and its *rokL6* single mutant Δ *rokL6*, which has an intact copy of *nagB*, grew well on all media, while *nagB* mutants were unable to grow on MM with mannitol (1%, wt/vol) and 5 mM GlcN or GlcNAc. The *rokL6-nagB* double mutant grew well on MM with 1% mannitol and 5 mM GlcN, but failed to grow when GlcNAc was used instead of GlcN (Fig. 1). Importantly, transformants expressing RokL6 in the Δ *nagB* Δ *rokL6* double mutant via the introduction of pCOM-*rokL6* failed to grow, similar to the *nagB* mutant. These data strongly suggest that the deletion of *rokL6* was the sole reason why Δ *nagB* Δ *rokL6* double mutants could grow on GlcN (Fig. 1).

Conservation and gene synteny of *rokL6*

The gene *rokL6* (sco1447) from *S. coelicolor* consists of 1,200 nucleotides and encodes a protein of 399 amino acids. RokL6 is characterized by an N-terminal winged helix-turn-helix DNA-binding site, which is found in various families of DNA-binding proteins and a sugar kinase domain annotated as a putative ROK-family regulator. Protein alignment shows a high amino acid sequence conservation of RokL6 in *Streptomyces* species (Fig. S1). Its genomic neighbors, sco1446 and sco1448, encode a putative integral membrane protein and a MFS transporter with unknown substrates, and the intergenic regions of

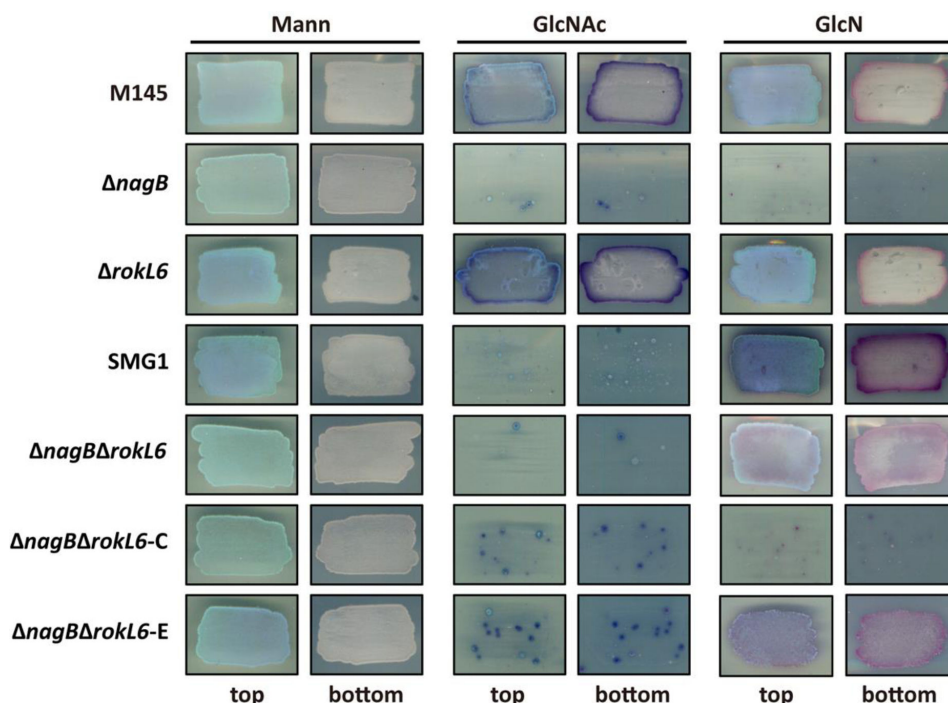


FIG 1 Sensitivity of *S. coelicolor* mutants to GlcN and GlcNAc. Spores (10^5 CFU) of *S. coelicolor* M145 or its mutant derivatives were streaked onto MM with either 1% (wt/vol) mannitol (Mann), 5 mM *N*-acetylglucosamine (GlcNAc), or 5 mM glucosamine (GlcN) and grown for 72 h at 30°C. Strains were *S. coelicolor* M145 (M145), its mutant derivatives Δ *nagB*, Δ *rokL6*, suppressor mutant SMG1, Δ *nagB* Δ *rokL6*, and the Δ *nagB* Δ *rokL6* mutant harboring either pCOM-*rokL6* (Δ *nagB* Δ *rokL6*-C) or empty vector pSET152 (Δ *nagB* Δ *rokL6*-E). Both top and bottom views are shown. Note that all strains without the gene *nagB* are sensitive to GlcNAc, while all strains lacking *rokL6* are resistant to GlcN.

rokL6-sco1446 and *rokL6*-sco1448 are 21- and 112-bp long, respectively (Fig. 2A). There is significant gene synteny for the genomic region surrounding *rokL6* and its orthologs in *Streptomyces*. In particular, sco1448 and its homologs often lie divergently transcribed from the corresponding ROK regulorencoding genes (Fig. 2B). The *rokL6*-sco1448 unit is also identified in many *Kitasatospora* species (Fig. S2).

Given the importance of ROK-family regulators in the control of sugar utilization (25) and the likelihood of the involvement of RokL6 in GlcN metabolism, we investigated its potential role in GlcN utilization by measuring the dry weights at different time points to compare the growth patterns of *S. coelicolor* M145 and its *rokL6* mutant in NMMP containing 1% GlcN as the sole carbon source. The mutant grew well in NMMP supplemented with either 1% glucose or with 1% GlcN (Fig. S3).

Transcriptome analysis of the *rokL6* mutant

To obtain further insights into the regulon of RokL6 and its relationship to GlcN metabolism, RNA sequencing (RNA-Seq) analysis was performed using RNA extracted from *S. coelicolor* M145 and its *rokL6* mutant grown on MM agar with either mannitol or mannitol + 50 mM GlcN as the carbon sources. Biomass was harvested at two time points corresponding to vegetative growth or sporulation. In total, 29 genes were significantly differentially expressed under at least one of the conditions analyzed, using an adjusted *P*-value < 0.01 and a log2 fold change >2 or <-2 as threshold (Table S1). Genes sco1446 and sco1448-sco1450, which flank *rokL6*, were upregulated in the *rokL6* mutant under all tested conditions (Fig. 3). In addition, sco0476 (for an unknown ABC transport protein)

