Hfq and sRNA 179 Inhibit Expression of the Pseudomonas aeruginosa cAMP-Vfr and Type III Secretion Regulons

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ABSTRACT Pseudomonas aeruginosa is a Gram-negative opportunistic pathogen causing skin and soft tissue, respiratory, and bloodstream infections. The type III secretion system (T3SS) is one important virulence factor. Production of the T3SS is controlled by ExsA, a transcription factor that activates expression of the entire T3SS regulon. Global regulators including Vfr, RsmA, and Hfq also contribute to regulation of the T3SS. Vfr is a cAMP-responsive transcription factor that activates exsA transcription. RsmA, an RNA-binding protein, inversely controls expression of the T3SS and the type VI secretion system (T6SS). Hfq is an RNA chaperone that functions by stabilizing small noncoding RNAs (sRNAs) and/or facilitating base pairing between sRNAs and mRNA targets. A previous study identified sRNA 1061, which directly targets the exsA mRNA and likely inhibits ExsA synthesis. In this study, we screened an sRNA expression library and identified sRNA 179 as an Hfq-dependent inhibitor of T3SS gene expression. Further characterization revealed that sRNA 179 inhibits the synthesis of both ExsA and Vfr. The previous finding that RsmA stimulates ExsA and Vfr synthesis suggested that sRNA 179 impacts the Gac/Rsm system. Consistent with that idea, the inhibitory activity of sRNA 179 is suppressed in a mutant lacking rsmY and rsmZ, and sRNA 179 expression stimulates rsmY transcription. RsmY and RsmZ are small noncoding RNAs that sequester RsmA from target mRNAs. Our combined findings show that Hfq and sRNA 179 indirectly regulate ExsA and Vfr synthesis by reducing the available pool of RsmA, leading to reduced expression of the T3SS and cAMP-Vfr regulons.

IMPORTANCE Control of gene expression by small noncoding RNA (sRNA) is well documented but underappreciated. Deep sequencing of mRNA preparations from Pseudomonas aeruginosa suggests that >500 sRNAs are generated. Few of those sRNAs have defined roles in gene expression. To address that knowledge gap, we constructed an sRNA expression library and identified sRNA 179 as a regulator of the type III secretion system (T3SS) and the cAMP-Vfr regulons. The T3SS- and cAMP-Vfr-controlled genes are critical virulence factors. Increased understanding of the signals and regulatory mechanisms that control these important factors will enhance our understanding of disease progression and reveal potential approaches for therapeutic intervention.

KEYWORDS Pseudomonas aeruginosa, sRNA, Hfq, type III secretion, Vfr, ExsA, RsmA

Pseudomonas aeruginosa is an important Gram-negative nosocomial pathogen that can cause skin and soft tissue, urinary tract, lung, and bloodstream infections (1). Many P. aeruginosa virulence factors are directly controlled by the cAMP-Vfr signaling
The CVS consists of the CyaA and CyaB adenylate cyclases, a cAMP degrading phosphodiesterase, and the transcription factor Vfr (virulence factor regulator). Vfr directly responds to increased intracellular cAMP pools to activate expression of the CVS regulon, which includes type IV pili, a type II secretion system, secreted factors (e.g., exotoxin A and protease IV), and the type III secretion system (T3SS). The T3SS is an important virulence factor that contributes to the pathogenesis of human and animal infections. The T3SS is a needle-like apparatus used to translocate at least four effectors with antihost properties into eukaryotic target cells. The combined activities of the effector proteins promote phagocytic evasion and systemic spread of the microorganism.

ExsA autoregulates its own transcription through the \( P_{\text{exsC}} \) promoter to generate a polycistrionic mRNA encoding \( \text{exsC}, \text{exsE}, \text{exsB} \), and \( \text{exsA} \) (Fig. 1). A 297-bp intergenic region separates \( \text{exsB} \) and \( \text{exsA} \) and contains a Vfr-dependent promoter \( P_{\text{exsA}} \) dedicated to \( \text{exsA} \) transcription (Fig. 1). Several other factors contribute to \( \text{exsA} \) transcription, including PsrA, which stimulates transcription from the \( P_{\text{exsC}} \) promoter, and MvaT/MvaU, VqsM, and Fis, which modulate \( P_{\text{exsA}} \) promoter activity.

Once transcribed, translation of ExsA appears to be inefficient, possibly owing to an inhibitory structure in the mRNA that reduces the efficiency of ribosomal recruitment.

The RNA helicase DeaD is required for ExsA synthesis in vivo and stimulates ExsA translation in vitro (22). DeaD likely functions by denaturing an inhibitory structure in the \( \text{exsA} \) mRNA leader region to enhance ribosomal recruitment (22). RsmA, a member of the \( \text{Escherichia coli} \) CsrA family of RNA-binding proteins, can have both positive and negative effects on gene expression through direct interactions with target mRNAs (23). RsmA has positive effects on the synthesis of Vfr and ExsA through mechanisms that remain to be defined (Fig. 1) (24). The activity of RsmA is controlled by RsmY and RsmZ, small noncoding RNAs (sRNAs) with multiple RsmA binding sites. RsmY and RsmZ function by sequestering RsmA from target mRNAs (25). Transcription of \( rsmY \) and \( rsmZ \) is controlled by the GacAS two-component system (23). Increased levels of RsmY and RsmZ lead to RsmA sequestration and reduced T3SS gene expression.

