Identification of BvgA-Dependent and BvgA-Independent Small RNAs (sRNAs) in *Bordetella pertussis* Using the Prokaryotic sRNA Prediction Toolkit ANNOgesic

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ABSTRACT Noncoding small RNAs (sRNAs) are crucial for the posttranscriptional regulation of gene expression in all organisms and are known to be involved in the regulation of bacterial virulence. In the human pathogen *Bordetella pertussis*, which causes whooping cough, virulence is controlled primarily by the master two-component system BvgA (response regulator)/BvgS (sensor kinase). In this system, BvgA is phosphorylated (Bvg+ mode) or nonphosphorylated (Bvg− mode), with global transcriptional differences between the two. *B. pertussis* also carries the bacterial sRNA chaperone Hfq, which has previously been shown to be required for virulence. Here, we conducted transcriptomic analyses to identify possible *B. pertussis* sRNAs and to determine their BvgAS dependence using transcriptome sequencing (RNA-seq) and the prokaryotic sRNA prediction program ANNOgesic. We identified 143 possible candidates (25 Bvg+ mode specific and 53 Bvg− mode specific), of which 90 were previously unreported. Northern blots analyses confirmed all of the 10 ANNOgesic candidates that we tested. Homology searches demonstrated that 9 of the confirmed sRNAs are highly conserved among *B. pertussis*, *Bordetella parapertussis*, and *Bordetella bronchiseptica*, with one that also has homologues in other species of the Alcaligenaceae family. Using coimmunoprecipitation with a B. pertussis FLAG-tagged Hfq, we demonstrated that 3 of the sRNAs interact directly with Hfq, which is the first identification of sRNA binding to Hfq. Our study demonstrates that ANNOgesic is a highly useful tool for the identification of sRNAs in this system and that its combination with molecular techniques is a successful way to identify various BvgAS-dependent and Hfq-binding sRNAs.

IMPORTANCE Noncoding small RNAs (sRNAs) are crucial for posttranscriptional regulation of gene expression in all organisms and are known to be involved in the regulation of bacterial virulence. We have investigated the presence of sRNAs in the obligate human pathogen *B. pertussis*, using transcriptome sequencing (RNA-seq) and the recently developed prokaryotic sRNA search program ANNOgesic. This analysis has identified 143 sRNA candidates (90 previously unreported). We have classified their dependence on the *B. pertussis* two-component system required for virulence, namely, BvgAS, based on their expression in the presence/absence of the phosphorylated response regulator BvgA, confirmed several by Northern analyses, and demonstrated that 3 bind directly to *B. pertussis* Hfq, the RNA chaperone involved in mediating sRNA effects. Our study demonstrates the utility of combining RNA-seq, ANNOgesic, and molecular techniques to identify small RNAs in the human pathogen *B. pertussis*, which may reveal their roles in pathogenesis.
identify various BvgAS-dependent and Hfq-binding sRNAs, which may unveil the roles of sRNAs in pertussis pathogenesis.

**KEYWORDS** small RNA, Hfq, BvgAS regulon, pertussis, RNA-seq, ANNOgesic

Small noncoding RNA (sRNA) regulators function as major players in posttranscriptional regulation, affecting a broad spectrum of cellular processes in bacterial physiology (reviewed in references 1–6). In many systems, sRNAs are specifically expressed as cells adapt to environmental conditions, including oxidative stress, DNA damage, iron and nutrition starvation, and lower temperatures. A major class of these sRNAs exerts their actions via base pairing with multiple target genes and modulate translation and/or mRNA stability (5, 7). This function can depend upon the RNA chaperone Hfq, a member of the Lsm/Sm family of proteins that bind sRNAs and mRNAs and facilitate their interaction (8–10). Disruption of the hfq gene shows pleotropic physiological effects in *Escherichia coli*, such as growth defects and increased sensitivity to stresses (11). In addition, hfq mutants exhibit significant virulence attenuation in a wide variety of pathogens, affecting invasion of epithelial cells, secretion of virulence factors, and survival in macrophages, suggesting that Hfq is essential for both physiological fitness and pathogenesis (9, 12). Hfq-dependent sRNAs have also been reported to be involved in bacteria/host interactions through modulating the expression of various virulence determinants, such as those involved in quorum sensing, type III secretion systems (T3SSs), iron transport, and biofilm formation (9, 12–14).

*Bordetella pertussis* is a Gram-negative bacterial pathogen that causes the highly contagious, acute respiratory illness whooping cough (pertussis). Despite a high coverage rate of vaccination (~85% worldwide according to WHO [15]), infection by this pathogen has resurged globally, in particular among vaccinated populations in developed countries. This resurgence has been attributed in part to the shorter-lived immunity achieved by the current acellular vaccine relative to the previously used whole-cell vaccine (16). Therefore, there is an urgent need for a comprehensive understanding of the molecular mechanisms of *B. pertussis* pathogenesis in order to develop better vaccines that engender longer-lasting and more effective immunity.

In *B. pertussis*, the majority of virulence genes are regulated by BvgAS, a master two-component system composed of BvgS, the histidine sensor kinase, and BvgA, the response regulator (17). Under standard growth conditions at 37°C, BvgS is autophosphorylated and transfers phosphate to BvgA via a phosphorelay. Phosphorylated BvgA (BvgA–P) dimers then activate virulence-activated genes (vags), such as pertussis toxin (ptx), filamentous hemagglutinin/adhesion (fha), and fimbriae (fim). This is known as the Bvg1 mode (18–20). BvgA–P dimers also activate the promoter for bvgR, which encodes a cyclic di-GMP (c-di-GMP) phosphodiesterase, which converts c-di-GMP to GMP (21, 22). The secondary messenger c-di-GMP, together with another response regulator RisA, is needed to activate a set of virulence-repressed genes (vrgs) (21). In the presence of BvgR, the level of c-di-GMP dramatically decreases, which in turn leads to a loss of RisA activation. As a consequence, vrgs are repressed in the Bvg1 mode. On the other hand, switching cells to a lower temperature (25°C) or growth in a high concentration (20 to 50 mM) of either nicotinic acid or magnesium sulfate (MgSO4) modulates gene expression, by inducing the Bvg2 mode (18–20, 23). Under these modulated conditions, BvgA is not phosphorylated. Without BvgA–P, BvgR remains inactive, and subsequently, the expression of vrgs increases. These observations suggest sophisticated networks among various regulators during the *B. pertussis* life cycle.

