

# Repression of the Antifungal Activity of *Pseudomonas* sp. Strain DF41 by the Stringent Response<sup>∇</sup>

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The stringent response (SR) enables bacteria to adapt to nutrient limitation through production of the nucleotides guanosine tetraphosphate and guanosine pentaphosphate, collectively known as (p)ppGpp. Two enzymes are responsible for the intracellular pools of (p)ppGpp: RelA acts as a synthetase, while SpoT can function as either a synthetase or a hydrolase. We investigated how the SR affects the ability of the biological control agent *Pseudomonas* sp. strain DF41 to inhibit the fungal pathogen *Sclerotinia sclerotiorum* (Lib.) de Bary. Strain DF41 *relA* and *relA spoT* mutants were generated and found to exhibit increased antifungal activity. Strain DF41 produces a lipopeptide (LP) molecule that is essential for *Sclerotinia* biocontrol. LP production and protease activity were both elevated in the *relA* and *relA spoT* mutants. Addition of *relA* but not *spoT* in *trans* restored the mutant phenotype to that of the parent. Next, we investigated whether an association exists between the SR and known regulators of biocontrol, including the Gac system and RpoS. A *gacS* mutant of strain DF41 produced less (p)ppGpp and exhibited a 1.7-fold decrease in *relA* expression compared to the wild type, suggesting that *relA* forms part of the Gac regulon. We discovered that *rpoS* transcription was reduced significantly in the SR mutants. Furthermore, *rpoS* provided in *trans* restored protease activity to wild-type levels but did not attenuate antifungal activity. Finally, *relA* expression was decreased in the mutants, indicating that the SR is required for maximum expression of *relA*.

A number of bacteria are able to antagonize the effects of fungal pathogens through a process known as biological control. *Pseudomonas* sp. strain DF41 is one such bacterium that has demonstrated excellent antifungal activity against *Sclerotinia sclerotiorum* (Lib.) de Bary in both greenhouse and field assays (2, 32). Strain DF41 produces several secondary metabolites that are believed to contribute to biocontrol, including hydrogen cyanide, protease, and a novel lipopeptide (LP) molecule (1, 2). LP production has been shown to be essential for strain DF41 biocontrol, as an LP-deficient mutant, DF41-1278, demonstrated greatly reduced *S. sclerotiorum* inhibition (1, 2). LP synthesis is somewhat unique in that it does not involve ribosome-generated proteins; instead, these compounds are synthesized on large, multimodular enzymes termed nonribosomal peptide synthetases (NRPS) (24).

Several regulators have been found to govern secondary metabolite production in biocontrol strains of *Pseudomonas*. For example, the Gac two-component signal transduction system, comprised of the sensor kinase GacS and the cognate response regulator GacA, is essential for biocontrol. A mutation in either *gacS* or *gacA* results in a loss of biocontrol activity in several pseudomonads, including strain DF41 (2, 13). The alternative stationary-phase sigma factor, RpoS, has also been implicated in secondary metabolite production; however, regulation by RpoS appears to differ among species of pseudomonads. For example, an *rpoS* mutant of *Pseudomonas chlororaphis* PCL1391 exhibited decreased production of phenazine-1-carboxamide compared to the wild type (10),

whereas in *Pseudomonas fluorescens* Pf-5, an *rpoS* mutation resulted in enhanced pyoluteorin and 2,4-diacetylphloroglucinol expression (31). Several other global and pathway-specific regulators have been identified that influence the biocontrol properties of pseudomonads (12).

In addition to these regulatory elements, environmental conditions can have a significant impact on bacterial metabolism. When bacteria such as strain DF41 colonize the plant environment, they experience dramatic fluctuations in many environmental conditions, including nutrient availability. One means by which bacteria are able to survive starvation is through induction of a global stress mechanism known as the stringent response (SR). During the SR, cells accumulate the nucleotides guanosine tetraphosphate (ppGpp) and guanosine pentaphosphate (pppGpp), collectively referred to as (p)ppGpp. In gamma- and betaproteobacteria, (p)ppGpp accumulation is controlled by two enzymes, RelA and SpoT (see reference 27 and references therein). RelA is a synthetase that generates (p)ppGpp when available amino acids are in limiting amounts. SpoT is a bifunctional enzyme that can act as either a hydrolase or a synthetase depending on the conditions present (27). (p)ppGpp exerts its influence on cell physiology by binding RNA polymerase (RNAP) near the catalytic site. This leads to increased transcription of certain genes, for example, those involved in amino acid biosynthesis, and decreased transcription of others, for instance, tRNA and rRNA genes (27). Accordingly, the SR enables bacteria to alter their gene expression to favor activities that promote survival under nutrient-limiting conditions.

Although the SR has been shown to affect antibiotic production in *Streptomyces* species (4, 11, 15, 18, 19), there is a paucity of knowledge regarding how this global stress response influences biocontrol traits in pseudomonads. The purpose of

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TABLE 1. Bacterial strains, plasmids, and primers used in this study