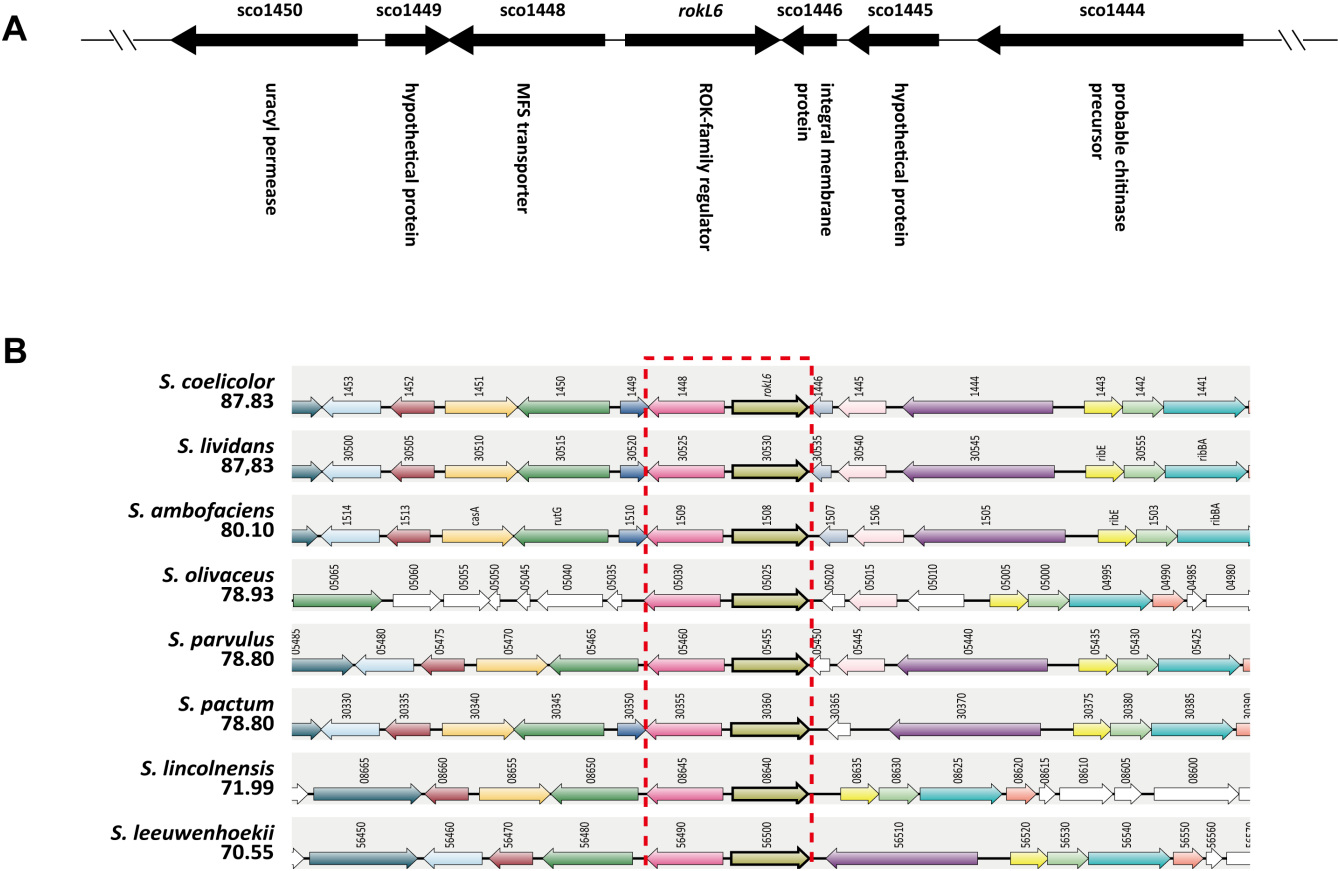


FIG 2 Genetic organization and gene synteny around *rokL6*. (A) Genetic organization of the genomic region around *rokL6*. All ORFs are depicted by solid arrows with their predicted gene products shown underneath. (B) Gene synteny of the region around *rokL6* in selected *Streptomyces* species. Synteny analysis was performed by SyntTax (scores are given). Homologous genes are presented in the same colors. Homologs of *rokL6* and its neighbor sco1448 are indicated in the dashed red box.

was downregulated at 24 h. In the absence of GlcN, transcription of the *pstSCA* operon (sco4140–4142), which is part of the PhoP regulon and related to the transport of inorganic phosphate (53), was significantly upregulated (greater than fourfold), while sco5338 (probable regulatory protein) and sco5339 (probable plasmid transfer protein) were downregulated in $\Delta rokL6$ during sporulation (Fig. 3A). In the presence of GlcN, the putative ABC transporter genes sco3704–sco3706 were upregulated in $\Delta rokL6$ during sporulation in the presence of GlcN; transcription of genes in the biosynthetic gene cluster for carotenoid biosynthesis (54), including *crtE* (sco0185), *crtI* (sco0186), *crtB* (sco0187), *crtY* (sco0191), *litQ* (sco0192), and the regulatory gene *litR* (sco0193), was all strongly downregulated in the *rokL6* mutant during vegetative growth (Fig. 3B). Indeed, compared to the wild-type M145, the production of carotenoids was significantly reduced in $\Delta rokL6$ in the presence of GlcN when plates were incubated in the light (Fig. S4), suggesting that RokL6 indeed plays a role in the control of carotenoid biosynthesis in *S. coelicolor*.

RokL6 specially binds to the intergenic region of *rokL6* and *sco1448*

To identify direct targets of RokL6 *in vivo*, we performed chromatin immuno-precipitation combined with sequencing. For this, we constructed a strain expressing RokL6 with a FLAG₃ tag, as described in the Materials and Methods section. We used