Our previous transcriptomic study identified >550 genes (~15% of the *B. pertussis* genome), which are affected by BvgAS (24). However, this number does not include any RNAs that were omitted from the annotation of the Tohama I reference genome, such as sRNAs and transcripts for small open reading frames (ORFs). Given that Hfq and sRNA regulators play a major role in the virulence of various pathogens (12, 25, 26), and that the hfq deletion mutation in the clinical strain *B. pertussis* 18323 affects the expression of
various virulence genes, including those within the T3SS (14), the involvement of sRNAs in gene regulation during *B. pertussis* pathogenesis seems likely.

Previous efforts to investigate sRNAs in *B. pertussis* have been limited. An *in silico* search for *B. pertussis* sRNAs using sRNA identification protocols and high-throughput technologies (SIPHT) (27) identified 14 sRNAs, which were then confirmed by Northern blotting (28). More recently, transcriptome sequencing (RNA-seq) analyses combined with terminator 5’-phosphate-dependent exonuclease (TEX)-treated and -untreated RNA samples were used to define the transcriptome architecture of *B. pertussis* Tohama I, revealing >500 “orphan” RNAs, which presumably include sRNAs; however, in this case, the candidates, which were collected using one condition, were not confirmed experimentally (29). Other work identified one of these candidates, RgtA, as a Bvg^- sRNA that is involved in glutamate metabolism (30). Finally, 9 additional sRNAs have been identified during the colonization of mouse trachea by the clinical strain *B. pertussis* 18323 (31).

Here, we conducted a genome-wide transcriptomic search for sRNAs using RNA-seq data that we previously used to define the BvgAS regulon in *B. pertussis* Tohama I (24). To process the data, we performed a computational analysis using the prokaryotic sRNA search program ANNOgesic, a comprehensive tool for generating bacterial genome annotations based on RNA-seq data (32). ANNOgesic was recently developed to surpass the limitations of current bacterial sRNA search programs. It can detect more than 20 genomics features, including sRNAs. The overall detection rate is ~80%; specifically, for sRNAs in *Helicobacter pylori* 26695, *Campylobacter jejuni* 81116, and *E. coli* K-12, the sensitivity has reached 90%, 84%, and 90%, respectively.

We picked several candidates to analyze by Northern blots and Hfq-binding studies. Our study demonstrates that combining RNA-seq, ANNOgesic, and molecular techniques is a successful approach for identifying various BvgAS-dependent and Hfq-binding sRNAs, which may reveal the roles of sRNAs in pertussis pathogenesis.

**RESULTS**

**RNA-seq analysis using ANNOgesic reveals sRNA candidates.** To search for *B. pertussis* sRNAs that were absent from the genomic annotation, we employed ANNOgesic, a recently developed RNA-seq-based genome annotation suite for bacteria and archaea, which contains a module for prokaryotic sRNA detection (32). This toolkit has not been used previously with *B. pertussis* RNA. The input fastq files were obtained from our previous RNA-seq study, which had identified the BvgAS-dependent regulon (GEO no. GSE98155) (24). Duplicates, collected under each condition, consisted of the following: (i) WT strain grown without MgSO4 (BvgA^+P is present), (ii) WT strain grown with MgSO4 (BvgA^+P is absent), (iii) the ΔbvgAS strain grown without MgSO4, and (iv) the ΔbvgAS strain grown with MgSO4.

To analyze the sequencing data, reads were first trimmed by Trimmomatic and aligned to the *B. pertussis* Tohama I reference genome with an RNA-seq pipeline, READemption (33), to generate the coverage wiggle (WIG) format files. These files were then processed by ANNOgesic to predict sRNAs from each condition.

In an ANNOgesic analysis “average coverage” is the number of mapped reads for an sRNA divided by the length of the designated gene (sRNA), a value that is similar to reads per kilobase per million (RPKM) mapped reads. As the average coverage value increases, the probability of a valid sRNA increases. Consequently, we used the average coverage value to evaluate the expression of a particular sRNA (see Table S1 in the supplemental material). After eliminating RNAs corresponding to the abundant SsrA and rRNAs (which had ANNOgesic rankings of 1 to 3), predicted sRNAs, whose best average coverage values were more than 30 in any of the 8 samples, were considered significant. This yielded the 143 candidates shown in Table S1.

Previously, various *B. pertussis* sRNAs present in the Tohama I strain have been reported. Early *in silico* analyses identified 14 sRNAs, which were confirmed by Northern analyses (28).
In a later global transcriptomics analysis of the strain grown under standard laboratory conditions, >500 orphan and antisense RNAs were identified; it was speculated that some of these RNAs would represent sRNA candidates (29). One of these candidates, RgtA, was shown to be dependent on the RNA chaperone Hfq, to be more abundant in the Bvg⁺ mode, and to be involved in glutamate transport (30).

To compare our ANNOgesic candidates to those in these previous reports, we classified any of our sRNA candidates as previously identified if they were transcribed in the same intergenic region with the same directionality. We found that 49 of our 143 candidates were previously identified in the study by Amman et al. (Table S1) (29), while 4 other candidates were found in the study by Hot et al. (28). Consequently, 90 possible sRNAs were newly identified by our ANNOgesic analysis.