Strain, plasmid, or primer	Genotype, phenotype, or sequence (5'–3') <sup>a</sup>	Source or reference
<b>Strains</b>		
<i>P. aeruginosa</i> strain PAO1	Wild type	17
<i>Pseudomonas</i> sp. strains		
DF41	Rif <sup>r</sup> ; wild type (canola root tip isolate)	32
DF41 <i>relA</i>	DF41 with a Gm <sup>r</sup> cassette inserted into the <i>relA</i> gene	This study
DF41 <i>relA spoT</i>	DF41 <i>relA</i> mutant with a Tet <sup>r</sup> cassette inserted into the <i>spoT</i> gene	This study
DF41-469	Rif <sup>r</sup> ; <i>gacS</i> ::Tn5-1063 genomic insertion	2
DF41-1278	Rif <sup>r</sup> ; <i>lp</i> ::Tn5-1063 genomic insertion	2
1278 <i>relA</i>	DF41-1278 with Gm <sup>r</sup> cassette inserted into the <i>relA</i> gene	This study
1278 <i>relA spoT</i>	1278 <i>relA</i> mutant with Tet <sup>r</sup> cassette inserted into the <i>spoT</i> gene	This study
<i>E. coli</i> strain DH5 $\alpha$	<i>supE44 ΔlacU169 φ80dlacZΔM15 hsdR17 recA1 endA1 gyrA96 thi-1 relA1</i>	Gibco
<b>Plasmids</b>		
pLP170	Promoterless <i>lacZ</i> transcriptional fusion	26
pRPOS- <i>lacZ</i>	<i>rpoS</i> promoter in pLP170	This study
pSW205	Promoterless <i>lacZ</i> translational fusion	9
pSW <i>relA</i>	<i>relA</i> promoter and first 21 codons of the open reading frame cloned into pSW205	This study
pCR2.1	TA cloning vector Amp <sup>r</sup>	Invitrogen
pCR <i>relA</i> -41	<i>relA</i> in pCR2.1	This study
pCR <i>spoT</i>	<i>spoT</i> in pCR2.1	This study
pEX18Ap	Suicide plasmid Amp <sup>r</sup>	16
pEX <i>relA</i>	<i>relA</i> in pEX18Ap	This study
pEX <i>relA</i> -800	pEX18Ap with an 800-bp deletion in <i>relA</i>	This study
pEX <i>relA</i> -Gent	pEX <i>relA</i> -800 Gm <sup>r</sup>	This study
pEX <i>spoT</i>	<i>spoT</i> in pEX18Ap	This study
pEX <i>spoT</i> -Tet	pEX <i>spoT</i> with Tet <sup>r</sup> cassette inserted into <i>spoT</i>	This study
pUCP22	Broad-host-range vector Amp <sup>r</sup> Gm <sup>r</sup>	36
pUCP <i>relA</i>	<i>relA</i> in pUCP22	This study
pUCP <i>spoT</i>	<i>spoT</i> in pUCP22	This study
pUCP22- <i>rpoS</i>	<i>rpoS</i> in pUCP22	26
pUCP23- <i>gacS</i>	<i>gacS</i> in pUCP23	26
pME- <i>rpoS</i>	<i>rpoS</i> in pME6010	This study
pME6010	pVS1-p15A shuttle cloning vector; Tet <sup>r</sup>	14
pME3219	pME6010 containing an <i>hcnA-lacZ</i> translational fusion	22
pRK600	Contains <i>tra</i> genes for mobilization; Chl <sup>r</sup>	8
pUCGm	Source of Gm <sup>r</sup> cassette	33
pFTC1	Source of Tet <sup>r</sup> cassette	5
<b>Primers</b>		
relA-FOR	ACCGTGGTAAAGGGTAGGCAAG	This study
relA-REV	GGGAAATCCCCCTGCTCTATG	This study
relAtransl-FRW	GGAATCCCCGCTTTTTTCAAGCCGAT	This study
relAtransl-REV	GAGGATCTCGCGGATCTCCA	This study
spoT-FOR	GCGTCACCGTTGAAGACTG	This study
spoT-REV	TTACTCGAGGACGACGATGG	This study
rpoSF	TACGTCAGTGCTTACGGCCA	This study
rsmZR	TATGACCCGCCACATTTTT	This study
P170fecorpoS	TGTGAATTTCGGGAGGGACA	This study
P170rxbarpoS	AGTCTAGAATCACCCTCCATTGCTT	This study

<sup>a</sup> Amp, ampicillin; Chl, chloramphenicol; Gm, gentamicin; Rif, rifampin; Tet, tetracycline.

this study was to investigate the impact of the SR on strain DF41 antifungal compound production. In addition, we examined whether there was a link between the SR and other known regulators of biocontrol, including the GacS/GacA two-component regulatory system and RpoS.

#### MATERIALS AND METHODS

**Bacterial strains and growth conditions.** All bacterial strains and plasmids used in this study are listed in Table 1. *Escherichia coli* strains were cultured on Lennox Luria-Bertani (LB) agar (Difco Laboratories, Detroit, MI). *Pseudomonas* sp. DF41 and its derivatives were routinely cultured at 28°C on King's B agar

plates supplemented with 2% glycerol (21). (p)ppGpp analysis was performed using cells grown in morpholinepropanesulfonic acid (MOPS; Sigma-Aldrich Canada, Oakville, Ontario, Canada) medium, as described by Cashel (3), supplemented with 1% Casamino Acids (Difco) and 400 μg/ml of DL-serine hydroxamate (Sigma). For LP analysis, cells were grown in M9 minimal medium (Difco) supplemented with 1% Casamino Acids, 1 mM MgSO<sub>4</sub>, and 0.2% glycerol. For β-galactosidase assays, strains were grown in M9 medium supplemented with 1 mM MgSO<sub>4</sub> and 0.2% glucose. *S. sclerotiorum* was maintained on potato dextrose agar (PDA; Difco). As required, media were supplemented with the following antibiotics from Research Products International Corp. (Mt. Prospect, IL): gentamicin (Gm; 20 μg/ml), tetracycline (Tc; 15 μg/ml), piperacillin (Pip; 100 μg/ml), and rifampin (Rif; 100 μg/ml) for strain DF41 and ampicillin (Amp; 100 μg/ml), Gm (15 μg/ml), and Tc (15 μg/ml) for *E. coli*.

**Nucleic acid manipulation.** Standard techniques were used for purification, cloning, and other DNA manipulations (30). PCR was performed under standard conditions suggested by Invitrogen Life Technologies data sheets supplied with the *Taq* polymerase.

**Sequence analysis.** PCR products were sequenced at the Centre for Applied Genomics at The Hospital for Sick Children (Toronto, Ontario, Canada), and the sequences were analyzed with the BLASTN and BLASTX databases.