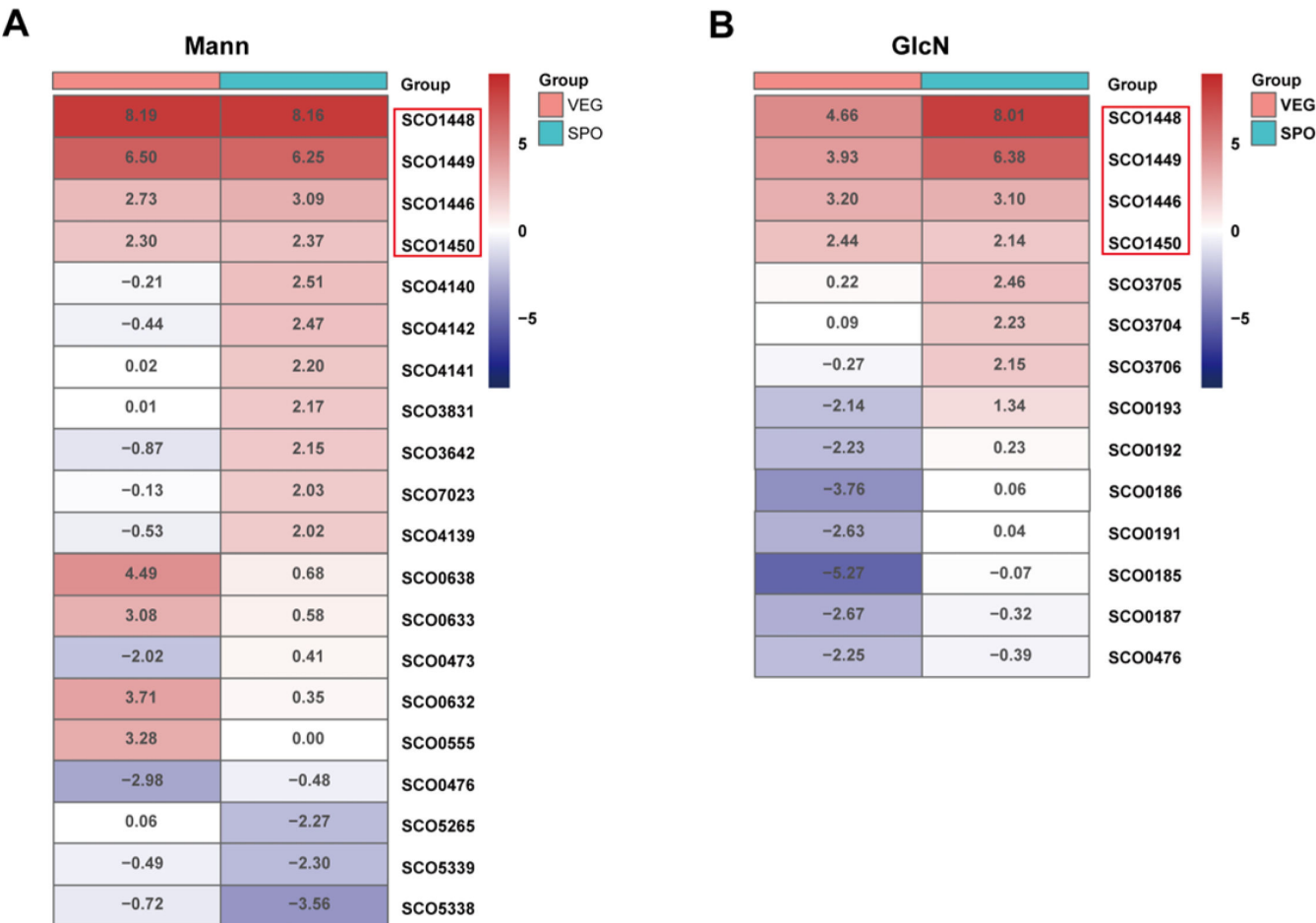


FIG 3 Heat maps of genes differentially expressed between the *rokL6* mutant and its parent *S. coelicolor* M145. Transcription patterns (expressed as log₂ fold changes $\Delta rokL6$ /wildtype) are presented for genes differentially expressed when grown on MM with mannitol (A) or on MM with mannitol and GlcN (B). Only genes with an adjusted *P*-value < 0.01 are shown. Navy, downregulated (log₂ fold change < -2) and brick red, upregulated (log₂ fold change > 2) in the *rokL6* mutant; intermediate log₂ fold changes are represented in white. Genes significantly differentially expressed under all growth conditions are highlighted with red boxes.

CRISPR-Cas9 technology to construct strain *S. coelicolor* M145-*rokL6*-FLAG, which has an engineered in-frame 3× FLAG epitope fused *in situ* and before the stop codon of *rokL6*. The functionality of RokL6-FLAG₃ was verified by introducing pCOM-FLAG into $\Delta nagB\Delta rokL6$. Indeed, while $\Delta nagB\Delta rokL6$ mutants grow well on MM with mannitol and GlcN, $\Delta nagB\Delta rokL6$ -FLAG expressing RokL6-FLAG₃ failed to grow (Fig. S5), showing that the RokL6-FLAG₃ was indeed expressed and active. ChIP-Seq analysis was performed with M145-*rokL6*-FLAG using cultures grown for 24 and 48 h on MM with mannitol

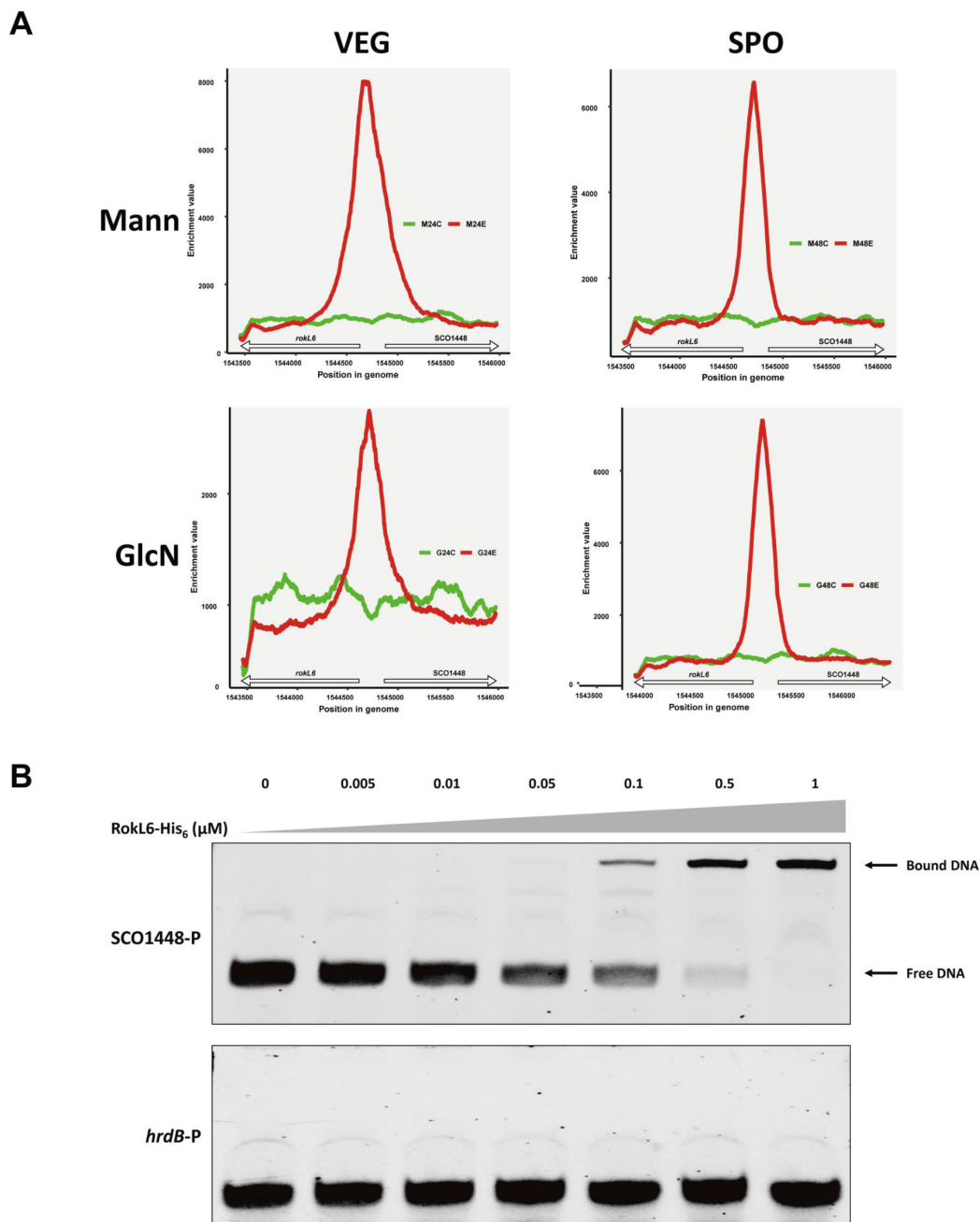


FIG 4 Identification of RokL6 binding sites *in vivo* and *in vitro*. (A) Enrichment peaks of RokL6 by ChIP-Seq. Red line, ChIP sample; green line, input chromosomal DNA used as a negative control. Genes flanking the peak summits are indicated. Numbers on the X-axis indicate genomic positions. Samples were collected on mannitol (Mann) or mannitol with GlcN (GlcN) at two time points, VEG and SPO. (B) EMSAs to establish direct binding *in vitro*. The binding of RokL6 to the intergenic sequence between *rokL6* and *sco1448* (*sco1448*-P) was tested by EMSAs, with the *hrdB* promoter (*hrdB*-P) sequence as the negative control. Concentrations are given in micromolar.

and MM with mannitol and GlcN. Under all conditions tested, only one major binding event was observed in all of the samples, namely to the intergenic region of *rokL6* and *sco1448* (Fig. 4A). Binding of RokL6 to the intergenic region shared between *rokL6* and *sco1448* was further verified *in vitro* by EMSAs. For this, C-terminally 6× His-tagged RokL6, RokL6-His₆, was expressed and purified from *E. coli* BL21(DE3). Compared to the probe for the negative control, *hrdB* promoter (*hrdB*-P), which exhibited no retardation in the presence of RokL6, the result shows that RokL6 caused a strong retardation on the probe corresponding to the promoter region of *rokL6*-*sco1448* (*sco1448*-P). Specifically, almost all *sco1448*-P was bound when 1 μM RokL6-His₆ was added in the EMSA (Fig. 4B).