Host factor for bacteriophage Qi\(\beta\) RNA replication (Hfq) was identified in \( E. coli \) for its essential role in bacteriophage replication (26). Hfq is an RNA-binding protein that functions as an RNA chaperone to stabilize sRNAs and/or facilitate imperfect base pairing between sRNAs and mRNA targets (27). The actions of Hfq and sRNA can affect translation and/or mRNA stability and result in negative or positive effects on gene expression. Hfq regulates \( \sim5\% \) of the \( P. aeruginosa \) genome, including genes involved
in stress responses, metabolism, and virulence (28–30). A *P. aeruginosa* hfq mutant has reduced alginate production, lipopolysaccharide (LPS), and quorum sensing and increased type III secreted products (29, 31). The mechanism of T3SS control by Hfq is poorly understood (31). sRNA PA0161 was recently identified using the GRIL-seq method and was proposed to interact with the *exsA* mRNA (32). In this study, we report that Hfq works with sRNA 0161 to directly inhibit *ExsA* synthesis and with sRNA 179 to indirectly regulate both *exsA* transcription and *ExsA* synthesis.

**RESULTS**

T3SS gene expression is derepressed in the absence of Hfq. Disruption of *hfq* in *E. coli* results in pleiotropic phenotypes resulting in delayed growth, decreased negative supercoiling, increased cell size, and sensitivity to UV light (33, 34). Previous studies found that *P. aeruginosa* hfq insertion and deletion mutants in strain PAO1 have a growth defect when cultured in LB medium (28, 35). In the current study, we constructed an Δ*hfq* deletion mutant (residues Δ2 to Δ82) in *P. aeruginosa* strain PA103 and observed a growth defect. In our subsequent studies with the PA103 Δ*hfq* strain, emergence of growth suppressor mutants was common. A recent study by Hill et al. also noted the emergence of growth suppressor mutants with the PAO1 *hfq* deletion strain (35), and sequencing of several of the suppressor mutants demonstrated involvement of multiple genetic loci. Although the nature of the PA103 Δ*hfq* suppressor mutants was not determined in the current study, each of the Δ*hfq* mutant phenotypes reported below could be complemented by expressing *hfq* in trans from a plasmid. For this reason, we believe that the growth phenotype of the suppressor mutants is unrelated to Hfq effects on expression of the Vfr and T3SS regulons as described below.

Previous studies found that Hfq contributes to *P. aeruginosa* T3SS gene expression (29, 31), but a regulatory mechanism was not described. To examine the contribution of Hfq to T3SS gene expression, a *P_{exsD}-lacZ* transcriptional reporter was integrated into the *ΦCTX* phage attachment site of wild-type (WT) strain PA103 and the Δ*hfq* mutant. The *P_{exsD}-lacZ* reporter is a measure of *ExsA*-dependent transcription and demonstrates strong induction when WT cells are cultured under inducing conditions for T3SS gene expression (low Ca²⁺, presence of EGTA) (Fig. 2) (36). *P_{exsD}-lacZ* reporter activity...
increased significantly in the Δhfq mutant under both noninducing (high Ca²⁺, no EGTA) and inducing conditions (Fig. 2). Corresponding immunoblots showed an increase in ExsA protein levels under both noninducing and inducing conditions in the Δhfq mutant compared to the wild type. Whereas T3SS gene expression was derepressed in the hfq mutant, expression of Hfq from a plasmid inhibited PexsD-lacZ reporter activity as well as ExoU and ExsA production in both the WT and hfq mutant backgrounds (Fig. 2). Hfq protein was detectable in the WT strain, absent in the hfq mutant, and significantly elevated in strains carrying the Hfq expression plasmid (Fig. 2). These findings demonstrate that Hfq has a negative effect on T3SS gene expression.

**Hfq regulates T3SS-mediated cytotoxicity.** To verify that Hfq negatively regulates T3SS gene expression, we infected Chinese hamster ovary (CHO) cells with strains (wild type and Δhfq) carrying either a vector control or the Hfq expression vector. Previous studies have shown that coculturing CHO cells with *P. aeruginosa* strain PA103 results in acute cytotoxicity that is T3SS-dependent (37). In addition, the kinetics are advanced in strains derepressed for T3SS gene expression (e.g., an exsD mutant) (36). CHO cell lysis was measured by lactate dehydrogenase release following 15, 30, 45, and 60 min of coculture with *P. aeruginosa* (Fig. 3A). Relative to the parental strain, cytotoxicity was reduced with the exsA mutant, encoding a positive regulator of T3SS gene expression (Fig. 3A and B). In the absence of either exsD or hfq, cytotoxicity displayed enhanced kinetics, consistent with depressed T3SS gene expression. Expression of Hfq in the wild-type background resulted in an ~4-fold reduction in cytotoxicity following 30 min of coculture relative to the vector control (Fig. 3A and B). Similarly, Hfq expression in the Δhfq deletion strain also reduced cytotoxicity compared to the vector control, but not to the same levels observed in the wild-type background (Fig. 3B). These results suggest that Hfq negatively regulates T3SS-mediated cytotoxicity.
Identification of sRNAs that regulate type III secretion. Hfq can inhibit protein synthesis by directly interacting with mRNA targets alone or by facilitating the base pairing of sRNAs with mRNA targets (27). To identify sRNAs that regulate the T3SS, we cloned and expressed a library of sRNAs in *P. aeruginosa* strain PAK and screened for effects on T3SS gene expression using the *P*nox-DlacZ transcriptional reporter. Gomez-Lozano et al. previously identified over 500 novel sRNAs in *P. aeruginosa* using three different methods to prepare transcript libraries (38). From this data set, we selected 240 sRNAs (referred to as Pants [P. aeruginosa novel transcripts]) in the original paper) that met two criteria, >200 nucleotides in length and known to originate from either the Watson or Crick strand of the genome (it has not been determined for all of the sRNAs). Each of the selected sRNAs was placed under the transcriptional control of an arabinose-inducible promoter. Six of the sRNAs (sRNA 18, 179, 182, 183, 214, and 351) inhibited *P*nox-DlacZ reporter activity greater than 2-fold in strain PAK (Fig. 4A and B).