**Generating *relA* and *relA spoT* mutants of strains DF41 and DF41-1278.** All primers used for the construction of mutant strains are listed in Table 1. The DF41 *relA* mutant strain was generated as follows. The *relA* gene of strain DF41 was amplified using primers *relA*-FOR and *relA*-REV. The resulting 2.4-kb fragment was then cloned into pCR2.1-TOPO (Invitrogen) to generate pCR*relA*-41. To liberate the insert, pCR*relA*-41 was digested with BamHI and EcoRI. The 2.4-kb fragment was subcloned into the same sites of the suicide vector pEX18Ap (16). Next, an 865-bp Sall fragment containing the Gm<sup>r</sup> cassette from pUCGm (33) was cloned into the Sall site of pEX*relA* to create pEX*relA*-Gent. Allelic exchange through triparental mating between the donor [*E. coli* DH5 $\alpha$ (pEX*relA*-Gent)], helper [*E. coli* DH5 $\alpha$ (pRK600)], and recipient (either DF41 or DF41-1278) strains was used to replace the wild-type *relA* gene with the mutated copy of the gene. Transconjugants were screened on LB agar supplemented with 100  $\mu$ g/ml Rif and 20  $\mu$ g/ml Gm. Bacteria that had undergone a double-crossover event were selected on LB agar containing 20% sucrose and 20  $\mu$ g/ml Gm. Verification of the mutation was achieved by PCR using the same primers. For creation of the DF41 *relA spoT* mutant, a 2.7-kb fragment containing the *spoT* gene was PCR amplified using primers *spoT*-FOR and *spoT*-REV. The resulting PCR product was subsequently cloned into pCR2.1-TOPO to generate pCR*spoT*. The *spoT* gene was then subcloned into pEX18Ap via SacI and XbaI sites. The resulting plasmid, pEX*spoT*, was digested with SmaI and ligated with a 2.0-kb SmaI fragment containing the Tc<sup>r</sup> marker from pFTC1 (5) to generate pEX*spoT*-Tet. Triparental mating was performed using *E. coli* DH5 $\alpha$ (pEX*spoT*-Tet), *E. coli* DH5 $\alpha$ (pRK600), and the DF41 *relA* or DF41-1278 *relA* strain. Transconjugants were screened on LB agar supplemented with 15  $\mu$ g/ml Tc and 100  $\mu$ g/ml Rif. Sucrose plates containing Tc were used to identify bacteria that had undergone a double-crossover event. To confirm the insertion of the Tc marker into the *spoT* gene, Southern blot analysis was performed.

**Construction of plasmids.** All primers are listed in Table 1. To generate the *relA* overexpression plasmid pUC*PreLA*, primers *relA*transFRW and *relA*-REV were used to amplify a 2.9-kb fragment. The PCR product was cloned into pCR2.1-TOPO to yield pCR*relA*-41. pCR*relA*-41 was then digested with EcoRI, and the 2.8-kb fragment containing the *relA* gene was cloned into pUCP22, placing the gene under the control of the plasmid-borne *lac* promoter. The *spoT* overexpression vector pUC*PSpoT* was created by first amplifying the *spoT* gene from strain DF41, using primers *spoT*-FOR and *spoT*-REV. The 2.8-kb fragment containing the gene was cloned into pCR2.1-TOPO to generate pCR*spoT*, which was subsequently digested with XbaI. The linearized plasmid was treated with Klenow DNA polymerase (Invitrogen) and digested again with BamHI. The 2.8-kb insert was cloned into the BamHI and SmaI sites of pUCP22 such that the transcription of *spoT* was dependent on the *lac* promoter. To create an *rpoS-lacZ* transcriptional fusion, the *rpoS* gene was amplified from DF41 genomic DNA by using primers *rpoS*F and *rsmZR*. The resulting 1.7-kb fragment was cloned into the pCR2.1-TOPO vector, resulting in pCR2.1-*rpoS*. Using pCR2.1-*rpoS* as the template, a 1.1-kb fragment was amplified using primers P170fecorpoS and P170rxbarpoS, which contain EcoRI and XbaI sites, respectively. The PCR product was digested with EcoRI and XbaI and cloned into the same sites of pLP170 (28), generating pRPOS-*lacZ*. For the *relA-lacZ* translational fusion pSW*relA*, a 644-bp fragment containing 72 nucleotides upstream of the ATG translational start site was PCR amplified from strain DF41 genomic DNA, using primers *relA*transFRW and *relA*transREV. The product was digested with EcoRI and SmaI and cloned into the same sites of pSW205 (9). To create pME-*rpoS*, containing *rpoS* constitutively expressed from the kanamycin promoter, a 1.3-kb KpnI-HindIII fragment was isolated from pUCP22-*rpoS* and ligated into the same sites of pME6010 (14).

**(p)ppGpp and *relA* expression analysis.** Determination of (p)ppGpp levels was performed as described by Cashel (3), with the following modifications. Cells were grown overnight in MOPS minimal medium (3) at 28°C, followed by 1/100 dilution of the culture in MOPS phosphate-free minimal medium containing 1 mg/ml of Casamino Acids and 100  $\mu$ Ci/ml of <sup>32</sup>P (Perkin Elmer, Waltham, MA). Three 200- $\mu$ l aliquots of culture were added to wells of a polystyrene microtiter plate (Costar; Corning Incorporated, Corning, NY) and grown at 28°C for an additional 8 h. DL-Serine hydroxamate was added to each well at a concentration of 400  $\mu$ g/ml and allowed to incubate for 2 h. To ensure that an equivalent number of cells was extracted for each strain, PA23 and its derivatives were grown on a parallel plate as described above, but excluding the <sup>32</sup>P. All of the

cultures showed equivalent turbidity levels (optical densities at 600 nm [OD<sub>600</sub>]). Thus, the three radiolabeled 200- $\mu$ l aliquots were pooled for each strain, and nucleotides were extracted with an equal volume of cold 13 M formic acid. A 20- $\mu$ l aliquot of the nucleotide sample was separated on polyethyleneimine-cellulose chromatography sheets (Sigma), using 1.5 M KH<sub>2</sub>PO<sub>4</sub> as the solvent, which was allowed to ascend the plate for 2.5 h. The spots were visualized by autoradiography. For the (p)ppGpp analysis, *Pseudomonas aeruginosa* strain PAO1 was included as a positive control, and the experiments were repeated five times. Expression of an *relA-lacZ* translational fusion was monitored in the DF41, DF41 *relA*, and DF41 *relA spoT* strains and in the *gacS* mutant strain DF41-469. Strains harboring pSW*relA* were grown for 4, 8, 16, and 24 h and then assayed for  $\beta$ -galactosidase activity (25).

**Antifungal assays.** To assess the ability of strain DF41 and its derivatives to inhibit fungal growth *in vitro*, radial diffusion assays were performed according to the method of Poritsanos et al. (26). Five replicates for each strain were analyzed, and the experiments were repeated three times.