**BvgAS dependence of predicted sRNAs.** To classify the BvgAS dependence of the 143 sRNAs predicted by ANNOgesic, we compared the average coverage of the following: set (i), WT without or with MgSO₄; and set (ii), WT versus the ΔbvgAS strain without MgSO₄. We considered a difference of 1.5-fold or greater in the coverage score in the average reads in either set as significant. There are 25 candidates that are more abundant in the presence of BvgA⁺P (see Table S2 in the supplemental material) and 53 candidates that are more abundant in the absence of BvgA⁺P (see Table S3 in the supplemental material).

As a comparison, we also used an independent method to visually identify sRNAs within our transcriptomic data set using Signal Map (Roche). Twenty sRNAs found by this visual analysis are shown in Fig. S1 in the supplemental material. Within this list, 17 sRNAs, S1 to S17, were among those predicted by ANNOgesic. About one-half of them (S2, S4, S6, S8, S10, S14, and S17) had been predicted, but not confirmed, in previous work. In addition, S9 has been recognized previously as a riboswitch and involved in flavin mononucleotide (FMN) biosynthesis (29, 34, 35), and we confirmed this analysis using the Rfam database (36).

The following three additional candidates shown in Fig. S1E were not predicted by ANNOgesic: S18, which had been previously identified as RgtA, a sRNA whose expression was reported to be activated by the B. pertussis response regulator RisA in the Bvg⁺ mode (30); and S19 and S20, which were previously unrecognized.

**Northern analyses confirm ANNOgesic sRNA predictions with high sensitivity.** Twelve of the sRNA candidates in Fig. 1 were chosen for validation by Northern blotting analysis, namely, S1, S2, S3, S4, S8, S9, S10, S11, S12, and S17, which were identified by ANNOgesic; and S18 and S20, which were identified by the Signal Map analysis (Table 1). These candidates were selected because they were carried wholly within intergenic regions (except S4), they represented very highly ranked ANNOgesic sRNAs (S1 to S4 and S17), and/or they represented a mixture of predicted Bvg⁺, Bvg⁻, or Bvg-independent transcripts. In addition, the 3’ end of S4 (Fig. S1) was antisense to the 3’ end of gene BP3392, the transposase present within a particular copy of an IS481 element. As hundreds of copies of this element are present within the B. pertussis genome (37), the presence of an sRNA that might regulate transcription from this element could provide an important regulatory feature. Finally, S18 and S20 represented two possible sRNAs that were found by the Signal Map analysis, but they were not predicted by ANNOgesic.

Northern analyses (probes listed in Table S4 in the supplemental material) confirmed the presence of all the candidates (Fig. 1, Table 1), indicating that ANNOgesic correctly predicted all 10 of its candidates that we tested. This finding indicates that the sensitivity, or the true positive rate, of ANNOgesic for detecting sRNA was quite high, namely, in this case 10 out of 10 or 100%. Although the presence of many different-sized species for S4 was unusual, heterogenous species were also observed in the S4 Signal Map (Fig. S1A).

The Bvg mode assigned by the average coverage of WT ± MgSO₄ or WT versus ΔbvgAS (Table 1; Table S2, S3) matched that observed with the Northern blot analyses except for 4 sRNAs, as follows: S1, S2, S8, and S12. However, in the cases of S1 and S12, the situation was complicated by the presence of multiple species, of which some were more abundant in one mode than another (Fig. 1, Table 1). Discrepancies
between the visual Signal Map results and ANNOgesic can arise because the Signal Map result is based on a visual examination that does not provide any statistical information, whereas ANNOgesic provides the statistical information of the expression level for each potential sRNA signal.

**TABLE 1** sRNA candidates in *B. pertussis*

| Name | Rank | ANNOgesic ID | Neighboring genes | Direction | Expression mode | ANNOgesic | Northern blot |
|------|------|--------------|-------------------|-----------|----------------|------------|--------------|
| S1   | 4    | 212          | BP2496 BP2497     | <         | Bvg(−)         | Bvg(−)/Bvg(+) |
| S2   | 5    | 128          | BP1348 dnaX       | >         | Bvg(−)         | Independent |
| S3   | 6    | 188          | ndk valS          | <         | Independent    | Independent |
| S4   | 7    | 296          | BP3391 BP3392    | >         | Independent    | Independent |
| S5   | 8    | 16           | phoB ubiE        | <         | Bvg(−)         | Bvg(+)     |
| S6   | 104  | 294          | <BP3385 BP3386   | <         | Independent    | Independent |
| S7   | 11   | 168          | BP1907 cyS       | >         | Independent    | Independent |
| S8   | 16   | 123          | BP1496 thrS      | >         | Independent    | Bvg(+)     |
| S9   | 17   | 45           | BP0470 ribB      | >         | Bvg(+)         | Bvg(+)     |
| S10  | 25   | 298          | BP3409 BP3410    | <         | Bvg(+)         | Bvg(+)     |
| S11  | 26   | 242          | BP2886 BP2887    | <         | Bvg(+)         | Bvg(+)     |
| S12  | 28   | 17           | ubiE BP0162      | >         | Independent    | Bvg(−)/Bvg(+) |
| S13  | 30   | 239          | panB BP2851      | <         | Bvg(−)         | Bvg(+)     |
| S14  | 34   | 95           | BP1164 BP1165    | >         | Independent    | Bvg(+)     |
| S15  | 86   | 288          | BP3352 BP3353    | <         | Independent    | Independent |
| S16  | 52   | 42           | <BP0405 <tRNA Val| <         | Bvg(−)         | Bvg(+)     |
| S17  | 9    | 276          | BP3151 BP3152    | <         | Bvg(+)         | Bvg(+)     |
| S18  | 16   | 123          | BP2735 BP2736    | <         | Bvg(−)         | Bvg(+)     |
| S19  | 28   | 17           | <dnaA rpmH       | >         | Independent    | Independent |
| S20  | 30   | 239          | <BP0920 cTB      | >         | Independent    | Independent |

*Names in bold were confirmed by Northern blot analyses.

**ANNOgesic rank, ID, and expression mode (see Table S1). S18, S19, and S20 were not identified by ANNOgesic so this information is not available for them.