As a secondary screen, the six sRNAs identified using strain PAK were tested for activity in strains PA14 and PA103 bearing the *P*nox-DlacZ reporter (Fig. 4B). We also included sRNA PA0161 as a positive control. sRNA PA0161 (not included in our original library) was recently identified using GRIL-Seq as an inhibitor of T3SS gene expression (32). Although sRNAs 18, 182, 183, and 214 possess regulatory activity in strain PAK, they had no effect on *P*nox-DlacZ reporter activity when expressed in strains PA14 and PA103 (Fig. 4B). sRNA 351 inhibited *P*nox-DlacZ reporter activity in both the PAK and PA14 backgrounds but was inactive in strain PA103. The Hfq expression vector inhibited *P*nox-DlacZ reporter activity in all three genetic back-

![Graph A](image1.png)

**FIG 4** Identification of small noncoding RNAs that alter T3SS gene expression. (A) *P. aeruginosa* strain PAK carrying a *P*nox-DlacZ transcriptional reporter was transformed with a vector control and the sRNA expression vectors indicated in Table S1 (note that only 237 of the 512 putative sRNAs were tested). Strains were cultured under inducing (+EGTA) conditions for T3SS gene expression in medium supplemented with 0.2% arabinose to induce sRNA expression and assayed for *P*nox-DlacZ reporter activity. The reporter values represent the activity of each sRNA relative to the vector control. (B) sRNAs that inhibited or stimulated *P*nox-DlacZ reporter activity greater than 2-fold (indicated by red lines in panel A) were selected for further analyses. Expression vectors for sRNA 18, 179, 182, 183, 214, and 483 were introduced into strains PA14 and PA103 carrying the *P*nox-DlacZ reporter and assayed as described in panel A for reporter activity. sRNA 0161, previously shown to inhibit T3SS gene expression (32), and the Hfq expression vector were included as controls. The reported values represent the activity of each sRNA or pHfq relative to each parental strain carrying the vector control. The reported values with the standard error represent the average of at least three experiments. *, *P* < 0.01.
Since only Hfq, sRNA 0161, and sRNA 179 inhibited T3SS gene expression in all three genetic backgrounds, the remainder of this study focused on those factors. sRNA 179 and Hfq function together to inhibit ExsA translation. To determine whether the inhibitory activities of sRNAs 0161 and 179 are Hfq-dependent, both sRNAs were expressed in P. aeruginosa strain PA103 and the Δhfq mutant. Whereas expression of either RNA in WT cells resulted in strong inhibition of P_{exsD-lacZ} reporter activity, ExsA production, and ExoU secretion, the inhibitory activities of sRNAs 0161 and 179 were largely suppressed in the Δhfq mutant (Fig. 5A). ExsA protein levels were reduced by...
~25% upon sRNA 0161 and 179 expression in the Δhfq mutant relative to the vector control. In contrast, ExsA protein was reduced by >90% upon sRNA 0161 and 179 expression in the WT background. ExoU protein levels followed a similar trend. These data are consistent with sRNAs 0161 and 179, both functioning as typical sRNAs that depend upon Hfq for facilitated base pairing with target mRNA(s).

The data presented thus far show that sRNAs 0161 and 179 expressed from a plasmid possess inhibitory activity. To examine the effects of sRNA 0161 and 179 expressed from their native chromosomal promoters, we constructed deletion mutants for each and a double mutant lacking both sRNA 0161 and 179. Deletion mutants lacking hfq, sRNA 0161, and both sRNAs (0161 and 179) carrying a vector control (pJN105) demonstrated significant depression of $P_{exsD}$$\cdot$$\text{lacZ}$ reporter activity (Fig. 5B). Although reporter activity was consistently elevated in the single Δ179 mutant, that difference did not meet a statistical test of significance. To determine whether sRNA 0161 and 179 account for all of the Hfq-dependent activity, the Hfq expression plasmid was introduced into each of the sRNA mutant backgrounds. Hfq expression resulted in significant inhibition of $P_{exsD}$$\cdot$$\text{lacZ}$ reporter activity in the single and double sRNA 0161 and 179 mutants (Fig. 5B). Finally, complementation analyses demonstrate that plasmid-expressed 0161 and 179 are sufficient to repress $P_{exsD}$$\cdot$$\text{lacZ}$ reporter activity in their cognate mutant backgrounds and in the double 0161 and 179 mutant (Fig. 5C). These combined findings demonstrate that (i) Hfq functions together with sRNAs 0161 and 179 to inhibit T3SS gene expression, (ii) sRNA 0161 and 179 activities are independent of one another, and (iii) Hfq also inhibits T3SS gene expression in a manner that does not rely upon sRNA 0161 or 179. Such activity could reflect sRNA-independent activity or involvement of an additional sRNA(s).