RokL6 binds to the inverted repeat *rokL6*-IR

Due to the high conservation in terms of both amino acid sequence and gene synteny of *rokL6* and *sco1448* with their orthologs, we reasoned that the orthologs of RokL6 may be autoregulators. To identify a putative RokL6 binding consensus, the intergenic DNA sequence of *S. coelicolor* *rokL6* and *sco1448*, and 10 additional orthologous gene pairs from other streptomycetes (Table 3) were analyzed for the presence of conserved motifs using MEME. This identified a highly conserved inverted repeat sequence of 23 nucleotides in all intergenic regions (Fig. 5A). The consensus of the motif was C(T)TAT-CAGG-seven nt-CCTGATAG(A), which contains an inverted repeat designated as *rokL6*-IR, a primary candidate for the RokL6 binding site.

To verify if this is indeed a *bona fide* RokL6 binding site, EMSAs were performed using 60 bp probes. These probes contained either the intact *rokL6*-IR sequence (probe RokL6-BS) or mutant versions in which one or two regions of *rokL6*-IR were replaced by *Eco*RI or *Bam*HI restriction sites (probe RokL6-M1 to M3). Compared to RokL6-BS, which was bound at all tested concentrations (0.1, 0.2, and 0.5 μM) of RokL6-His₆, the binding signals of RokL6-His₆ to the mutated probes RokL6-M1 and RokL6-M2 were weakened, and to RokL6-M3 were fully abolished (Fig. 5B). These data show that the inverted repeat sequence *rokL6*-IR is indeed an essential part of the RokL6 binding site.

To determine if RokL6 has other binding sites on the *S. coelicolor* genome, the sequence of *rokL6*-IR was used to scan the *S. coelicolor* genome using the PREDetector algorithm (49). In addition to the promoter regions of *rokL6* and *sco1448*, one putative motif was identified with a relative high score upstream of *sco0137* encoding a PTS EII with an unknown substrate (Table S2). EMSAs showed that binding between RokL6 and the promoter of *sco0137* (*sco0137*-P) *in vitro* was very weak (Fig. S6A). The putative PTS transporter EII genes *sco0137* and *sco0136* are co-expressed from a single transcriptional unit; to see if the operon would be the major GlcN transporter, a mutant was created in both the wild-type strain *S. coelicolor* M145 and in the *nagB* mutant. For this, the region from nt position +10 of *sco0137* to nt position +1,542 of *sco0136* was replaced by *aac* (3)/IV (see Materials and Methods). The deletion of *sco0136*-*sco0137* in the *nagB* mutant

TABLE 3 Alignment of palindromic sequences found upstream of genes encoding RokL6 and its orthologs in streptomycetes

Microorganism	Gene name	Pos ^a	Sequence ^b
<i>S. coelicolor</i>	<i>rokL6</i>	-64	cgtaaa <u>ctatcagggag</u> Cctgcctgatagatagtggatta
<i>Streptomyces lividans</i>	SLIV_30530	-64	cgtaaa <u>ctatcagggag</u> Cctgcctgatagatagtggatta
<i>Streptomyces</i> CCM_MD2014	NI25_31970	-65	cgtaaa <u>ctatcagggag</u> Cctgcctgatagatagtgcatta
<i>Streptomyces ambofaciens</i>	SAM40697_1335	-64	cgtaaa <u>ctatcaggaac</u> Cctgcctgatagataggggatat
<i>Streptomyces pactum</i>	B1H29_30360	-65	cgtaaa <u>ctatcaggaac</u> Cctgcctgatagataggcgatta
<i>Streptomyces parvulus</i>	SPA_05455	-65	cgtaaa <u>ctatcagggag</u> Cctgcctgatagatagtggatta
<i>Streptomyces aquilus</i>	EJC51_09795	-51	cgtaaa <u>ctatcagggag</u> Cctgcctgatagatagagcgta
<i>Streptomyces chromofuscus</i>	IPT68_06320	-55	ggtaaa <u>ttatcagggag</u> Cctgcctgatagataggggcgt
<i>Streptomyces cacaioi</i> subsp. <i>asoensis</i>	G9272_09180	-74	cgtaaa <u>ctatcaggaac</u> Cctgcctgataaatagagggcga
<i>Streptomyces venezuelae</i> ATCC 21113	DEJ44_04995	-64	ggtaaa <u>ctatcaggaac</u> Cctgcctgataaatagaccgcag
<i>Streptomyces cadmiisoli</i>	DN051_31430	-72	ggtaaa <u>ctatcaggaac</u> Cctgcctgatagataggggaccc

^aPosition, distance of the central nucleotide (shown in uppercase) of the consensus sequence, relative to the translational start codon of the gene.

^bBases within palindromic sequences are underlined.