**Neighboring genes of the sRNA; > corresponds to top strand gene; < corresponds to bottom strand gene.

**Directionality of the sRNA transcription; >, top strand; <, bottom strand.

**Expression mode of sRNA in RNA-seq or Northern blot analyses. Bvg(−), observed in the presence of BvgA−; Bvg(+), observed in the absence of BvgA−; 0, not detected; CONST, similar levels in BvgA(−) and BvgA(+).

**Northerns were performed for sRNAs in bold.
Gene conservation among *B. pertussis* sRNAs. Gene conservation among closely related organisms is an important consideration for the validation of potential sRNAs and has been used as a method to identify sRNAs in various bacterial genomes (38). Consequently, we examined the sequence conservation of the sRNAs that were predicted by ANNOgesic using NCBI BLAST together with the multiple ortholog alignment tool of BioCyc. This strategy revealed that except for S4, all of these sRNAs are highly conserved in sequence and genomic location among *B. pertussis*, *B. bronchiseptica*, and *B. parapertussis* (Fig. 2, Fig. S2A to H). For S2, extensive conservation extends to other *Bordetella* spp. and even to *Achromobacter xylosoxidans* within the Alcaligenaceae family (Fig. S2B). In addition, for several of the sRNAs, including the previously unidentified sRNA S3, there is also limited homology outside the *Bordetella* genus (Fig. 2; Fig. S2A, D, and E). These observations suggest that these sRNAs may have similar functions in various related species. For S4, which is antisense to the 3'-end of a particular IS481 element transposase, conservation was seen only among various *B. pertussis* genomes.

Identification of 3 Hfq-associated sRNAs. The bacterial RNA chaperone Hfq is a major player in posttranscriptional gene regulation, modulating the stability of mRNAs via base pairing with trans-encoded sRNAs (8, 9). As Hfq is intimately involved in sRNA regulation, we would expect that at least some pertussis sRNAs would be dependent on *B. pertussis* Hfq (HfqBp). In fact, previous work has demonstrated that an *hfq* deletion mutation in the clinical strain *B. pertussis* 18323 affected the expression of various virulence genes, including those associated with the T3SS (14).

Based on this assumption, we investigated whether any of our identified sRNAs were associated with Hfq using a pulldown assay in a strain in which the chromosomal *hfq* of *B. pertussis* was tagged with a 3× FLAG epitope, generating Hfq<sup>Bp</sup>-FLAG. Hfq of the *Bordetella* genus lacks a C-terminal extension found in many other proteobacteria, including *E. coli*, *Salmonella enterica*, and *Yersinia pestis* (Fig. 3A). As it was important to confirm that neither the natural truncation nor a C-terminal tag eliminated Hfq function, we investigated the ability of a tagged Hfq<sup>Bp</sup> to complement an *E. coli* *hfq* deletion strain, which had previously been used to screen the effect of *E. coli* *hfq* mutants on sRNAs (39).

The *E. coli* strain used to assay the tagged Hfq<sup>Bp</sup> carries a translational fusion of *rpoS-lacZ* in the chromosome under the control of the arabinose inducible promoter P<sub>BAD</sub> (Fig. 3B). Complementation by a plasmid-borne *hfq* is assayed by an increase in the *rpoS-lacZ* fusion
protein, as manifested by an increase in β-galactosidase. This arises because translation of the rpoS mRNA is positively regulated by Hfq together with the sRNAs DsrA, RprA, and ArcZ (40–42).

We assayed the complementation ability of two C-terminally tagged HfqBp proteins, namely, HfqBp-FLAG and His-tagged HfqBp-HIS; the isogenic E. coli WT and hfq deletion strains served as controls. We tested HfqBp containing the FLAG tag as well as one with a His tag to ask whether the nature of the tag would affect function. As expected, the β-galactosidase level was very low in the E. coli hfq deletion mutant (Fig. 3C). In contrast, when either HfqBp-FLAG or HfqBp-HIS was present in the E. coli hfq deletion strain, the level of β-galactosidase was comparable to that obtained in the E. coli WT strain. These results indicated that the C-terminally tagged HfqBp proteins are active in this construct.

Based on this result, we used B. pertussis containing the hfqBp-flag within its chromosome to coimmunoprecipitate RNA with Hfq. Both the WT and the hfqBp-flag strains were grown with and without MgSO4. After coimmunoprecipitation of the HfqBp-FLAG complexes with an anti-FLAG antibody, RNA was extracted for Northern blot analyses to monitor the levels of sRNAs. Using this system, we confirmed the HfqBp dependence of particular species within three sRNAs. The Bvg-independent S4, which contains heterogeneous transcripts, had species of ~350 nucleotide (nt), 200 nt, 150 nt, and 75 nt long that precipitated with tagged Hfq, while the Bvg S17 species of ~50 nt long and the Bvg- S18 (RgtA) species of ~200 nt long were enriched in the Hfq pulldown (designated by short lines in Fig. 4).

Identification of various genes in Tohama I that are affected by Hfq. We next investigated whether the expression of particular B. pertussis genes was dependent on Hfq. Total RNA samples, collected from WT or the isogenic hfq deletion strain after growing in media without MgSO4, were analyzed using real-time quantitative PCR (RT-qPCR) (Fig. 5). When comparing Δhfq versus WT, genes with a fold change (FC) of ≥2 or ≤0.5 and a P value of <0.05 were considered to be significant (see Table S5 in the supplemental material). We selected a variety of genes that previous analyses had assigned as Bvg- mode,
Bvg\(^{-}\) mode, or Bvg independent (24, 43). We included 5 Bvg\(^{+}\) mode genes that encode known or putative transcriptional regulators (BP2142 [GntR family member], BP3239 and BP2878 [putative LysR family members], \textit{brpL} [an ECF sigma factor], and BP0319 [a putative regulator]) based on our previous observation that the BvgAS regulon controls dozens of proteins involved in transcription (24). Our list also included 3 classic \textit{vrgs} (\textit{vrg18}, \textit{vrg24}, and \textit{vrg73}) and 4 proteins involved in transport, as follows BP3831 (amino acid ABC transporter substrate-binding protein), BP3862 (peptide ABC transporter substrate-binding protein), \textit{vag8} (autotransporter), and \textit{sphB1} (autotransporter subtilisin-like protease). Finally, we included 4 flagellar genes, namely, \textit{flia} (sigma factor), \textit{flf} (flagellar M-ring protein), \textit{flID} (flagellar hook-associated protein 2), and \textit{flIO} (flagellar protein), based on the recent discovery that \textit{B. pertussis} can be motile (44) and our previous finding that all of these genes are BvgA independent in \textit{B. pertussis} (24) despite their being Bvg\(^{2}\) in \textit{B. bronchiseptica} (17).