**Hfq impacts ExsA synthesis at the posttranscriptional level.** Based on the previous observation that T3SS genes demonstrate increased mRNA levels in an hfq mutant (31), we hypothesized that Hfq, sRNA 0161, and/or sRNA 179, either directly or indirectly, regulate ExsA synthesis. Because ExsA autoregulates its own transcription, we established an experimental system to uncouple exsA transcription from ExsA synthesis. The native 101-nt upstream untranslated region (corresponding to the exsA transcription start site from the $P_{exsA}$ promoter) and coding sequence for exsA were placed under the transcriptional control of a rhamnose-inducible promoter ($P_{rha}$) and integrated into the chromosome at the Tn7 transposon insertion site of an ΔexsA strain. Titration of rhamnose identified conditions (0.005%) where $P_{exsD}$$\cdot$$\text{lacZ}$ reporter activity and ExsA expression levels were similar to the WT PA103 strain (Fig. 6A; see Fig. S2 in the supplemental material). Expression of the ExsA-dependent $P_{exsD}$$\cdot$$\text{lacZ}$ reporter was entirely dependent upon addition of rhamnose to the culture medium and remained responsive to calcium chelation by EGTA (Fig. 6A), thus validating the system.

The Tn7-rhamnose system was then used to determine whether Hfq, sRNA 0161, and/or sRNA 179 affect ExsA synthesis at the posttranscriptional level. Expression of Hfq, sRNA 0161, and sRNA 179 each resulted in strong inhibition of $P_{exsD}$$\cdot$$\text{lacZ}$ reporter activity and ExsA protein levels (Fig. 6B, lanes 1 to 4), and the activity of the sRNAs was entirely dependent upon Hfq (lanes 5 to 8). To verify that the reduction in ExsA synthesis was unrelated to changes in exsA mRNA levels, qRT-PCR for exsA was performed on RNA harvested from the samples shown in lanes 1 to 4. Whereas ExsA protein levels were reduced ~4-fold in the Hfq, 0161, and 179 expression strains relative to the vector control, exsA mRNA levels were only slightly reduced (~25%) (Fig. 6C). Some degradation of the exsA mRNA was expected owing to the lack of ribosomal protection that occurs during active translation. These findings are consistent with Hfq and sRNAs 0161 and 179 controlling ExsA synthesis at a posttranscriptional level.

**sRNA 0161 and the exsA mRNA interact with each other and Hfq.** Our finding that sRNA 0161 inhibits ExsA synthesis and the previous demonstration of a specific interaction between sRNA 0161 and the exsA mRNA by GRIL-seq (32) (Fig. 6B) was suggestive of a direct effect on ExsA translation. To measure the interaction in vitro,
sRNA 0161 was in vitro synthesized, radiolabeled at the 5’ end, and incubated with an unlabeled portion of the exsA mRNA leader region prior to electrophoresis on a nondenaturing gel. The exsA probe (exsA 101) consists of the native 101 untranslated region and 20 bp downstream of the start codon. An sRNA 0161-exsA complex was observed when 20 nM unlabeled exsA mRNA leader region was added to the reaction (Fig. S2). We were unable to detect an interaction between sRNA 179 and the exsA mRNA using the same approach. We also assayed for Hfq binding and found that Hfq interact with sRNA 0161 and/or the exsA mRNA (Fig. S3). These data are consistent with
the previous GRIL-Seq findings of Zhang et al. (32) and suggest that Hfq directly interacts with sRNA 0161 and the 5' untranscribed region (UTR) of exsA to reduce translation. Furthermore, the inability to detect an interaction between sRNA 179 and exsA suggested that the mechanism of inhibition by sRNA 179 may be indirect and distinct from the action of sRNA 0161.

Hfq inhibits the Vfr-cAMP signaling system. We noted that Hfq appeared more effective at inhibiting \( P_{\text{exsA}}-\text{lacZ} \) reporter activity in the WT background (Fig. 2) compared to the exsA mutant carrying the rhamnose-inducible \( exsA \) allele (Fig. 6B). One difference between these strains is that \( exsA \) transcription is no longer controlled by the Vfr-dependent \( P_{\text{exsA}} \) promoter. This raised the possibility that Hfq also impacts the cAMP-Vfr system. Indeed, \( P_{\text{exsA}}-\text{lacZ} \) reporter activity and Vfr protein levels by immunoblot. \( *, P < 0.001 \). (C) Twitching motility zones for \( P. aeruginosa \) strains PA103, PAK, and PA14 carrying a vector control (pJN105) or an Hfq expression vector. \( *, P < 0.05 \).