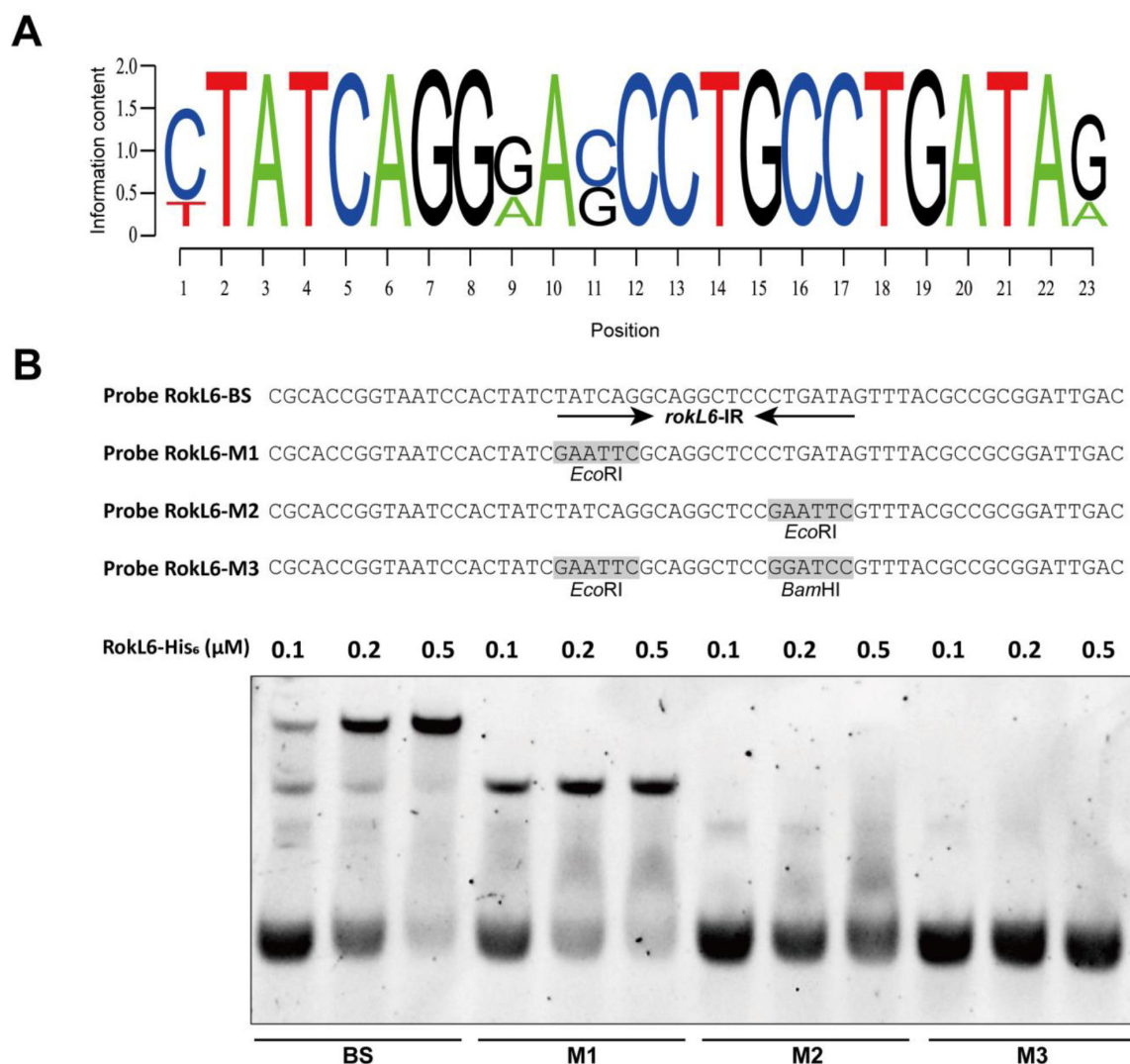


FIG 5 Identification of the RokL6 binding motif. (A) Sequence Logo representation of consensus sequence of RokL6 binding sites (analyzed by WebLogo). (B) RokL6 binding sites confirmation by EMSA. EMSAs of RokL6-His₆ with four probe sequences (RokL6-BS, RokL6-M1, RokL6-M2, and RokL6-M3) were tested. The palindromic sequences (*rokL6-IR*) in probe RokL6-BS are indicated by arrows. For each probe tested, the concentrations of RokL6-His₆ were 0.1, 0.2, and 0.5 μM.

did not relieve GlcN toxicity (Fig. S6B). This strongly suggests that *sco0136-sco0137* are at least not solely responsible for GlcN transport in *S. coelicolor*.

Identification of the transcription start sites for *rokL6* and *sco1448* and repression by RokL6

5' rapid amplification of cDNA ends (5' RACE) was applied to identify the promoters of both *rokL6* and *sco1448*. Fragments of 5' cDNA of *rokL6* and *sco1448* were PCR-amplified from the *S. coelicolor* genome and sequenced (Fig. S7). The TSS of *rokL6* was localized to a G located 67 nt upstream of the *rokL6* translation start codon (TSC), and the *sco1448* TSS was mapped to a G located 27 nt upstream of the *sco1448* TSC (Fig. 6A). The RokL6 binding site *rokL6-IR* encompasses nt positions -7 to +14 relative to the *rokL6* TSS, which is immediately downstream of the putative -10 region of the *rokL6* promoter. Suggestively, the *rokL6-IR* is located exactly between the -10 and -35 sequences of the *sco1448* promoter. This places the RokL6 binding site in the ideal position to repress both genes at the same time. Binding of the repressor close to the -10 or -35 consensus boxes for the RNA polymerase sigma factor is common, as it interferes with the recognition,

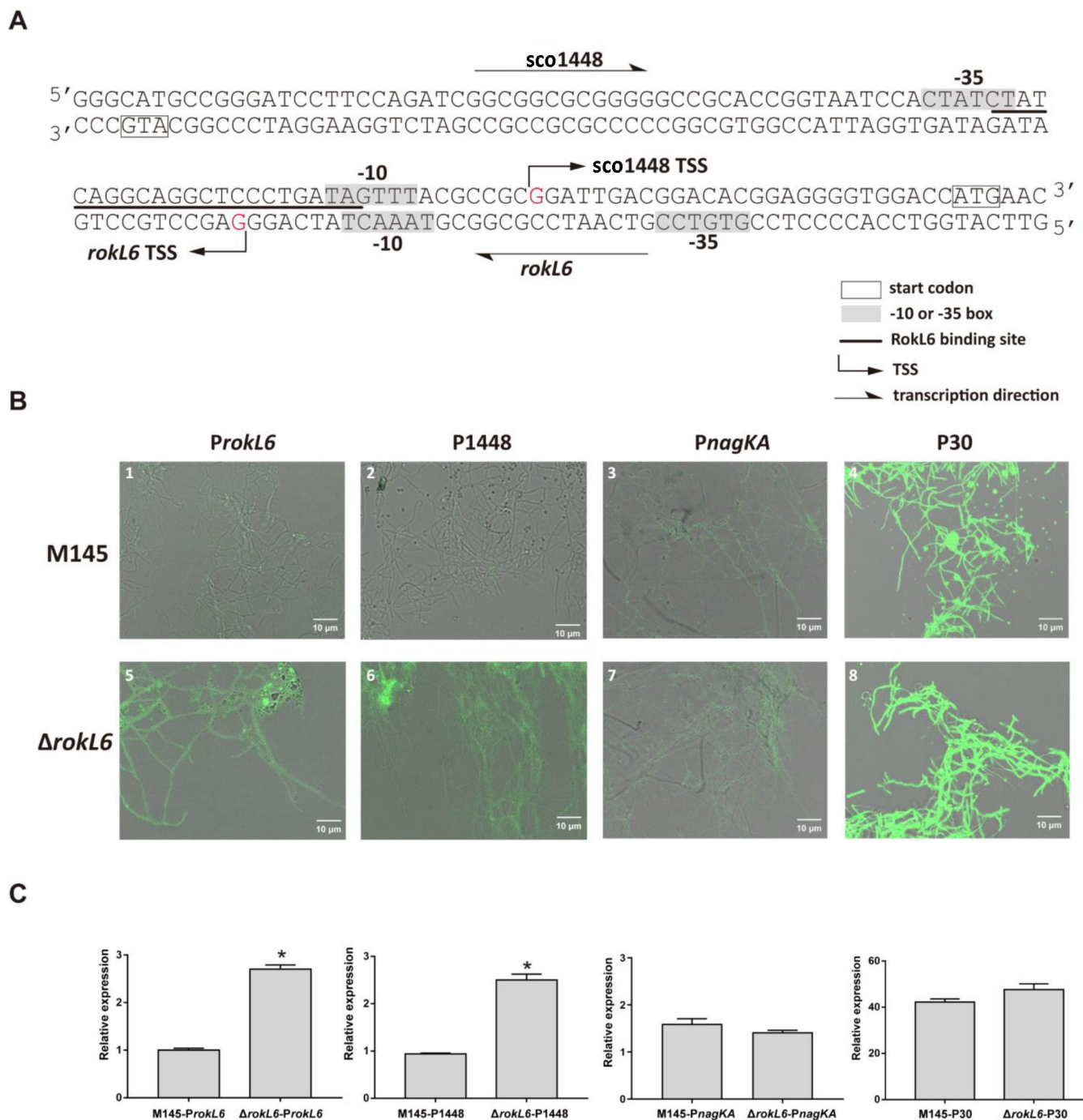


FIG 6 RokL6 acts as a transcriptional repressor of *sco1448* and as an auto-repressor. (A) Nucleotide sequences of *rokL6* and *sco1448* promoter region and RokL6-binding site. Number, distance (nt) from the respective TSC; bent arrows, TSS; boxes, start codons; gray shading, predicted -10 or -35 boxes based on 5'RACE (Fig. S7); underlined, RokL6 binding site; and straight arrow, direction of transcription. (B) Fluorescence intensity of eGFP measured based on confocal fluorescence micrographs. Mycelia of the strains expressing eGFP from different promoters were imaged using confocal microscopy. Strains analyzed as indicated: 1, M145-*ProkL6*; 2, M145-P1448; 3, M145-*PnagKA*; 4, M145-P30; 5, Δ*rokL6*-*ProkL6*; 6, Δ*rokL6*-P1448; 7, Δ*rokL6*-*PnagKA*; and 8, Δ*rokL6*-P30. (C) Relative transcription levels of eGFP. Transcription levels of eGFP gene expressed from different promoters (*ProkL6*, P1448, *PnagKA*, and P30) in M145 and Δ*rokL6* were measured by qPCR analysis. Relative transcription levels were normalized to the transcription level of the gene for eGFP with *ProkL6* in M145, which was set as 1. Data were calculated from triplicate biological experiments and presented as mean ± SD (**P* < 0.05).

binding of RNA polymerase, and the formation of the transcriptional initiation complex (55). This suggests that RokL6 represses the transcription of both *sco1448* and *rokL6* by

binding to the core promoter regions to prevent their transcription from proceeding properly.