**HPLC analysis.** LP was extracted from cultures and analyzed by high-performance liquid chromatography (HPLC) as described by Berry et al. (2), with the following modifications. To determine the efficiency of the extraction, surfactin (Sigma) was used as an internal control. Cell-free supernatants were spiked with a 500- $\mu$ l aliquot of surfactin (1-mg/ml stock concentration), which was extracted with the LP. The chromatograms obtained for each strain were normalized using the surfactin peak height. We have previously shown that the peak at 28 min corresponds to the DF41 LP (2); consequently, the amount of LP present in the extracts was determined by measuring the height of the 28-min peak.

**Bioluminescence.** Five-milliliter cultures were grown in M9 minimal medium plus 1 mM MgSO<sub>4</sub> plus 0.2% glucose and assayed by the OD<sub>600</sub> to determine the number of cells present. The cultures were centrifuged and resuspended in an equal volume of 1 $\times$  phosphate-buffered saline (PBS). A 1-ml aliquot of the cell suspension was mixed with 1 ml of 2 $\times$  523 medium supplemented with 10 mg/liter of sodium citrate (20). A 2.5- $\mu$ l aliquot of a 10% *n*-decanol solution was added to each tube, which was mixed for 15 s with a vortex mixer and measured for bioluminescence 90 s later by use of a BG-P luminometer (GEM Biomedical Inc., Hamden, CT). Bioluminescence was expressed in relative light units (RLU) by dividing the total bioluminescence signal by the OD<sub>600</sub>.

**HCN analysis.** Qualitative determination of hydrogen cyanide production was performed using Cyantesmo paper (Macherey-Nagel GmbH & Co., Duren, Germany). Plasmid pME3219, harboring an *hcnA-lacZ* translational fusion (22), was mobilized into the DF41, DF41 *relA*, and DF41 *relA spoT* strains. Strains were grown for 24 h and then assayed for  $\beta$ -galactosidase activity. Samples were analyzed in triplicate, and experiments were repeated three times.

**Protease production.** Cultures were grown in M9 minimal medium supplemented with 1 mM MgSO<sub>4</sub>, 0.2% glucose, and 1.5% skim milk (Difco) for 5 days at 28°C to induce protease production. A 200- $\mu$ l aliquot of cell-free supernatant was analyzed for the activity of this enzyme in a 0.65% solution of casein according to the method of Cupp-Enyard (7). Tyrosine, which is released upon the hydrolysis of casein by the protease enzyme, is able to react with the Folin-Ciocalteu reagent (Sigma) to produce a blue chromophore (7). This chromophore is measured spectrophotometrically at a wavelength of 660 nm. To determine the amount of tyrosine liberated, a standard curve was generated using pure tyrosine at the following concentrations: 0.055, 0.111, 0.221, 0.442, and 0.553  $\mu$ M. Each strain was assayed in triplicate, and experiments were performed three times.

***rpoS* expression in DF41, DF41 *relA*, and DF41 *relA spoT* strains.** Expression of the *rpoS* gene was monitored using pRPOS-*lacZ*. Cultures of the DF41, DF41 *relA*, and DF41 *relA spoT* strains harboring pRPOS-*lacZ* were grown for 4, 8, 16, 24, and 48 h and then measured for  $\beta$ -galactosidase activity.

**Nucleotide sequence accession numbers.** The GenBank accession numbers for the sequences of the DF41 *rpoS*, *relA*, and *spoT* genes are EU595545.1, HQ615419, and HQ615420, respectively.

## RESULTS

**Generation of DF41 *relA* and *relA spoT* mutant strains and (p)ppGpp analysis.** To determine the impact of the SR on the biocontrol activity of strain DF41, *relA* and *relA spoT* mutants were created. Primers designed from the *relA* and *spoT* sequences of *Pseudomonas aeruginosa* PAO1 were used to amplify these genes from strain DF41 genomic DNA. Sequence analysis revealed the DF41 alleles to be 99% identical to the *relA* and *spoT* genes of *P. aeruginosa* PAO1 (GenBank ac-

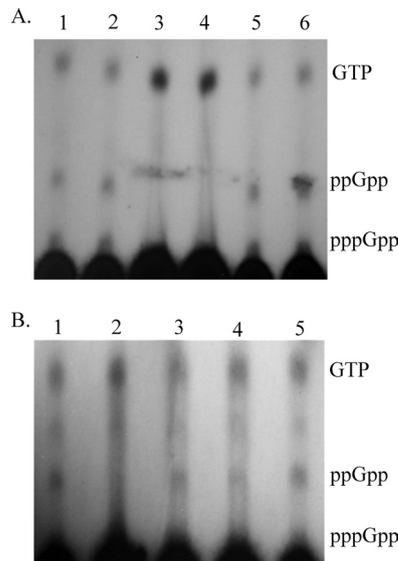


FIG. 1. (p)ppGpp analysis of *Pseudomonas* sp. strain DF41 and its SR derivatives after serine hydroxamate induction. Cells were labeled with  $^{32}\text{P}$ , and nucleotides were extracted and separated by thin-layer chromatography. (A) Lane 1, *Pseudomonas aeruginosa* strain PAO1; lane 2, *Pseudomonas* sp. DF41; lane 3, DF41 *relA* mutant; lane 4, DF41 *relA spoT* mutant; lane 5, DF41 *relA*(pUCPreLA) mutant; lane 6, DF41 *relA spoT*(pUCPreLA) mutant. (B) Lane 1, DF41; lane 2, DF41-469 (*gacS* mutant); lane 3, DF41-469(pUCP23-*gacS*); lane 4, DF41-469(pUCPreLA); lane 5, PAO1.

cession no. AE004091.2), 85% identical to those of *P. fluorescens* Pf0-1 (GenBank accession no. CP000094.2), and 84% identical to those of *P. fluorescens* Pf-5 (GenBank accession no. CP000076.1). DF41 *relA* and *relA spoT* mutants were generated through allelic exchange. Double-crossover mutations were confirmed by PCR and Southern blot analysis (data not shown). We were unable to isolate an *spoT* single null mutant, indicating that the absence of SpoT in an *relA*<sup>+</sup> background is likely lethal, similar to what has been reported for other bacteria (27, 37).