FIG 7 Hfq and sRNA 179 inhibit the cAMP-Vfr signaling system. (A) WT PA103 or an hfq mutant transformed with the indicated expression vectors were assayed for Vfr-dependent \( P_{\text{exsA}}-\text{lacZ} \) transcriptional reporter activity. WT PA103 carrying a promoterless \( \text{lacZ} \) transcriptional reporter was included as a control for background activity. \( *, P < 0.05 \). (B) The indicated strains were transformed with a vector control or an Hfq expression vector and assayed for \( vfr^{-}\text{lacZ} \) translational reporter activity and Vfr protein levels by immunoblot. \( *, P < 0.001 \). (C) Twitching motility zones for \( P. aeruginosa \) strains PA103, PAK, and PA14 carrying a vector control (pJN105) or an Hfq expression vector. \( *, P < 0.05 \).
significant inhibition of twitching motility in strains PA103, PAK, and PA14 (Fig. 7D). These combined data suggest that Hfq/sRNAs have two effects on T3SS gene expression, inhibition of Vfr synthesis resulting in reduced $P_{\text{exsA}}$ promoter activity and inhibition of ExsA synthesis.

**sRNA 179 requires RsmY and RsmZ to control T3SS gene expression.** Previous studies have shown that the small RNA-binding protein RsmA is required for the synthesis of both Vfr and ExsA (39). Our finding that Hfq and sRNA 179 overexpression inhibits Vfr and ExsA synthesis suggested involvement of the Rsm system. RsmA availability is controlled by the small noncoding RNAs RsmY and RsmZ (40). To examine whether Hfq and sRNA 179 function through the Rsm system to control T3SS gene expression, we expressed Hfq, sRNA 0161, and sRNA 179 in $\Delta rsmY$, $\Delta rsmZ$, and $\Delta rsmYZ$ mutants and measured $P_{\text{exsD}}$-lacZ reporter activity. Although no significant differences were observed between the WT strain and the $\Delta rsmY$ and $\Delta rsmZ$ single mutants, $P_{\text{exsD}}$-lacZ reporter activity was significantly derepressed in the $\Delta rsmYZ$ mutant owing to increased RsmA availability (Fig. 8A). Overexpression of Hfq, sRNA 1061, and sRNA 179 resulted in strong inhibition of $P_{\text{exsD}}$-lacZ reporter activity when expressed in each of the strains tested, with one exception; sRNA 179 lacked activity in the $\Delta rsmYZ$ mutant.

We next tested the hypothesis that sRNA 179 stimulates rsmY and/or rsmZ transcription. Both transcriptional reporter and reverse transcription-quantitative PCR (qRT-PCR) data demonstrate that sRNA 179 expression results in a significant increase in $P_{\text{rsmY}}$-lacZ reporter and RsmY RNA levels (Fig. 8B to D). The effect of sRNA 179 is specific, as reporter activity and RsmY levels were unaffected by Hfq or sRNA 0161 expression, and Hfq, sRNA 0161, and sRNA 179 had no effect on $P_{\text{rsmZ}}$-lacZ reporter and RsmZ RNA levels. These combined findings demonstrate that sRNA 179 has indirect effects on both exsA transcription and ExsA translation through modulation of RsmY levels, thus impacting RsmA availability, which itself has positive effects on Vfr and ExsA synthesis.

**DISCUSSION**

The previous finding that Hfq influences *P. aeruginosa* T3SS gene expression led us to investigate the mechanism (29, 30). Potential roles for Hfq include sRNA and mRNA stability control and mRNA translational control with sRNA-assisted targeting (41). We hypothesized the involvement of an sRNA(s) in the regulation of the T3SS. Screening an sRNA expression library resulted in the identification of sRNA 179 as an Hfq-dependent inhibitor of both the T3SS and CVS regulons. sRNA 179 is 134 nt long and located downstream of acp1, a putative acyl carrier protein. The reported 5’ end of sRNA 179 immediately follows the acp1 stop codon (38) and may be generated by a processing event. sRNA 179 lacks an intact open reading frame containing a stop codon. The finding that sRNA 179 activity is Hfq-dependent supports our model that the RNA is the relevant gene product and is consistent with a recent paper showing that sRNA 179 coprecipitates with Hfq (42). sRNA 179 indirectly inhibits ExsA and Vfr synthesis at the posttranscriptional level and likely works through the Rsm system. Data supporting this include (i) the lack of inhibitory activity when sRNA 179 is overexpressed in a $\Delta rsmYZ$ mutant (Fig. 8A), (ii) increased $P_{\text{rsmY}}$-lacZ transcriptional reporter activity upon sRNA 179 overexpression (Fig. 8B), and (iii) increased RsmY RNA levels upon sRNA 179 overexpression (Fig. 8C). Our working model is that sRNA 179-mediated increase in RsmY expression titrates RsmA from some target mRNAs, leading to reduced T3SS and CVS expression, as both ExsA and Vfr synthesis are dependent upon RsmA (Fig. 1) (39). The RsmA regulon consists of $\sim$500 target genes and includes the T6SS (43). To examine whether sRNA 179 has a general effect on the RsmA regulon, we immunoblotted culture supernatants for the T6SS secretion substrate Hcp1 upon sRNA 179 overexpression but observed no change relative to the parental strain (data not shown). This was not entirely unexpected, as we previously demonstrated that T3SS gene expression is more sensitive to changes in RsmA availability relative to T6SS gene expression (39).