To determine if this is indeed the case, promoter activity assays were performed using eGFP as the reporter gene cloned in integrative vector pSET152, using promoters *ProkL6* (*rokL6* promoter) and P1448 (*sco1448* promoter), with P30 and the *DasR*-controlled *nagKA* promoter (*PnagKA*) as the controls. The resulting vectors pEGFP-*rokL6*, pEGFP-1448, pEGFP-*nagKA*, and pEGFP-P30 were introduced into *S. coelicolor* M145 and Δ *rokL6*. The recombinant strains were grown on MM agar with 1% mannitol and analyzed for eGFP expression levels by fluorescence microscopy and qPCR. When eGFP was expressed from either the *nagKA* promoter or the artificial promoter P30, fluorescence was unchanged between M145 and Δ *rokL6*. Conversely, when expressed from either *ProkL6* or P1448, eGFP levels were significantly higher in Δ *rokL6* as compared to the parental strain M145 (Fig. 6B). This result was confirmed by qPCR for the transcription of the gene for eGFP, with enhanced expression of eGFP in the *rokL6* mutant as compared to the parent, while again no significant differences in expression levels were observed for either *PnagKA* or P30. However, the transcript levels of the gene for eGFP from both *ProkL6* and P1448 were significantly upregulated in Δ *rokL6* (Fig. 6C), suggesting that *RokL6* indeed represses the transcription of *sco1448* and at the same time acts as an autoregulator.

Overexpression of transporter SCO1448 relieves GlcN toxicity in *nagB* mutants

The data above demonstrate that putative transporter protein SCO1448 is the primary target of *RokL6*. We therefore hypothesized that SCO1448 may function as an exporter of one or more toxic metabolic intermediates that accumulate in *nagB* mutants grown on media containing GlcN. Inactivation of the repressor gene *rokL6* results in enhanced expression of SCO1448, alleviating GlcN toxicity, presumably by exporting a toxic intermediate. If this is indeed the case, expressing *sco1448* from a *RokL6*-independent (constitutive) promoter should have the same effect as deleting *rokL6*. To test this, *sco1448* was placed under the control of the strong and constitutive P30 promoter (see Materials and Methods for details) and introduced into Δ *nagB* to obtain SCO1448 overexpression strain, Δ *nagB*SCO1448-O. As a control, we used empty plasmid pSET152, to obtain Δ *nagB*-E. Transcription levels of *sco1448* were measured, showing that in both the de-repressed Δ *nagB* Δ *rokL6* and the overexpressing Δ *nagB*SCO1448-O strains, SCO1448 was highly expressed (Fig. 7A). Importantly, *RokL6*-independent expression of SCO1448 fully alleviated the toxicity of GlcN to *nagB* mutants, while Δ *nagB* or Δ *nagB* with the empty plasmid were still sensitive to GlcN (Fig. 7B).

Finally, we evaluated the viability of *sco1448* knock-out mutants in the presence of GlcN. For this, the coding regions of *sco1448*, spanning nucleotides + 6 to +1,023 (relative to the translational start site), were substituted with apramycin resistance cassette *aac* (3)/IV in *S. coelicolor* M145, Δ *nagB*, SMG1 (Δ *nagB* with a suppressor mutation in *rokL6*), and Δ *nagB* Δ *rokL6*. Homologous recombination of the gene was achieved by the introduction of vector pKO-1448, which was constructed by cloning the resistance cassette between the upstream and downstream flanking regions of *sco1448* in the unstable multi-copy plasmid pWHM3. Correct recombination events were confirmed by resistance to apramycin and sensitivity to thiostrepton (for loss of the plasmid) and by PCR, thus obtaining *sco1448* knock-out mutants, Δ *sco1448*, Δ *nagB* Δ *sco1448*, SMG1 Δ *sco1448*, and Δ *nagB* Δ *rokL6* Δ *sco1448*. As expected, when *sco1448* was deleted in suppressor mutant SMG1 or in Δ *nagB* Δ *rokL6*, neither strain could grow in the presence of GlcN (Fig. 7C). This again provides compelling evidence that *sco1448* is solely responsible for alleviating the toxicity of GlcN to *nagB* mutants. Taken together, our data show that *RokL6* directly represses the expression of *sco1448* and that, in turn, *sco1448* is responsible for alleviating GlcN toxicity to *nagB* mutants. It is likely that SCO1448 exports toxic metabolic intermediates derived from GlcN-6P, which accumulate in *nagB* mutants of *S. coelicolor* when grown on GlcN.