Of the 23 \textit{B. pertussis} Tohama I genes we tested, the absence of HfqBp upregulated mRNA levels of 8 genes with a mean FC of \(\approx 2\) and downregulated 2 genes with a mean FC of \(\approx 0.5\) (Fig. 5, Table S5). For BP3862, encoding a peptide ABC transporter substrate-binding protein, the \(P\) value was \(>0.05\) and for BP3841 (not annotated), there was 1 analysis using 2 biological replicates, so statistics could not be performed. Consequently, we could not conclude that these two genes were significantly affected.

The two identified downregulated genes were the Bvg\(^{+}\) genes \textit{vag8} (autotransporter)
and BP1005 (conserved hypothetical protein). The identified upregulated genes were two Bvg− mode genes (BP2171 [ccoQ, encoding a Cytochrome c oxidase subunit] and BP2878 [encoding a putative LysR-family transcriptional regulator]) and the Bvg− mode genes vrg24, vrg73, BP3871 (cold shock like protein), and BP3831 (encoding an amino acid ABC transporter substrate binding protein) (Fig. 5). BP3831 is the Bvg− mode gene that has been shown to be a likely target of S18 (RgtA) (29). The finding that BP3831 is affected by an hfq deletion together with our demonstration that S18 (RgtA) interacts directly with Hfq (Fig. 4) suggests the interaction between RgtA and BP3831 mRNA is mediated by Hfq.

We compared these findings to a previous comprehensive study of genes affected by a hfq deletion in the B. pertussis clinical strain 18323 (14). In this case, FCs of Δhfq/WT were determined by microarray analyses in exponentially growing and stationary-phase cells as well as cells at 0 and 12 days after mouse lung infection. Among our significantly affected genes, BP1005 and vag8 were also downregulated under all growth conditions of B. pertussis 18323, and BP3871 and BP3831 were also upregulated during exponential growth of B. pertussis 18323. However, two upregulated genes (BP2171 and BP2878) were downregulated in the clinical strain during exponential growth and at 12 days postinfection, respectively, and two others (vrg24 and vrg73) were not dysregulated under any condition. It appears then that the B. pertussis hfq regulon is sensitive to the particular strain and growth conditions, suggesting myriad ways that sRNAs can influence B. pertussis gene expression.

**DISCUSSION**

Transcriptomic analysis by ANNOgesic reveals new B. pertussis sRNAs. While sRNA regulators have been investigated intensively in various bacterial pathogens, B. pertussis has remained an incomplete research area in this regard. A previous study using SIPHT identified 14 sRNAs, which were then confirmed by Northern blotting analyses (28). A later RNA-seq analysis, using samples collected under one condition, defined the transcriptome architecture of B. pertussis Tohama I and revealed >500 orphan and antisense RNAs, which could include sRNAs (29). One of the candidates, RgtA, was confirmed as a Bvg− mode sRNA that is involved in glutamate metabolism (30). Finally, 9 additional sRNAs have been identified during the colonization of mouse trachea by the clinical strain B. pertussis 18323 (31).

In this study, we tested the ability of the recently developed prokaryotic genome annotation toolkit ANNOgesic to correctly identify sRNAs within a B. pertussis RNA-seq data set. This analysis integrates high-throughput RNA sequencing data to predict unannotated features that have not been readily identified previously. In addition to predicting sRNA and sRNA targets from RNA-seq data, ANNOgesic offers a variety of tools for detecting the transcription start site (TSS), processing site, transcript, promoter, terminator, untranslated region (UTR), small open reading frame (sORF), operon, riboswitch, CRISPR, and circular RNA in bacteria and archaea (32), making it a pioneer in the field. It is modular, has comprehensive documentation, and is easy to use. In recent work, ANNOgesic has been used to help define the transcriptome map of Bacteroides thetaiotaomicron (45).

A recent comparison of different available sRNA prediction tools (46) indicated that three methods (APERO, ANNOgesic, and TLA from RNA-eXpress) outperformed others. Within these three tools, APERO worked for paired-end sequencing data sets only, while TLA was not specific for bacteria. Therefore, ANNOgesic was an excellent choice for sRNA detection in this study. Our results validated this choice. Not only did ANNOgesic provide a high true positive rate on sRNA prediction but also its results inferred correctly the expression mode of those sRNAs in most cases.

Using ANNOgesic, we found 143 putative sRNAs, including 90 candidates that had not been reported previously. Northern blot analyses confirmed all 10 of the sRNA candidates that we tested. Furthermore, our homology searches indicated that 9 of the confirmed candidates are highly conserved within various Bordetella spp. (Fig. 2, Fig. S2), a feature that is consistent with their providing an important function within the Bordetella genus. Overall, we conclude that ANNOgesic is a highly useful tool for the
identification and classification of sRNAs in this system. We expect that the vast majority of the 90 previously unrecognized \textit{B. pertussis} sRNAs that were predicted by ANNOgesic will be valid.