Next, we analyzed (p)ppGpp levels in the aforementioned strains. In the presence of DL-serine hydroxamate, which induces the SR, no (p)ppGpp was detected in the *relA* and *relA spoT* mutants, unlike the case in strain DF41, which produced both nucleotides (Fig. 1A). The presence of pUCPreLA, a vector harboring the strain DF41 *relA* gene, restored (p)ppGpp production in both mutants (Fig. 1A, lanes 5 and 6). The presence of *spoT* on plasmid pUCPspoT, however, did not alter (p)ppGpp production in the mutants (data not shown).

**SR affects strain DF41 antifungal activity.** When we assessed the ability of the mutants to inhibit *S. sclerotiorum* *in vitro*, we observed 1.5-fold and 1.8-fold increases in antifungal activity for the DF41 *relA* and DF41 *relA spoT* strains, respectively, compared to the parent (Table 2). The presence of pUCPreLA was able to restore the antifungal activities of both mutants to that of the wild type (Table 2). Production of many secondary metabolites begins at the transition between the logarithmic and stationary phases, also known as the idiophase. As such, it is important that there were no differences in growth rate between the wild type, the SR mutants, and the complemented strains (see Fig. 4A).

TABLE 2. Antifungal activity of *Pseudomonas* sp. strain DF41 and its derivatives after 5 days of growth

Strain	Zone of fungal growth inhibition (mm) <sup>a</sup>
DF41(pUCP22)	4.58 (1.0)
DF41 <i>relA</i> (pUCP22)	7.25 (0.5) <sup>b</sup>
DF41 <i>relA spoT</i> (pUCP22)	8.30 (1.5) <sup>c</sup>
DF41 <i>relA</i> (pUCPreLA)	4.75 (1.2) <sup>d</sup>
DF41 <i>relA spoT</i> (pUCPreLA)	5.50 (1.2) <sup>d</sup>

<sup>a</sup> Mean (standard deviation) obtained for five replicates.

<sup>b</sup> Significantly different from the wild type ( $P < 0.005$ ).

<sup>c</sup> Significantly different from the wild type ( $P < 0.05$ ).

<sup>d</sup> Not significantly different from the wild type.

**Secondary metabolite production by strain DF41 and its SR mutant derivatives.** Because LP production is essential for strain DF41 biocontrol (1, 2), we investigated whether the ppGpp-deficient strains exhibited elevated LP expression. A derivative of strain DF41, termed DF41-1278, is LP deficient due to a Tn5-1063 (*luxAB*) insertion in the NRPS locus responsible for LP synthesis (1, 2). Consequently, it is possible to monitor transcription of the NRPS genes in strain DF41-1278 by using a bioluminescence assay (1). Two SR mutants of strain DF41-1278 were created, designated the 1278 *relA* and 1278 *relA spoT* mutants. After 12 h of growth, NRPS transcription in the 1278 *relA* and 1278 *relA spoT* strains was elevated 5.6-fold and 6.1-fold, respectively, over that of the wild type (Fig. 2). Addition of *relA* *in trans* restored transcription to levels close to wild-type levels (Fig. 2). By 24 h, all strains exhibited equal levels of transcription (data not shown). Next, we determined the level of LP present in DF41, DF41 *relA*, and DF41 *relA spoT* culture supernatants through HPLC analysis. A single peak eluting at 28 min (Fig. 3) was observed for all strains except for strain DF41-1278 (Fig. 3B). We demonstrated previously that this 28-min peak corresponds to the DF41 LP molecule (2). For the SR mutants, the 28-min peak was 1.5- to

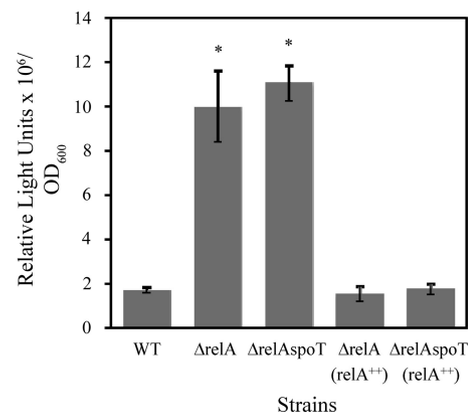


FIG. 2. Transcription of NRPS genes in *Pseudomonas* sp. strain DF41-1278 and in the 1278 *relA*, 1278 *relA spoT*, and *relA*-complemented mutants. Column 1, DF41-1278(pUCP22); column 2, 1278 *relA*(pUCP22) strain; column 3, 1278 *relA spoT*(pUCP22) strain; column 4, 1278 *relA*(pUCPreLA) strain; and column 5, 1278 *relA spoT*(pUCPreLA) strain. Bioluminescence was monitored after 12 h of growth in M9 minimal medium supplemented with 1 mM MgSO<sub>4</sub> and 0.2% glucose. For strains that differ significantly from the wild type, columns are labeled with asterisks (\*,  $P < 0.005$ ).