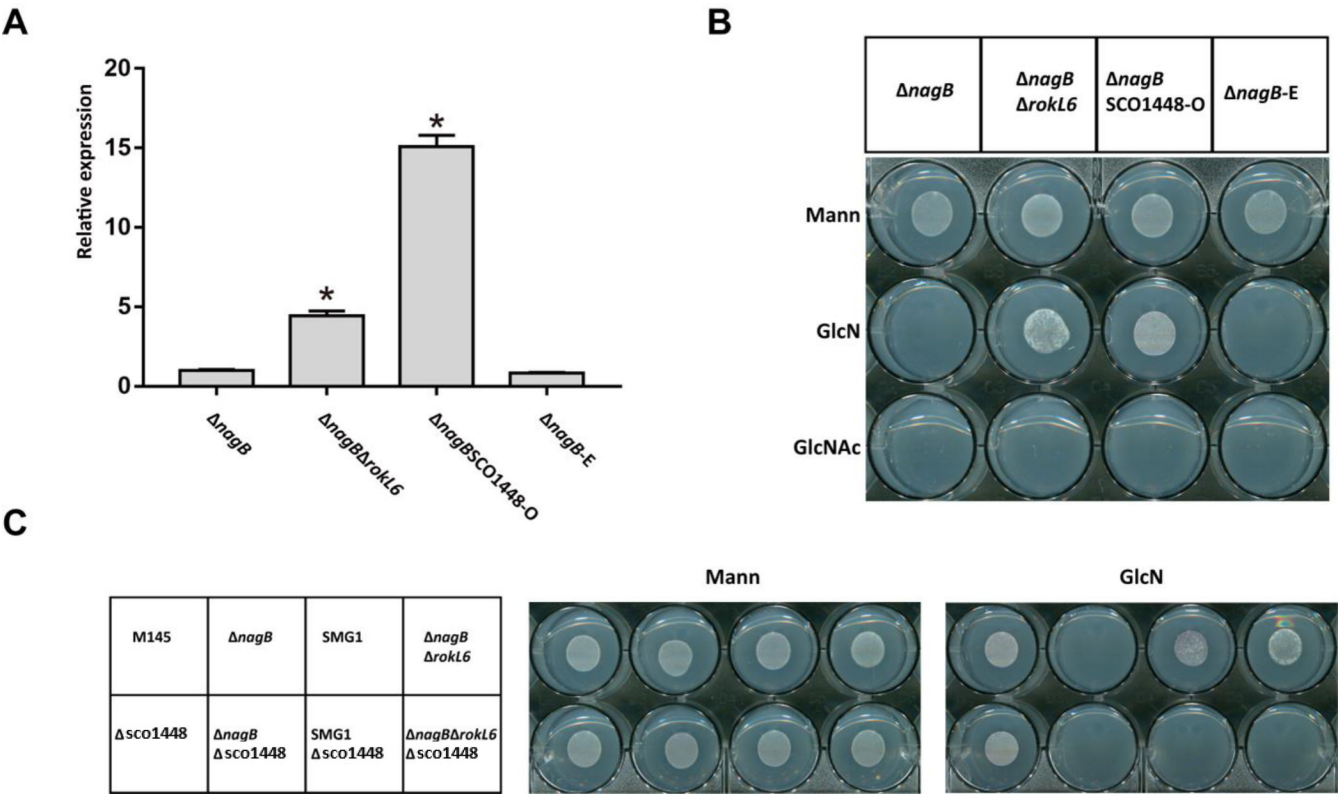


FIG 7 Constitutive expression of *sco1448* relieves the toxicity of *nagB* mutants to GlcN. (A) *sco1448* transcription levels analyzed by qPCR of the strains as follows: $\Delta nagB$, $\Delta nagB\Delta rokL6$, $\Delta nagB$ with overexpressed *sco1448* ($\Delta nagBSCO1448-O$), and $\Delta nagB$ complemented with empty pSET152 ($\Delta nagB-E$). qPCR data were calculated from three independent experiments and presented as mean \pm SD (* $P < 0.05$). (B) Growth of $\Delta nagBSCO1448-O$ on MM with GlcN(Ac). Spores of $\Delta nagB$, $\Delta nagB\Delta rokL6$, $\Delta nagBSCO1448-O$, and $\Delta nagB-E$, as indicated on the top, were spotted on MM with mannitol (Mann), MM with 1% mannitol and 5 mM GlcN (GlcN), and MM with 1% mannitol and 5 mM GlcNAc (GlcNAc) to determine their growth. (C) Growth of *sco1448* knock-out strains on GlcN. For media see panel B.

DISCUSSION

The aminosugar *N*-acetylglucosamine is a preferred nutrient for *Streptomyces* and also acts as a signaling molecule for the nutritional status of the environment. While GlcNAc metabolism and transport in streptomycetes have been well studied, little is known about how GlcN is metabolized in *Streptomyces*. *S. coelicolor nagB* mutants, which cannot convert GlcN-6P into the glycolytic intermediate Fru-6P, fail to grow on either GlcNAc or GlcN, indicating that toxic intermediates are produced when GlcN-6P is not actively metabolized by NagB. We used this feature to select for suppressor mutants, whereby it is important to note that some of the *nagB* suppressors selected on GlcNAc fail to confer resistance to GlcN and vice versa. In this work, we focused on the GlcN-specific gene *rokL6* (*sco1447*), the mutation of which relieves GlcN toxicity in *nagB* mutants.

RokL6 is an ROK-family regulator, and members of this family of regulators often play a role in the control of sugar metabolism. Systems-wide analysis using RNA-Seq, qPCR, ChIP-Seq, and EMSAs showed that RokL6 (SCO1447) directly represses the transcription of *sco1448*, which is annotated as an MFS sugar transporter. SCO1448 plays a key role in alleviating GlcN toxicity in the absence of NagB activity. Upregulation of *sco1448* is sufficient to relieve GlcN toxicity to *S. coelicolor nagB* mutants, while conversely, deletion of *sco1448* in *S. coelicolor ΔnagBΔrokL6* or the suppressor mutant SMG1 abolishes the acquired GlcN resistance (Fig. 7). Thus, the key to GlcN resistance lies in the expression of SCO1448. Still, a lot is unclear about the *rokL6-sco1448* gene pair. In particular, why would streptomycetes have a cryptic exporter to protect themselves from toxic intermediates related specifically to GlcN? The system is likely important, as its gene synteny and also (the location of) the RokL6 binding site are highly conserved in

Streptomyces and its sister genus *Kitasatospora* (Fig. 2B and Table 3). The high conservation of the binding site predicts that the expression of *sco1448* is repressed by RokL6 in many if not all streptomycetes. While the biological significance is not yet understood, the proteins likely play an important role in preventing the accumulation of excess toxic intermediates under specific growth conditions in the natural environment. How the repression by RokL6 is relieved in the cell and what the possible ligands are that control its DNA binding, which likely facilitates the derepression of *sco1448*, remains to be elucidated.

An important question is also what is the exact nature of the toxic molecule(s) that accumulate(s) in *nagB* mutants, which are likely derived from GlcN-6P. Mutants that lack a functional NagA enzyme accumulate high levels of GlcNAc-6P, which is lethal in *E. coli* and *Bacillus subtilis* (21, 22, 56). However, *S. coelicolor nagA* null mutants are able to grow in the presence of either GlcNAc or GlcN, and deletion of *nagA* from *nagB* null mutants of *S. coelicolor* also alleviates the toxicity of either aminosugar (23). Still, there are no metabolic routes that point to a key role for NagA in GlcN metabolism. Why then would mutation of *nagA* prevent toxicity of not only GlcNAc but also GlcN in *nagB* mutants? This again shows that there are still significant gaps in our understanding of aminosugar metabolism in *Streptomyces*. This is in fact quite surprising for such an important central metabolic pathway.

ChIP-Seq and EMSA assays demonstrated that RokL6 binds directly to the palindromic *rokL6*-IR identified from the intergenic region of *rokL6* and *sco1448*, with the sequence 5'-C(T)TATCAGG-seven nt-CCTGATAG(A)-3'. Furthermore, the precise transcription start sites for *rokL6* and *sco1448* were identified by 5' RACE, showing that the two genes are transcribed from overlapping promoters. RokL6 strategically binds to a site that is located precisely in between the -35 and -10 boxes for *sco1448* and downstream of the -10 box for *rokL6*, allowing RokL6 to inhibit the transcription of both genes at the same time. The transcriptional repression of *sco1448* and autoregulation of *rokL6* by RokL6 were further demonstrated by promoter activity assays and qPCR (Fig. 6), showing that RokL6 indeed inhibits the transcription of both genes.

The RokL6 binding site is distinct from operators characterized for ROK-family regulators in, e.g., *E. coli* and Firmicutes. The operator consensus sequences of NagC and Mlc in *E. coli* or XylR in firmicutes are typically composed of two A/T-rich inverted repeats separated by a spacer of 5–9 bp (18, 57, 58). The binding consensus sequences of RokB from *S. coelicolor* and RokA from *Streptococcus pneumoniae* are also enriched with T and A at the 5'-end and 3'-end (59, 60), similar to CysR from *Corynebacterium glutamicum* (61). The binding target of CsnR from *S. lividans* shares some similarity with that of RokL6, containing the sequence 5'-AGG-seven nt-CCT-3' in the binding consensus. In contrast, the binding consensus of Rok7B7 of *Streptomyces avermitilis* is 5'-TTKAMKHSTTSAV-3' and is unrelated to that of RokL6 (18, 25).