sRNA 179 inhibits the T3SS when overexpressed in the WT, $\Delta rsmY$, and $\Delta rsmZ$ single
mutants but lacks activity when expressed in the ΔrsmYZ mutant (Fig. 8A). These data suggest involvement of both rsmY and rsmZ in T3SS gene expression by sRNA 179 and may reflect the redundant function that RsmY and RsmZ play in RsmA sequestration. The homeostatic relationship between RsmA availability and RsmY/RsmZ-mediated seque-

FIG 8 sRNA 179 modulates the Rsm system by increasing RsmY transcription. (A) The indicated strains carrying a $P_{\text{exsD-lacZ}}$ reporter were transformed with either the vector control (pJN), the Hfq expression vector, or sRNA 0161 and 179 expression vectors. Strains were cultured under inducing conditions for T3SS gene expression supplemented with 0.05% or 0.2% arabinose to induce Hfq or sRNA expression, respectively, and assayed for $P_{\text{exsD-lacZ}}$ reporter activity as reported in Miller units. The reported values with the standard error represent the average of at least three experiments. *, $P < 0.05$. (B) WT carrying either a $P_{\text{rsmY-lacZ}}$ or $P_{\text{rsmZ-lacZ}}$ transcriptional reporter and the indicted expression vectors were assayed for reporter activity as in panel A. *, $P < 0.005$. (C and D) Comparison of RsmY (C) and RsmZ (D) levels in RNA samples harvested from the indicated strains. RNA samples were quantified by qRT-PCR and normalized to the $\text{rimM}$ housekeeping gene. The reported values are relative to each strain carrying the vector control.
tration of RsmA is a complicating factor. Because RsmA has a positive effect on rsmY and rsmZ transcription (39), deletion of either may result in a compensatory effect wherein reduced sequestration of RsmA stimulates transcription of the remaining gene. Transcription of both rsmY and rsmZ is directly controlled by the GacAS two-component regulatory system (44). It is unclear how sRNA 179 stimulates rsmY transcription, but modulation of GacAS signaling seems unlikely since rsmZ transcription is unaffected by sRNA 179. In addition to GacAS, other factors, including HptB, MvaT, and MgtE, also influence rsmYZ transcription (45–47). HptB is a promising candidate, as both HptB and sRNA 179 effect rsmY transcription only. While it is clear that rsmYZ are required for sRNA 179 activity, they may not be sufficient. Our finding that F_{rsmZ-lacZ} reporter activity and RsmZ levels are unchanged when sRNA 179 is overexpressed in the rsmY background (Fig. 8B and D) suggest additional mechanisms of inhibition by sRNA 179.

In addition to sRNA 179, sRNA 0161 was recently identified with GRIL-seq as another inhibitor of the T3SS (32), and we confirmed that sRNA 0161 is also Hfq-dependent (Fig. 4A). The GRIL-seq data suggest that sRNA 0161 directly targets the exsA leader region through imperfect base pairing and is consistent with our data showing that sRNA 0161 inhibits ExsA synthesis at the posttranscriptional level (Fig. 5B and C). In addition, Hfq coprecipitates with the exsA leader region (42). The simplest model to account for the inhibitory activity of sRNA 0161 is occlusion of the ribosome binding site. Alternative mechanisms may involve RsmA and DeaD, both positive regulators of ExsA synthesis (22, 39). Rather than blocking ribosome access, sRNA 0161 may instead prevent RsmA and/or DeaD from binding to the exsA leader region. Another possibility is that RsmA and/or DeaD activate by displacing/preventing sRNA 0161 from base pairing with the exsA leader region. Testing these models will be the subject of future studies.

sRNA 0161 and 179 function independently from one another, and either is sufficient to inhibit T3SS gene expression (Fig. 4C). The primary implication is that together they provide both direct (through the activity of sRNA 1061) and indirect (through sRNA 179) Hfq-dependent mechanisms to inhibit ExsA synthesis. Transcription and synthesis of exsA and ExsA are major inputs for upstream regulatory factors (17). Although seemingly redundant, sRNA 0161 and sRNA 179 may differentially coordinate and/or respond to distinct signals that influence T3SS expression. Future studies to understand transcriptional control of sRNA 0161 and sRNA 179 will be important.

Regulatory control of the T3SS by Hfq likely extends beyond the mechanisms described here. Expression of Hfq in an sRNA 0161 179 double mutant still resulted in significant inhibition of T3SS gene expression (Fig. 4B). This may reflect sRNA-independent activity or involvement of an additional sRNA(s) in control of the T3SS. There are over 500 predicted sRNAs in P. aeruginosa (38), only a small fraction of which have been characterized. Most sRNAs require a chaperone to facilitate interactions with their target mRNA. sRNAs that sequester Hfq (CrcZ) and RsmA (RsmY and RsmZ) result in indirect regulation of the T3SS and other virulence factors controlled by Hfq, RsmA, and Vfr (30, 48). Finally, Hfq is known to stabilize RsmY (49). Hfq stabilization protects RsmY from RNAse E-mediated cleavage and thus enhances RsmA sequestration by RsmY. Regulation of the T3SS is complex, with numerous regulators that directly or indirectly alter exsA transcription and ExsA translation (18–20, 36). Hfq can now be added to that list of regulators, highlighting the complexity of regulatory mechanisms that tightly control T3SS gene expression.