Despite this success, the program did miss RgtA (S18), which was identified in a previous study (30), and S19 and S20, which we found by a visual analysis of our data set. It is possible that ANNOgesic missed these RNAs because we set a stringent cutoff of a best average coverage value greater than 30 in at least one sample. In addition, in the case of S19, ANNOgesic may have assigned this RNA as a part of the leader region of \textit{rpmH} rather than a unique sRNA. For example, various sRNAs, which are transcribed from the leader region (5' UTR) of genes, such as \textit{grAL} in \textit{E. coli} and EutX, a part of the riboswitch of \textit{eut} genes in \textit{Enterococcus faecalis}, were also not initially recognized as sRNAs (47, 48).

**Dependence of \textit{B. pertussis} sRNAs on Hfq.** The RNA chaperone Hfq is found in multiple bacterial species, from nitrogen-fixing symbionts to \textit{Enterobacteriaceae} pathogens, and is a major player in posttranscriptional gene regulation, modulating the stability of mRNAs mostly via base-pairing trans-encoded sRNAs (5, 8–10). Hfq\textsubscript{Bp} has been shown to play an essential role in the expression of various virulence genes (14, 49), as an isogenic \textit{hfq} deletion mutation of the \textit{B. pertussis} clinical strain 18323 produces smaller amounts of toxins (49), and the transcriptional levels of the T3SS and other virulence factors (\textit{vag8}, \textit{bsp22}, and \textit{tcFA}) are downregulated by an \textit{hfq} deletion (14).

Hfq\textsubscript{Bp} (8.8 kDa) is smaller than \textit{E. coli} Hfq (11.2 kDa) because the C-terminal domain (CTD) of \textit{E. coli} Hfq is missing in Hfq\textsubscript{Bp} and in other members of the betaproteobacteria phyla (Fig. 3A) Nonetheless, previous work has shown that a truncated \textit{E. coli} Hfq, harboring residues 1 to 65, maintains an active RNA chaperon activity, binding single-stranded RNA similarly to WT (50). In addition, we find that Hfq\textsubscript{Bp} is functionally active in \textit{E. coli} (Fig. 3C). Other work has found that the CTD of \textit{E. coli} Hfq is essential for the rapid release of the double-stranded RNA products, although its presence does not change the rate of RNA binding, suggesting that the CTD may be important for rapid recycling of Hfq (10, 50). The evolutionary significance for why Hfq proteins within the betaproteobacteria phyla lack the CTD has yet to be revealed.

Using an \textit{in vivo} pulldown assay, we found 3 sRNAs that interact with Hfq\textsubscript{Bp}: S4, which is expressed independently of the Bvg mode; S17, which is enhanced in the Bvg\textsubscript{1} mode; and RgtA (S18), which is enhanced in the Bvg\textsubscript{2} mode. To our knowledge, this is the first demonstration of the direct interaction of any \textit{B. pertussis} sRNAs with Hfq\textsubscript{Bp}. Previous reports have indicated that the expression of RgtA is Hfq dependent (30), and we have confirmed that this sRNA binds directly to Hfq\textsubscript{Bp}.

S4 has been shown to be associated with the expression of adhesin and virulence factors, and it is thought that it transposes into the gene. S4 RNA is quite heterogenous (Fig. 1), and the 3' end(s) of S4 have not been precisely mapped, but the RNA-seq data indicate some S4 RNAs may contain 3' ends that are antisense to the BP3392 transcript (Fig. S1A), suggesting that S4 might be able to regulate expression of the IS481 transposase RNA. Previous work has detected various antisense RNA signals that appeared to arise from the activities of inward and outward promoters of the IS481 element (29). However, in this case, the S4 transcript is derived from a promoter that is independent of the IS element. These observations, therefore, raise the possibility that S4 might affect the level of the transposase by a different manner.

In the past few years, it has become increasingly apparent that both the Bvg\textsuperscript{+} and Bvg\textsuperscript{−} modes contribute to the lifestyle of \textit{B. pertussis}; Bvg\textsuperscript{+} contributes through the expression of adhesin and virulence factors and the Bvg\textsuperscript{−} through the expression of genes involved in survival, transmission, and/or persistence (24, 51, 52). The known roles of bacterial sRNAs in regulating lifestyle and pathogenesis suggest that these
factors will be important determinants for *B. pertussis* growth under various conditions and during infection. Our work here establishes the use of ANNOgesic as a valuable tool for detecting valid sRNAs as well as the presence of Hfq-binding sRNAs in both the Bvg^+^ and Bvg^-^ modes. We expect that future work will identify the targets of these sRNAs and how they affect the *B. pertussis* life cycle.

**MATERIALS AND METHODS**

**Bacterial strains and cell culture.** *B. pertussis* Tohama I BP536 (53) and its ΔbvgAS derivative have been described (24). To create *B. pertussis* hfq::3xflag allelic exchange was performed utilizing an 898-bp fragment containing, in order, 161-bp of the upstream flanking sequence, the entire 234-bp hfq gene, a 99-bp sequence (GTCGACAAGCTTGCGGCCGCACTCGAGGACTACAAAGACGATGACGACAAGGACTACAAAGAGTATGCAGACGACTAGTACAAAGA) (33) and 404-bp downstream flanking sequence. This fragment was synthesized and cloned between the KpnI and BamHI sites of the allelic exchange plasmid pSS4894 (21) by GenScript USA, Inc. The resulting plasmid pQC2314 was then transformed into *E. coli* strain RH03 (54) to create a donor strain for conjugation. Using wild-type *B. pertussis* BP536 as a recipient, allelic exchange was carried out as described previously (21) to obtain the *B. pertussis* strain QC4594 (B. pertussis BP536, hfq::3xflag) which was further verified by PCR/sequencing using primers outside the allelic exchange construct.

