



An RNA-centric global view of *Clostridioides difficile* reveals broad activity of Hfq in a clinically important gram-positive bacterium

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The gram-positive human pathogen *Clostridioides difficile* has emerged as the leading cause of antibiotic-associated diarrhea. However, little is known about the bacterium's transcriptome architecture and mechanisms of posttranscriptional control. Here, we have applied transcription start site and termination mapping to generate a single-nucleotide-resolution RNA map of *C. difficile* 5' and 3' untranslated regions, operon structures, and noncoding regulators, including 42 sRNAs. Our results indicate functionality of many conserved riboswitches and predict *cis*-regulatory RNA elements upstream of multidrug resistance (MDR)-type ATP-binding cassette (ABC) transporters and transcriptional regulators. Despite growing evidence for a role of Hfq in RNA-based gene regulation in *C. difficile*, the functions of Hfq-based posttranscriptional regulatory networks in gram-positive pathogens remain controversial. Using Hfq immunoprecipitation followed by sequencing of bound RNA species (RIP-seq), we identify a large cohort of transcripts bound by Hfq and show that absence of Hfq affects transcript stabilities and steady-state levels. We demonstrate sRNA expression during intestinal colonization by *C. difficile* and identify infection-related signals impacting its expression. As a proof of concept, we show that the utilization of the abundant intestinal metabolite ethanolamine is regulated by the Hfq-dependent sRNA CDIF630nc_085. Overall, our study lays the foundation for understanding clostridial riboregulation with implications for the infection process and provides evidence for a global role of Hfq in posttranscriptional regulation in a gram-positive bacterium.

small RNA | Hfq | differential RNA-seq | *C. difficile* | transcript termination site

Antibiotic-resistant bacteria are a major global threat to human health, endangering our ability to perform a range of modern medical interventions. The obligate anaerobe, spore-forming *Clostridioides difficile* has become the leading cause of antibiotic-associated diarrhea over the past two decades (1). Increasing numbers of multiresistant clinical isolates (2) and recurrent infections are key challenges in the treatment of *C. difficile* infections (CDI), often leaving fecal microbiota transfer as the only clinical option (3, 4).

The clinical challenges posed by CDI have prompted much effort to understand how this pathogen regulates virulence in response to environmental conditions. As a result, there is a comprehensive body of literature focusing on toxin production and sporulation control by several global metabolic regulators including CcpA, CodY, Rex, and PrdR (5, 6). Furthermore, several specialized and general sigma factors including TcdR (6), Spo0A (7), SigD (8, 9), SigH (10), and SigB (11–13) are linked to virulence and metabolism, although the exact molecular mechanisms often remain unknown. Most of this knowledge has been accumulated through detailed studies of individual genes and promoters, whereas RNA-sequencing (RNA-seq)-based annotations

of the global transcriptome architecture, which have accelerated research in the gram-positive pathogens *Listeria* (14), *Staphylococcus* (15), and *Streptococcus* (16), have only recently become available for *C. difficile* (17).

This paucity of global knowledge about RNA output in *C. difficile* readily extends to posttranscriptional control of gene expression. The bacterium is of particular scientific interest, being the only gram-positive species thus far in which deletion of *hfq* seems to have a large impact on gene expression and bacterial physiology (18–20). Specifically, deletion of *hfq* increases sporulation (21), a crucial pathogenic feature of this bacterium that enables transmission between hosts. In gram-negative bacteria, Hfq commonly exerts global posttranscriptional control by facilitating short base-pairing interactions of small regulatory RNAs (sRNAs) with *trans*-encoded messenger RNAs (mRNAs) (22, 23), but its role in gram-positive bacteria remains controversial. For example, functional studies on Hfq in *Bacillus subtilis*, a model bacterium for Firmicutes, revealed in vivo association with a subset of sRNA (24), but comparative analyses of a wild-type and *hfq* knockout strain revealed only moderate effects on sRNA and mRNA transcript levels (25) and the absence of any significant growth defect (26), which lead to the conclusion that Hfq plays a minor role in posttranscriptional regulation in *B. subtilis*.

Significance

Clostridioides difficile is the leading cause of healthcare-associated diarrhea worldwide following antibiotic treatment. Consequently, there is medical need for novel antibacterial agents acting against *C. difficile* that leave the resident microbiota unharmed. The development of such narrow-spectrum antibiotics requires precise knowledge of the mechanisms that fine-tune gene expression to orchestrate the genomic output at each locus in the genome. We address this issue by defining the global transcriptome architecture of *C. difficile* including noncoding regulatory elements, many of which are expressed during gut colonization. Our analysis of these regulators provides evidence for the global function of Hfq in sRNA binding and stabilization in a gram-positive bacterium.

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Previous efforts using *in silico* methods (27) and RNA-seq (28) have predicted >100 sRNA candidates in *C. difficile* and identified widespread RNA binding by Hfq (20), which suggests the existence of a large posttranscriptional network in *C. difficile*. However, under which conditions these sRNAs are expressed, which targets they regulate, and whether they depend on Hfq remain fundamental open questions.