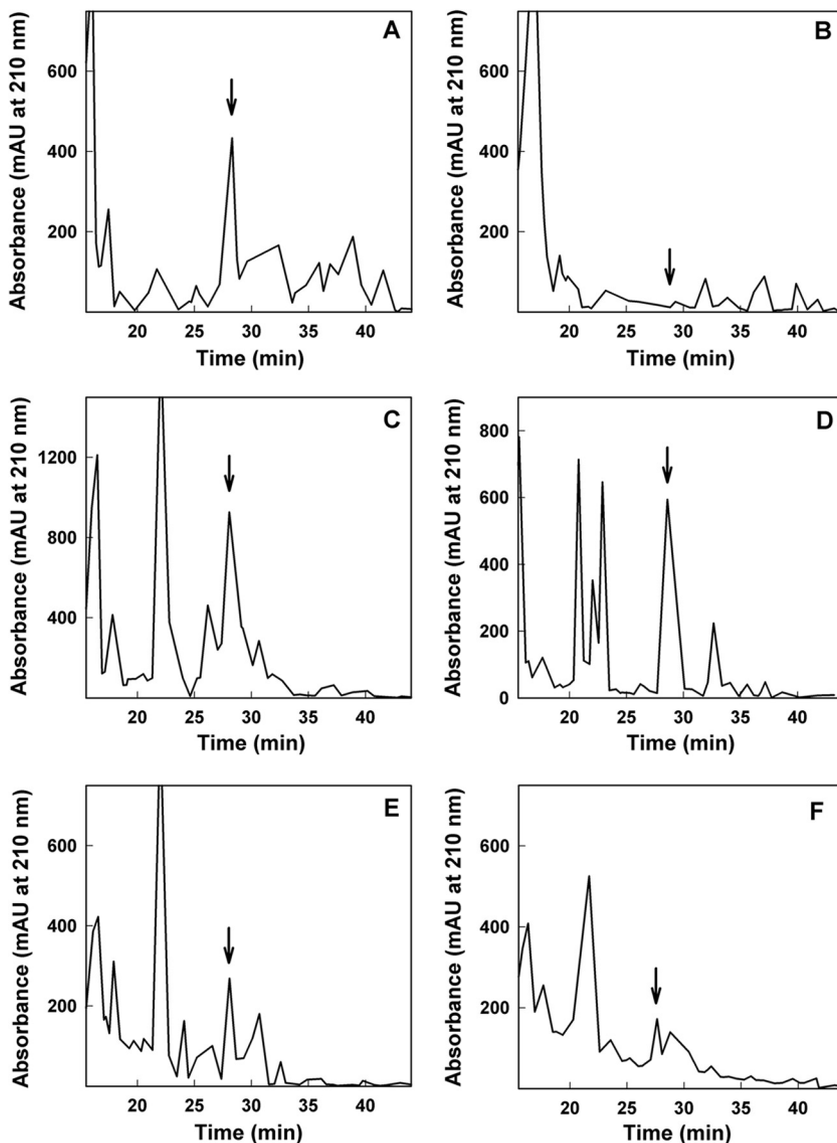


FIG. 3. HPLC analysis of LP produced by *Pseudomonas* sp. strain DF41, the stringent response mutants, and strains harboring pUCPreLA. LP was extracted from cell-free supernatants of 4-day-old cultures and separated by HPLC. Chromatograms depict the following strains: DF41(pUCP22) (A), DF41-1278(pUCP22) (B), DF41 *relA*(pUCP22) (C), DF41 *relA spoT*(pUCP22) (D), DF41 *relA*(pUCPreLA) (E), and DF41 *relA spoT*(pUCPreLA) (F). Peaks containing the strain DF41 LP molecule are indicated with arrows.

2.0-fold larger (Fig. 3C and D) than that for the wild type (Fig. 3A). When *relA* was expressed in *trans*, LP production by the SR mutants was markedly reduced (Fig. 3E and F).

Besides LP molecules, strain DF41 liberates the volatile antibiotic HCN (1, 2). Using Cyantesmo paper, we determined that the wild type and the SR mutants all produced HCN (data not shown). Furthermore, no differences in *hcnA-lacZ* expression were observed between the DF41 (888 ± 166 Miller units), DF41 *relA* (1,163 ± 124 Miller units), and DF41 *relA spoT* (1,152 ± 113 Miller units) strains.

Quantitative analysis of protease production revealed that the DF41 *relA* and DF41 *relA spoT* mutants produced 2-fold more protease than the wild type (Table 3). Addition of pUCPreLA restored the protease activity of the SR mutants to wild-type levels (Table 3).

TABLE 3. Protease activity of *Pseudomonas* sp. strain DF41 and derivatives harboring the *relA* overexpression plasmid pUCPreLA and the *rpoS* overexpression plasmid pUCP22-*rpoS*

Strain	Protease activity (U of enzyme/ml) <sup>a</sup>
DF41(pUCP22).....	0.752 (0.04)
DF41 <i>relA</i> (pUCP22).....	1.480 (0.01) <sup>b</sup>
DF41 <i>relA spoT</i> (pUCP22).....	1.487 (0.02) <sup>b</sup>
DF41 <i>relA</i> (pUCPreLA).....	0.882 (0.10) <sup>c</sup>
DF41 <i>relA spoT</i> (pUCPreLA).....	0.914 (0.20) <sup>c</sup>
DF41 <i>relA</i> (pUCP22- <i>rpoS</i> ).....	0.677 (0.10) <sup>c</sup>
DF41 <i>relA spoT</i> (pUCP22- <i>rpoS</i> ).....	0.735 (0.02) <sup>c</sup>

<sup>a</sup> Mean (standard deviation) for five replicates.

<sup>b</sup> Significantly different from the wild type (*P* < 0.005).

<sup>c</sup> Not significantly different from the wild type.

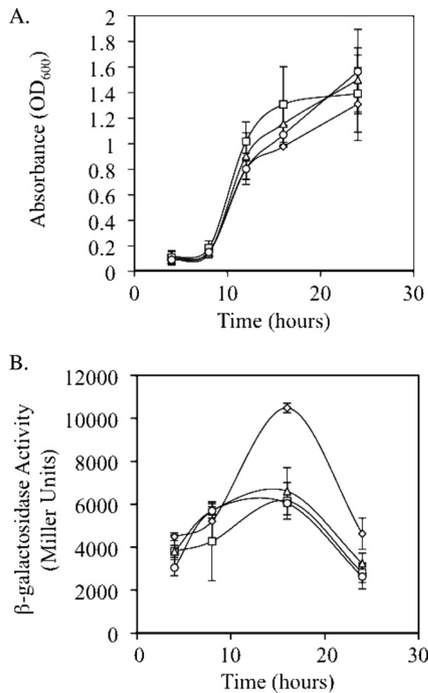


FIG. 4. Ability of GacS and the stringent response to regulate *relA* translation. Growth curves for each strain are depicted in panel A, while translational analysis of *relA* is shown in panel B. Strains used were as follows: *Pseudomonas* sp. strain DF41 (diamonds), DF41 *relA* mutant (squares), DF41 *relA spoT* mutant (triangles), and DF41-469 (circles). Strains in panel B harbor the *relA* translation fusion pSW*relA*. Strains were grown in M9 minimal medium supplemented with 1 mM MgSO<sub>4</sub> and 0.2% glucose. Note that *relA* expression was markedly reduced in both the SR mutants and the *gacS* mutant.