To delete hfq in *B. pertussis*, we used the allelic exchange plasmid pSS4661, which contains a vegetative origin of replication (oriV); an origin of transfer (oriT); the genes encoding antibiotic resistance to ampicillin (bla), kanamycin (kan), bleomycin (ble), and streptomycin (str); a gene for the endonuclease I-SceI cleavage site; and a multiple cloning site. Using genomic DNA of *B. pertussis* BP536 as the template, the upstream flanking region (amplified by primers 3533-hfqUSOE-BSaNott [TATAAGGTCTCGGCGGGTGCAAGGCTTGAGCTATTCGTCGCTTC] and 3534-hfqUSJE [TATAAGGTCTCGGCGGGTGCAAGGCTTGAGCTATTCGTCGCTTG]) and the downstream flanking region (amplified by primers 3535-hfqDSEE-BSaNott [TATAAGGTCTCGGCGGGTGCAAGGCTTGAGCTATTCGTCGCTTG] and 3536-hfqDSOE-BSaBamHI [TATAAGGTCTCGGCGGGTGCAAGGCTTGAGCTATTCGTCGCTTG]) were digested with Bsal and ligated together into a vector cleaved with NotI and BamHI (Bsal recognition sites are in bold and cohesive ends created by digestion are underlined). The net result was an insertion of a NotI-BamHI fragment, containing flanking sequences of the hfq gene and an in-frame deletion of codons 4 to 71 of the hfq gene itself, inserted between the NotI and BamHI sites of pSS4661, yielding plasmid pQC1798. To perform allelic exchange, plasmid pQC1798 was transformed into the *E. coli* donor strain S17.1 (55) with selection on LB agar plus 20 μg/ml kanamycin. In a subsequent conjugation, the recipient was *B. pertussis* BP536 that had been grown for 3 days at 37°C on Bordet Gengou (BG) agar supplemented with 50 μg/ml streptomycin and 50 mM MgSO₄ to induce the Bvg^-^ mode, thus preventing Ptx-driven I-SceI expression upon transfer of pQC1798. Plasmid cross-in-conjugation was performed by swapping the *E. coli* donor strain together with the Bordetella recipient strain onto BG agar supplied with 50 mM MgSO₄. After incubation at 37°C for 3 h, bacteria were recovered by swabbing and were reswabbed onto BG agar plus 20 μg/ml kanamycin, 50 μg/ml streptomycin, and 50 mM MgSO₄. Exconjugants, obtained after 3 to 4 days of incubation at 37°C, were restreaked onto BG agar plus 50 μg/ml streptomycin but without MgSO₄ to allow Ptx-driven I-SceI expression, the cleavage of the I-Sce restriction site, and plasmid cross-out, resulting in *B. pertussis* strain QC2910 (*B. pertussis* BP536, Δhfq) which was further verified by PCR using primers outside the allelic exchange construct and sequencing of the PCR product.

To obtain cells for RNA isolation, cells were grown on BG agar (56) containing streptomycin (50 μg/ml) and ampicillin (50 μg/ml) at 37°C in the absence (Bvg^-^ mode) or in the presence (Bvg^+^ mode) of 50 mM MgSO₄. After 3 days, cells were scraped from plates and resuspended in PLB medium (56) with or without MgSO₄ to obtain an initial optical density at 600 nm (OD₆₀₀) between 0.1 and 0.2. Cells (in duplicates) were incubated at 37°C with shaking at 250 rpm until the OD₆₀₀ reached a value between 0.5 and 0.6. Aliquots were then collected by centrifugation.

The gene encoding hfq^pp^::flag or hfq^pp^-his was inserted into the *E. coli* Δhfq strain SG30200 (57) as follows. Gene fragments harboring hfq^pp^ with either a 3× FLAG or His-tag fused at the C terminus before the hfq stop codon were generated by PCR using primers that were designed to give PCR products with 50 nt upstream and 40 nt downstream, which were then introduced into DJS2814 (MG1655 lacX74 malP:: lacI^Δ^, Δhfq::tpA_minumanar-kan-Paad-ccdB cassette mini λ·tet; gift of Susan Gottesman, National Cancer Institute, Bethesda, MD) to replace the Δhfq::tpA_minumanar-kan-Paad-ccdB cassette. Recombinants were selected by growth on arabinoose at 37°C (to select against the mini-λ). The resulting mutants, which were verified by PCR and sequencing, yielded either KM3032 (harboring *Bp* hfq-his) or KM3033 (harboring *Bp* hfq-flag). After P1 lysates of both KM3032 and KM3033 were prepared, the fragments harboring either *Bp* hfq-his or *Bp* hfq-flag were replaced with the Δhfq::cat-sacB cassette in SG30200 (MG1655, Δhfq::cat-sacB, ΔpurK-kan) by P1 transduction, generating the final strains as either KM3040 (MG1655, *Bp* hfq-his) or KM3041 (MG1655, *Bp* hfq-flag). The strains were verified by PCR and checked by the loss of the kanamycin cassette. PM1409 (58), which contains *E. coli* hfq and is otherwise isogenic with SG30200, served as the positive control. It is referred to as WT *E. coli* in Fig. 3.

**Pipeline for sRNA prediction by ANNOgesic.** Illumina reads in fastq format from our previously published RNA-seq analyses (24) were first trimmed by Trimomatic (59) version 0.36 and then processed by the RNA-seq pipeline READemption (33) version 0.45. Reads longer than 12 nt were aligned to the reference genome
sequences for 8, pertussis Tohama I (GenBank accession no. NC_002929.2) using segemehl (60) version 0.2.0. The subcommands “gene quanti” and “coverage” in READemption were used to calculate the gene expression.

miRNA predictions were made by ANNONtastic (32) version 0.6.27 using the subcommand “sra.” The input files included the coverage wiggle files generated by READemption, the GFF annotation format, and the reference genome file (NC_002929.2). Default parameters were used, and the predicted sRNAs were ranked by the average coverage among the replicates. The average coverage score equal or greater than 30 in at least one condition was used as the cutoff.