ChIP-Seq experiments identified a single binding site in all samples. Using that binding consensus as input, scanning the *S. coelicolor* genome using PREDetector revealed one additional sequence with similarity to the RokL6 consensus, namely upstream of the *sco0137-sco0136* operon that encodes PTS transporter EIIC enzymes. Its control by RokL6 and the fact that in *E. coli* and *B. subtilis*, GlcN is transported via the PTS (62–64), suggested a possible role for the *sco0137-sco0136* operon in GlcN transport. We tested whether the inactivation of the operon would affect GlcN toxicity in *nagB* mutants, but this is not the case (Fig. S6B). Therefore, PTS transporter *sco0136-sco0137* is at least not the only transporter for GlcN uptake in *S. coelicolor*, and more studies are required to understand how GlcN is internalized in streptomycetes.

In terms of indirect effects of the deletion of *rokL6*, besides *sco1448*, also the adjacent *sco1446* and *sco1448-sco1450* were upregulated in *rokL6* mutants. Other genes whose transcription was affected were those for transporters *sco0476*, *sco3704-sco3706*, and *sco4140-sco4142*, as well as genes of the carotenoid biosynthetic gene cluster (BGC). Upregulation of the latter BGC was validated by enhanced pigmentation of mycelia grown in the light (Fig. S4). However, our experiments strongly suggest that the

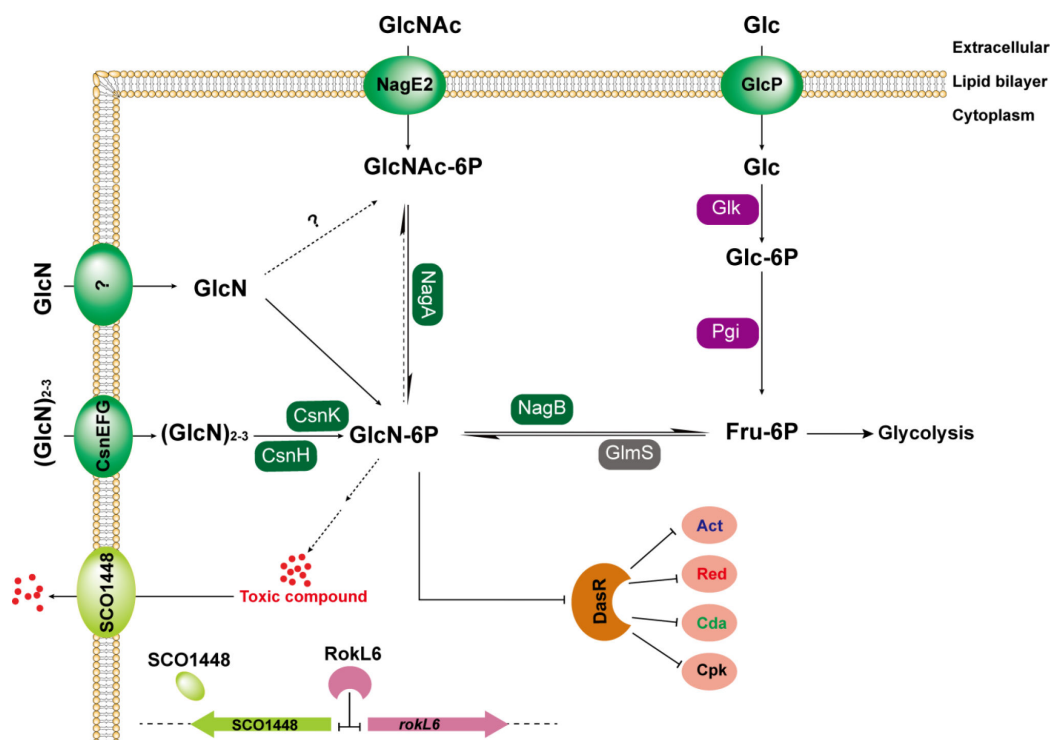


FIG 8 Model for aminosugar metabolism and the roles of RokL6 and SCO1448 in *S. coelicolor*. Metabolism of GlcNAc is well understood in *S. coelicolor*. GlcN-6P is an allosteric inhibitor of the global repressor DasR, which represses the biosynthesis of antibiotics. Based on our data, we propose that toxic substances are formed from GlcN metabolism, particularly from GlcN-6P, which accumulates in GlcN-grown *nagB* mutants; these toxic substances are likely exported by the MFS transporter SCO1448. We show that this transporter is transcriptionally repressed by the ROK-family regulator RokL6, and that enhanced expression of *sco1448* relieves GlcN toxicity. Metabolic routes are presented by black arrows with the enzymes as indicated/unknown routes by dotted arrows. Unknown transporters and enzymes are indicated by question marks. For details, see the text. Glc, glucose; 6P, 6-phosphate; GlcP, glucose permease; Glk, glucokinase; Pgi, glucose-6-phosphate isomerase; GlmS, glucosamine-fructose-6-phosphate aminotransferase; Act, actinorhodin; Red, prodiginines; Cda, calcium-dependent antibiotic; and Cpk, cryptic polyketide.

intergenic region between *rokL6* and *sco1448* is the primary binding site for RokL6, and the other genes whose transcription was affected in *rokL6* mutants are likely controlled indirectly.

Our current understanding of aminosugar metabolism and transport in *Streptomyces* is shown schematically in Fig. 8, including possible functions for RokL6 and SCO1448. GlcN-6P is derived either from the internalization of GlcN or the deacetylation of GlcNAc-6P. High accumulation of GlcN-6P and/or their metabolic derivatives is lethal to *S. coelicolor*, in line with observations in *E. coli* and *B. subtilis*. The MFS transporter SCO1448 likely serves to facilitate the export of toxic intermediate(s), and in turn, the transcription of *sco1448* is repressed by RokL6 (Fig. 8). Constitutive expression of *sco1448* does not relieve the toxicity of GlcNAc, suggesting that GlcNAc and GlcN toxicity are not caused by the same molecule(s).

Taken together, this study characterizes the function of the ROK-family regulator RokL6, including its regulons, binding sites, and its roles in the relief of GlcN toxicity in *S. coelicolor*. These findings shed new light on aminosugar sensitivity and on the control of aminosugar metabolism in *Streptomyces*, which plays a central role in their life cycle.

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AUTHOR CONTRIBUTIONS

Chao Li, Conceptualization, Formal analysis, Investigation, Methodology, Validation, Visualization, Writing – original draft, Writing – review and editing | Mia Urem, Conceptualization, Data curation, Formal analysis, Investigation, Methodology, Validation, Writing – original draft, Writing – review and editing | Chao Du, Data curation, Investigation, Methodology, Software, Validation, Writing – review and editing | Le Zhang, Investigation, Methodology, Supervision, Validation, Writing – review and editing | Gilles P. van Wezel, Conceptualization, Funding acquisition, Investigation, Project administration, Supervision, Writing – original draft, Writing – review and editing

DATA AVAILABILITY

Clean RNA-Seq reads and gene read-counts tables are available at GEO database (65)with accession number [GSE234437](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE234437). Clean ChIP-Seq reads and binding region identification (peak calling) files are available at GEO database with accession number [GSE234438](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE234438).

ADDITIONAL FILES

The following material is available [online](#).

Supplemental Material

Supplemental information (AEM01674-23-s0001.pdf). All supplemental tables and figures.

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