#### *relA* expression and (p)ppGpp production are reduced in a *gacS* mutant of strain DF41.

To ascertain whether a link exists between the Gac system and the SR, (p)ppGpp production was assessed in a *gacS* mutant of strain DF41 called DF41-469. As observed in Fig. 1B, (p)ppGpp levels were markedly reduced in the *gacS* mutant. Production of these nucleotides was increased by the addition of either *gacS* or *relA* in *trans* (Fig. 1B). Next, we generated an *relA* translational fusion and monitored its activity in strains DF41 and DF41-469 (*gacS*). Although there were no differences in growth rate between the two strains (Fig. 4A), a 1.7-fold reduction in *relA* expression was observed in the *gacS* mutant compared to the wild type at 16 h (Fig. 4B). Addition of pUCP23-*gacS* in *trans* was unable to complement the DF41 *relA* and DF41 *relA spoT* mutants for any of the aforementioned phenotypes, including LP production and antifungal and protease activities (data not shown).

**The stringent response is required for maximal *relA* expression.** Expression of an *relA* translational fusion was monitored in the DF41, DF41 *relA*, and DF41 *relA spoT* strains. In strain DF41, *relA* expression reached its maximum at 16 h, followed by a sharp decline by 24 h (Fig. 4B). In the SR mutants, *relA* expression peaked at 16 h; however, expression was reduced 1.6-fold in these strains. Thus, it appears that in strain DF41, the SR is required for maximal *relA* expression.

**RpoS overexpression complements *rpoS* transcription and protease activity in SR mutants.** In strain DF41, *rpoS* was

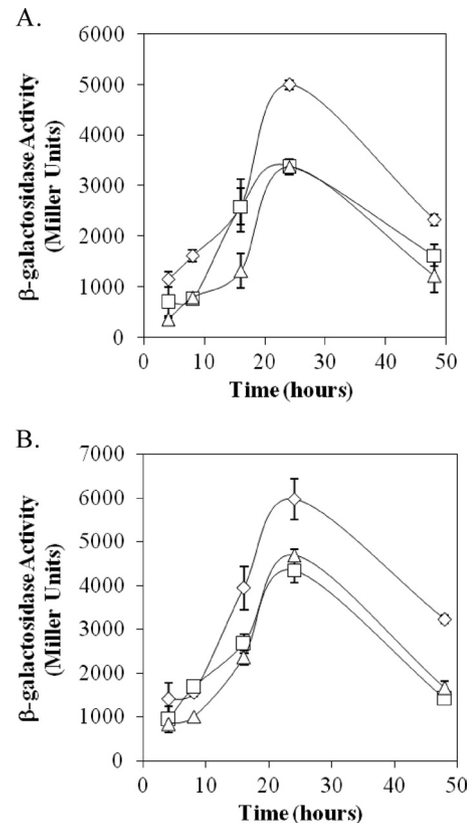


FIG. 5. Expression of *rpoS-lacZ* in strain DF41 and its derivatives. Strains used were as follows: *Pseudomonas* sp. strain DF41 (diamonds), DF41 *relA* mutant (squares), and DF41 *relA spoT* mutant (triangles). All strains harbored the *rpoS-lacZ* reporter plasmid pRPOS-*lacZ* together with empty vector pME6010 (A) or pME-*rpoS* (B). The results shown in panels A and B are from the same experiment. Note that the presence of pME-*rpoS* in the SR mutants (B) restored *rpoS* activity to a level close to that of DF41(pME6010) (A).

found to be under the control of GacS (2). To determine the impact of the SR on the transcription of this sigma factor, the  $\beta$ -galactosidase activity of an *rpoS-lacZ* transcriptional fusion was monitored in strain DF41 and its derivatives. At 24 h, *rpoS* transcription was reduced almost 1.5-fold in the ppGpp-deficient strains compared to the wild type (Fig. 5A). To determine the effect of constitutively expressed *rpoS* on transcription of this sigma factor, pME-*rpoS* was mobilized into the DF41, DF41 *relA*, and DF41 *relA spoT* strains, all of which carry the *rpoS-lacZ* transcriptional fusion plasmid. As shown in Fig. 5B, the presence of pME-*rpoS* in the SR mutants elevated *rpoS-lacZ* expression to a level close to that of DF41 carrying the empty vector (Fig. 5A). Conversely, when pME-*rpoS* was introduced into the DF41-1278, 1278 *relA*, and 1278 *relA spoT* strains, no change in NRPS gene transcription was observed (data not shown). To see if constitutively expressed *rpoS* would alter phenotypic traits displayed by the SR mutants, the DF41 *relA*(pME-*rpoS*) and DF41 *relA spoT*(pME-*rpoS*) strains were analyzed for LP production and antifungal and protease activities. No difference in LP production and antifungal activity was observed (data not shown); however, protease activity exhibited by the ppGpp-deficient strains was restored to wild-type levels upon the addition of *rpoS* in *trans* (Table 3).

## DISCUSSION

The SR is a global regulatory mechanism that enables bacteria to alter a broad range of physiological traits, including growth rate, motility, persistence, biofilm formation, virulence, and secondary metabolite production, in response to nutrient limitation (27). In light of this, we were interested to learn how the SR affects the antifungal activity of DF41. This is particularly important because when DF41 is used as a biocontrol agent against *Sclerotinia* stem rot of canola, it is applied as a foliar spray (2, 32). The aerial surfaces of the plant represent a harsh environment for bacteria, as they must endure exposure to high and low temperatures, UV radiation, and desiccation (23). Additionally, nutrients are scarce, so bacteria on the plant surface are expected to undergo the SR (23).