MATERIALS AND METHODS

Strain and plasmid construction. The P. aeruginosa strains and plasmids used in this study are listed in Table S1, and the cloning details and primers are provided in Tables S2 and S3. Routine cloning was performed with E. coli Top10 or DH5α cultured in LB-Lennox medium with tetracycline (12 μg/ml) or gentamicin (15 μg/ml) as required. Primers were used to amplify the sRNAs represented in the library (listed in Table S3), followed by cloning into the XbaI/Sacl restriction sites of pUN105. The Δhfq mutant was generated by PCR amplification of 5’ and 3’ flanking regions using primers N1/N2 and C1/C2 using PA103 genomic DNA as a template. The C1/C2 and N1/N2 PCR products were sequentially cloned into the HindIII/EcoRI and EcoRI/BamHI sites, respectively, of pBluescript SK. REPORTER ACTIVITY AND RSMZ LEVELS ARE UNCHANGED WHEN SRNA 179 IS OVEREXPRESSION IN THE RSMY BACKGROUND (FIG. 8B AND D) SUGGEST ADDITIONAL MECHANISMS OF INHIBITION BY SRNA 179.

In addition to sRNA 179, sRNA 0161 was recently identified with GRIL-seq as another inhibitor of the T3SS (32), and we confirmed that sRNA 0161 is also Hfq-dependent (Fig. 4A). The GRIL-seq data suggest that sRNA 0161 directly targets the exsA leader region through imperfect base pairing and is consistent with our data showing that sRNA 0161 inhibits ExsA synthesis at the posttranscriptional level (Fig. 5B and C). In addition, Hfq coprecipitates with the exsA leader region (42). The simplest model to account for the inhibitory activity of sRNA 0161 is occlusion of the ribosome binding site. Alternative mechanisms may involve RsmA and DeaD, both positive regulators of ExsA synthesis (22, 39). Rather than blocking ribosome access, sRNA 0161 may instead prevent RsmA and/or DeaD from binding to the exsA leader region. Another possibility is that RsmA and/or DeaD activate by displacing/preventing sRNA 0161 from base pairing with the exsA leader region. Testing these models will be the subject of future studies.

sRNA 0161 and 179 function independently from one another, and either is sufficient to inhibit T3SS gene expression (Fig. 4C). The primary implication is that together they provide both direct (through the activity of sRNA 1061) and indirect (through sRNA 179) Hfq-dependent mechanisms to inhibit ExsA synthesis. Transcription and synthesis of exsA and ExsA are major inputs for upstream regulatory factors (17). Although seemingly redundant, sRNA 0161 and sRNA 179 may differentially coordinate and/or respond to distinct signals that influence T3SS expression. Future studies to understand transcriptional control of sRNA 0161 and sRNA 179 will be important.

Regulatory control of the T3SS by Hfq likely extends beyond the mechanisms described here. Expression of Hfq in an sRNA 0161 179 double mutant still resulted in significant inhibition of T3SS gene expression (Fig. 4B). This may reflect sRNA-independent activity or involvement of an additional sRNA(s) in control of the T3SS. There are over 500 predicted sRNAs in P. aeruginosa (38), only a small fraction of which have been characterized. Most sRNAs require a chaperone to facilitate interactions with their target mRNA. sRNAs that sequester Hfq (CrcZ) and RsmA (RsmY and RsmZ) result in indirect regulation of the T3SS and other virulence factors controlled by Hfq, RsmA, and Vfr (30, 48). Finally, Hfq is known to stabilize RsmY (49). Hfq stabilization protects RsmY from RNAse E-mediated cleavage and thus enhances RsmA sequestration by RsmY. Regulation of the T3SS is complex, with numerous regulators that directly or indirectly alter exsA transcription and ExsA translation (18–20, 36). Hfq can now be added to that list of regulators, highlighting the complexity of regulatory mechanisms that tightly control T3SS gene expression.

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and 179 allelic exchange vectors were generated using the primer pairs listed in Tables S2 and S3 and cloned by isothermal assembly into pEXG2Tc. The ΔΔfq, Δf0161, and Δ179 mutants were generated by allelic exchange and ssaB-mediated resolution as previously described (51). The hfq gene from P. aeruginosa from strain PA103 was cloned into the XbaI and SpeI sites of pJN105 (52).

**Growth curves.** PA103 strains were grown overnight at 37°C in LB containing gentamicin (80 µg/ml) as required. The strains were diluted to an absorbance (A600) of 0.01 in a 96-well plate. A600 was measured every hour using a Tecan plate reader (Tecan Trading AG, Switzerland) until the stationary phase. Background absorbance measured from wells with only LB was subtracted, and absorbance was plotted against time.