The expression profiles of the sRNAs were designated Bvg+, Bvg−, or Bvg independent based on the average coverage of the WT ± MgSO4 or WT-MgSO4/[Δ]bvgAS-MgSO4 data sets. An FC of ≥1.5 or ≤0.67 in either data set was designated Bvg+ or Bvg−, respectively.

**Visualization of sRNAs by Signal Map.** The transcriptomic data for the previously reported RNA-seq analyses used for this work are available in the NCBI database (GEO no. GSE98155) (24). A transcriptomic analysis of the fastq files generated from these data were first performed using the open source software Rockhopper with the default parameter. This analysis generated a strand-specific and normalized wiggle file from each fastq file. The wiggle files were then converted into the GFF format, and SignalMap software (Roche) was used to visualize with the reference genome annotation (NC_002929.2).

**RNA isolation and Northern blot analysis.** Total RNA was isolated by the hot-phenol method as described (61, 62). Briefly, each frozen cell pellet was suspended with 600 μl ice-cold phosphate-buffered saline (PBS) and 100 μl 8X lysis solution (320 mM Na-acetate, 8% SDS, and 16 mM EDTA [pH 8]), and 800 μl UltraPure 25:24:1 phenol/chloroform/isoamyl alcohol (Invitrogen) were added. The mixture was incubated at 65°C for 5 min with maximum shaking, and the aqueous phase was separated by centrifugation. The aqueous phase was then reextracted using 25:24:1 phenol/chloroform/isoamyl alcohol twice. The RNA was precipitated by adding 3 volumes of ethanol to the aqueous phase and chilled at −70°C overnight. The RNA precipitate was collected by centrifugation at 4°C and washed with 70% ethanol. Each RNA pellet was dissolved in diethyl pyrocarbonate (DEPC)-treated water.

For Northern blot analyses, RNA (7 μg) was electrophoresed on 10% polyacrylamide, 7 M urea gels in 1× Tris-borate-EDTA (TBE) buffer at 180 V for ~1 h. The RNA was then electroblotted (200 V for 1.5 h at 4°C) onto a Zeta probe GT blotting membrane (Bio-Rad). Membranes were cross-linked by UV light, and the RNA was hybridized with biotinylated oligomers (Table S4) (Integrated DNA Technologies [IDT]) overnight at 42°C. Chemiluminescent signals were visualized with the Amersham Imager 600 (GE Healthcare) (61). Size marker lanes contained the Biotinylated sRNA ladder (Kerafast). Northern analyses were performed at least in duplicate.

**RT-qPCR analysis.** Real-time PCR analyses were performed as previously described (24). Oligonucleotides used to amplify target genes and the internal control are available upon request. In each RT-qPCR analysis, which contained biological duplicates or triplicates as indicated in Table S5, the cycle in which fluorescence was detected (the quantitation cycle [Cq]) was observed for the gene or reference gene rpoD (Cqref) [ΔCq = Cqgene − Cqref]; the ΔCq was determined as Cqref − Cqgene for the RNA from the Δhfq strain − ΔCq for the RNA from the WT strain; the fold change (FC) was determined as 2ΔΔCq. For the statistical analyses, the mean FC and standard error (SE) from the values for a particular gene were first determined using the website https://ncalculators.com/statistics/standard-error-calculator.htm. The t statistic was calculated as the mean FC/SE. The P value was then determined using the one-tailed hypothesis and a “degrees of freedom” value of n − 1 for the number of replicates (n) performed for a gene analyzed in a single RT-qPCR analysis and (n1 − 1) + (n2 − 1) + ... for genes in which multiple analyses were performed using the website https://www.socscistatistics.com/pvalues/tdistribution.aspx. These values are listed in Table S5.

**Hfq coimmunoprecipitation.** Wild-type B. pertussis Tohama I BP336 and the 3× FLAG-tagged hfq strain (QC4594) were collected by centrifugation at 4°C to an equivalence of an OD600 of 40 to 50. Pellets were stored at −80°C. Lysates were obtained as previously described (39, 63). Briefly, cells were resuspended with 0.8 ml of lysis buffer (20 mM Tris-Cl [pH 8], 150 mM NaCl, 1 mM MgCl2, 1 mM DTT, 1× complete protease inhibitor cocktail [Roche], and 0.05 U SUPERaseIn [ThermoFisher]) and then disrupted with 1.2 ml glass beads (0.3 mm diameter, Sigma) by sonication. Cleared lysates were collected by centrifugation in an Eppendorf microcentrifuge at 14,000 rpm at 4°C for 10 min. The supernatant was mixed with 50 μl EZview red anti-FLAG M2 affinity gel (Sigma) by rotation for 1 h at 4°C. The beads were collected by centrifugation and then washed sequentially with low salt buffer (20 mM Tris-Cl [pH 8], 150 mM NaCl, 2 mM EDTA, 1% Triton X-100, and 0.1% SDS), high salt buffer (20 mM Tris-Cl [pH 8], 500 mM NaCl, 2 mM EDTA, 1% Triton X-100, and 0.1% SDS), LiCl buffer (10 mM Tris-Cl [pH 8], 250 mM LiCl, 1 mM EDTA, 1% deoxycholate, and 1% NP-40), and twice with Tris-EDTA (TE) buffer (10 mM Tris-Cl [pH 8] and 1 mM EDTA). RNA was eluted from the beads by the addition of 0.3 ml of elution buffer (100 mM NaHCO3 and 1% SDS, containing 0.05 U SUPERaseIn). The solution was extracted with 25:24:1 phenol/chloroform/isoamyl alcohol and precipitated with ethanol, and the RNA was dissolved in diethyl pyrocarbonate (DEPC)-treated water. For the Northern analysis, 5.5 μg and 0.16 μg of the supernatant and eluant were used, respectively, and samples were treated as described above. Northern analyses were performed at least in duplicate.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available online only.

**SUPPLEMENTAL FILE 1**, XLSX file, 0.1 MB.

**SUPPLEMENTAL FILE 2**, PDF file, 2.1 MB.
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We declare no conflicts of interest.

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