Characterization of DF41 *relA* and *relA spoT* SR mutants revealed a lack of (p)ppGpp production (Fig. 1) and enhanced inhibition of *S. sclerotiorum* (Table 2), indicating that the SR negatively regulates strain DF41 biocontrol. We next addressed whether the SR affects LP expression. HPLC analysis of culture extracts showed that LP levels were elevated between 1.5- and 2.0-fold in the DF41 *relA* and *relA spoT* mutants compared to those in strain DF41 (Fig. 3). Moreover, at 12 h, transcription of the NRPS biosynthetic genes was increased over 5 times in the SR mutant background relative to that in the wild type (Fig. 2). These findings suggest that in the absence of (p)ppGpp, the NRPS genes are induced earlier, leading to enhanced LP production. A similar finding was reported for *Streptomyces clavuligerus*, in which an *relA* mutant produced elevated levels of the antibiotics clavulanic acid and cephamycin C (11). Moreover, transcription of the clavulanic acid and cephamycin C biosynthetic genes was increased substantially in the *relA*-deficient strain (11). With respect to SR control over antibiotic production, *Streptomyces* species are probably the most well-studied group of bacteria (4, 11, 15, 18, 19). Depending on the strain of *Streptomyces* in question, the antibiotics produced, and the growth media employed, the SR has been found to either positively (4, 15, 18, 19) or negatively (11) regulate antibiotic production. Hence, for *Streptomyces* spp., the SR appears to have a variable effect on antibiotic production. For biocontrol strains of *Pseudomonas*, little is known about SR control over secondary metabolite production and fungal antagonism. One exception, however, is *Pseudomonas* sp. strain MIS38. Therein, a transposon insertion in *spoT* resulted in elevated (p)ppGpp levels and reduced production of the LP arthrfactin (35), consistent with the inverse relationship between (p)ppGpp levels and LP expression observed for strain DF41.

In pseudomonads, regulation of biocontrol factors is governed by a complex network that functions at both the transcriptional and posttranscriptional levels. At the top of the hierarchy, the GacS/GacA two-component signal transduction system is essential for biological control in *Pseudomonas* spp. (12, 13). Mutation of either component typically leads to a complete loss of antagonistic activity (13), which was found to be the case for strain DF41 (2). Because the SR inhibits expression of factors that are also under Gac control, we were interested in determining whether a connection exists between the two systems in strain DF41. In this study, we observed a 1.7-fold decrease in *relA* expression in the *gacS*-negative back-

ground (Fig. 5). (p)ppGpp analysis also revealed a dramatic reduction in the level of this nucleotide in the *gacS* mutant, which was increased upon the addition of *gacS* in *trans* (Fig. 1B). These findings indicate that the expression of *relA* is influenced by the Gac system, although the molecular mechanism(s) underlying this regulation has yet to be uncovered. In other pseudomonads, the Rsm system forms part of the Gac regulatory cascade (12). After Gac activation, a series of regulatory RNAs are produced that antagonize the effects of RsmA-like proteins, which bind to and block translation of biocontrol mRNAs (12). The only Rsm component that has been identified in strain DF41 thus far is a homolog of the regulatory RNA RsmZ (T. R. de Kievit and C. Berry, unpublished data). The presence of RsmZ suggests that a cognate repressor protein(s) and additional regulatory RNAs likely exist, some or all of which may play a role in *relA* regulation.

During the SR, RNAP binding to (p)ppGpp and additional effectors, such as DksA, leads to activation and repression of different subsets of genes (27). For genes that are activated, increased expression is believed to occur by both direct and indirect mechanisms (see reference 27 and references therein). The direct effect involves RNAP in complex with (p)ppGpp increasing transcription from a given promoter. Indirect effects arise because (p)ppGpp facilitates RNAP binding to alternative sigma factors, leading to induction of these regulons. In this study, the sigma factor gene *rpoS* was found to be upregulated during the SR, while the NRPS genes were repressed. To better understand how the SR affects transcription of *rpoS* and the NRPS genes, we expressed *rpoS* constitutively from the kanamycin promoter in the SR mutants. The presence of plasmid pME-*rpoS* in the *relA* and *relA spoT* mutants increased *rpoS* transcription to near wild-type levels (Fig. 5). In strain DF41, *rpoS* is positively autoregulated, as evidenced by the increased transcription brought about by pME-*rpoS* (Fig. 5B). It is possible that in the SR mutants, the elevated level of plasmid-encoded RpoS enabled it to bind RNAP in the absence of (p)ppGpp, resulting in increased *rpoS* transcription. On the other hand, addition of *rpoS* had no effect on NRPS gene transcription. In this situation, (p)ppGpp may affect transcription from these promoters directly by decreasing NRPS transcription. Alternatively, the SR may exert its effects indirectly, through an as yet unidentified regulator of NRPS gene expression. Clearly, a great deal of work remains in order for us to understand exactly how the SR affects expression of these and other genes in strain DF41. When we examined whether the presence of *rpoS* in *trans* altered the SR mutant phenotype, the only trait that was affected was protease activity, which was restored to wild-type levels (Table 3). A protease-encoding gene(s) has not yet been identified in strain DF41; however, the results of the present study suggest that this gene(s) is under RpoS control.

In summary, we demonstrated that the SR negatively affects the antifungal activity of strain DF41 *in vitro*. Whether SR mutants would demonstrate enhanced biocontrol *in planta* has not been established. When an organism encounters a stress, several mechanisms may be induced to mount an appropriate response. For instance, bacteria frequently produce long-chain polymers of phosphate residues, termed poly(P)s, when starved for nutrients (29). These molecules are synthesized by polyphosphate kinase, the product of the *ppk* gene,

while their hydrolysis to  $P_i$  residues depends upon the *ppx*-encoded exopolyphosphatase (29). It was reported that in *Streptomyces lividans*, expression of the antibiotics actinorhodin and undecylprodigiosin was enhanced in the *ppk* mutant under phosphate-limiting conditions, presumably as a result of reduced endogenous  $P_i$  levels (6). Furthermore,  $P_i$ -mediated inhibition is believed to occur independently of (p)ppGpp, as *relA* transcription in the poly(P)-deficient strain was similar to that in the wild type (6). Taken together, these findings suggest that the regulation of antibiotic production is complex and requires multiple pathways to respond appropriately to the prevailing conditions. Synthesis of antifungal factors, including LP and protease, is an energetically costly process. When nutrients are scarce, the SR enables strain DF41 to shift resources from secondary metabolite production to activities required for coping with starvation. Because biocontrol bacteria are often forced to survive under nutrient-depleted conditions, it is essential to understand the impact of the SR on secondary metabolism, as this directly impacts antagonism.

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