**β-galactosidase assays and immunoblots.** PA103 strains were cultured at 37°C overnight in LB containing 80 µg/ml gentamicin as required. The next day, strains were diluted to an absorbance (A600) of 0.1 in tryptic soy broth (TSB) supplemented with 100 mM monosodium glutamate and 1% glycerol. Arabinose was added to induce Hfq (0.05%) or sRNA (0.2%) expression from the Pm promoter, and rhamnose (0.005%) was added to induce exsA expression from the Pm promoter as appropriate. Cultures were incubated at 37°C and harvested when the A600 reached 1.0. β-galactosidase activity was determined as previously described with the substrates ortho-nitrophenyl-galactopyranoside (ONPG) (36) or chlorophenol red-β-D-galactopyranoside (CPRG) (18) as previously described. CPRG activity was determined by measuring product formation at 578 nM and defined as CPRG units (i.e., A570/culture A600/time [min]/culture vol [ml]) × 1,000. Statistical analyses were determined with one-way analysis of variance (ANOVA) using GraphPad Prism version 5.0c for Mac OS X (GraphPad, La Jolla, CA). Immunoblots using rabbit immune serum to ExsA, Hfq, ExoU, PcrV, and Hcp1 were performed as previously described (36).

**Cytotoxicity assays.** Chinese hamster ovary (CHO) cells (ATCC CCL-61) were cultured in Ham’s F-12 medium (Invitrogen Corp., Carlsbad, California) supplemented with 10% fetal calf serum, 50 µg/ml penicillin and streptomycin, 2 mM L-glutamine, 0.12% sodium bicarbonate, and 2.5 mM HEPES at 37°C in 5% CO2. For cytotoxicity assays, CHO cells were seeded at 8 × 104 cells/well into 24-well tissue culture plates (80 mm). Cells were incubated at 37°C for 30 min and then mixed with 2 µl of the supernatant was transferred to a 96-well plate and assayed for lactate dehydrogenase (LDH) release using the CytoTox 96 nonradioactive cytotoxicity assay (Promega, Madison, WI). The percent cytotoxicity was calculated by subtracting the optical density at 490 nm (OD490) of an uninfected control from each sample and using WT PA103 as the positive control normalized to 100%.

**Hfq purification.** E. coli Tuner(DE3) expressing histidine-tagged Hfq(pET23bHfq) was cultured overnight on LB with ampicillin (200 µg/ml). Cell suspensions were prepared and used to start a 2-liter culture at an A600 of approximately 0.1 in LB with ampicillin (200 µg/ml). At an A600 of 0.5, isopropyl-β-D-1-thiogalactopyranoside (IPTG) was added at 1 mM to induce protein production. Once the culture reached an A600 of 3.0, cells were harvested by centrifugation and suspended in nickel nitrotriacetic acid (Ni-NTA) binding buffer (20 mM Tris-HCl [pH 7.9], 300 mM NaCl, 5 mM imidazole) supplemented with 1 complete protease inhibitor cocktail (PIC) tablet (Roche). Cells were lysed by passage through a microfluidizer at 17,000 lb/in2 (Microfluidics, Westwood, MA). Cell lysates were cleared by centrifugation and immediately loaded onto a 1-ml Hit Trap His column (GE Healthcare Life Sciences) and eluted with binding buffer containing 300 mM imidazole. The peak elution fractions were pooled and dialyzed against Ni-NTA binding buffer containing 5 mM diethanolamine (DTT) for 4 h at 4°C. The resulting protein was flash-frozen in 1-ml aliquots. Two 1-ml frozen aliquots of Hfq were buffer-exchanged by diluting to 10 ml with Ni-NTA binding buffer and then concentrated to 1 ml using an Amicon Ultra centrifugal filter (nominal molecular weight limit, 10,000 Da) and repeated five times. The concentrated Hfq samples were then combined and exposed to 0.15 µL of prequillibrated Talon beads (Clontech) for 10 min, rocking at 4 degrees. Beads were then washed with 5 ml of Ni-NTA binding buffer three times and eluted with 10 ml of binding buffer containing 0.5 M imidazole. The eluted Hfq was concentrated again and buffer-exchanged three times to storage buffer (20 mM Tris-HCl [pH 7.9], 300 mM NaCl, 1 mM DTT). Protein concentration was determined using the Bradford assay.

**Electrophoretic mobility shift assays.** RNA was generated from DNA templates encoding the exsA UTR region and sRNA 179 by in vitro transcription and end-labeled with [γ-32P] ATP as previously described (24). Purified Hfq, at the indicated concentrations was incubated with the RNA probes in 1× binding buffer (10 mM Tris-HCl [pH 7.5], 10 mM MgCl2, 100 mM KCl, 3.25 ng/µl total yeast tRNA [Life Technologies], 10 mM DTT, 5% [vol/vol] glycerol, and 0.1 unit RNaseOUT [Life Technologies]). Reaction mixtures were incubated at 37°C for 30 min and then mixed with 2 µl of gel loading buffer II (Life Technologies) and immediately subjected to electrophoresis on 10% (wt/vol) native polyacrylamide glycin gel (10 mM Tris-HCl [pH 7.5], 380 mM glycin, and 1 mM EDTA) at 4°C. Imaging was performed using an FLA-7000 phosphorimager (Fujiﬁlm), and the images were analyzed using MultiGauge v3.0 software.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**FIG S1**, TIF file, 0.3 MB.

**FIG S2**, TIF file, 0.3 MB.

**TABLE S1**, DOCX file, 0.03 MB.

**TABLE S2**, DOCX file, 0.01 MB.
TABLE S3, DOCX file, 0.1 MB.

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