Another important feature of posttranscriptional control in *C. difficile* are *cis*-regulatory RNA elements. Pioneering work has deciphered the function of genetic switches in the 5' untranslated region (UTR) of the *flgB* operon, containing the early-stage flagellar genes, and the cell wall protein-encoding gene *cwpV* (29–31). Moreover, cyclic di-GMP-responsive riboswitches were shown to regulate biofilm formation and toxin production (9, 32–34). However, other members of the many different riboswitch classes have not been systematically identified. Combined with the nascent stage of sRNA biology in *C. difficile*, this argues that global approaches are needed to understand the full scope of posttranscriptional regulation in this important human pathogen.

In the present study, we applied recently developed methods of bacterial RNA biology (35, 36) to construct a global atlas of transcriptional and posttranscriptional control in *C. difficile*. We provide an example for a gram-positive bacterium with Hfq-dependent sRNA-based gene regulation and identify the ethanolamine utilization pathway as a target of sRNA-mediated regulation in *C. difficile*. Overall, we provide evidence for extensive Hfq-dependent posttranscriptional regulation and lay the foundation for future mechanistic studies of RNA-based gene regulation in *C. difficile*.

Results

High-Resolution Transcriptome Maps of *C. difficile* 630. For generating high-resolution transcriptome maps of *C. difficile*, we chose the reference strain 630 (DSM 27543, CP010905.2). Being widely used by the *C. difficile* community, this strain offers the most comprehensive genome annotation. Using two global RNA-seq approaches, we analyzed RNA samples from three different conditions: late-exponential and early stationary growth in tryptone-yeast (TY) broth and late-exponential growth in brain heart infusion (BHI) broth (Fig. 1A). The resulting genome-wide maps provide single-nucleotide-resolution transcriptional start sites (TSSs) and termination sites (TTs) (Fig. 1B). We have used this information to annotate 5' and 3' UTRs and operon structures, to correct previous open reading frame (ORF) annotations, to add previously overlooked small genes, and to annotate sRNA loci. Inspired by other gene expression databases such as SalCom (37), AcinetCom (38), and Theta-Base (39), we have launched the interactive web browser “Clost-Base” (<https://www.helmholtz-hiri.de/en/datasets/clostridium>) for our transcriptome data. This online resource for *C. difficile* allows visualization of the transcriptomic data in the context of annotated coding and noncoding genes as well as transcript features (e.g., TSS) that we have experimentally determined in this work. The browser enables search queries and retrieval of primary sequences for any annotated feature in the database to serve as a resource for the *C. difficile* community.

Genome-wide Annotation of TSSs. Differential RNA-seq (dRNA-seq) (40, 41) was performed to capture 5' ends of transcripts. In brief, one-half of the sample remains untreated (Tex⁻) to capture both, primary (5'-PPP) and processed (5'-P or 5'-OH) 5' ends of transcripts; the other half is treated with terminator exonuclease (Tex⁺), leading to specific degradation of processed (5'-P or 5'-OH) RNAs, thus enriching primary transcripts and enabling TSS annotation (Fig. 1B). Conversely, relative read enrichment in the Tex⁻ complementary DNA (cDNA) libraries indicate RNA processing sites (Fig. 1B). The *C. difficile* 630 genome comprises annotations for 3,778 coding sequences (CDS) as well as transfer RNAs (tRNAs), ribosomal RNAs (rRNAs), and the housekeeping

RNAs 6S RNA, RNaseP, tmRNA, and SRP. By identifying 2,293 TSSs, we were able to define transcriptional units of individual genes and polycistronic operons for approximately one-half of the genome (Dataset S1).

Benchmarking our global data with previous studies of individual genes (10, 30, 42), our dRNA-seq results are consistent with published TSSs of *sigH*, *cwpV*, *cwpI9*, and *spo0A* (Fig. 1C). For annotation purposes, we assigned TSSs to one of five classes according to their genomic location and expression level: pTSS (primary TSS of a gene or operon), sTSS (secondary TSS showing lower expression level compared to pTSS for the same gene or operon), iTSS (internal TSS located inside a gene), aTSS (antisense to a gene within 100-nt distance), and oTSS (orphan TSS, no nearby gene) (Fig. 1D). Naturally, some of these TSS annotations overlap (e.g., among the 1,627 pTSS, 126 are located within a gene (iTSS), and 46 are transcribed antisense (aTSS) to a gene). However, in contrast to other bacteria where antisense transcription is a pervasive transcriptome feature (43), it only accounts for ~6% of TSS events in *C. difficile*.

Internal TSSs were abundant and accounted for 20% of all start sites, many of which seem to uncouple downstream genes within operons. However, they may also indicate misannotated translational start codons or protein isoforms. One example is the putative hydrolase (CDIF630_02227) encoded between *ermB1* and *ermB2* of the erythromycin resistance cassette, which belongs to a conserved family of ATPases involved in plasmid partitioning. In addition to a very weakly expressed primary TSS upstream of the annotated start codon, we detected a strongly expressed iTSS located 14 nt inside the coding sequence. Manual checking for start codons (AUG, UUG, and GUG) and a ribosome-binding site (RBS) downstream of the iTSS identified a start codon 43 nt downstream of the existing ORF annotation, placing an RBS in optimal distance to the start codon and generating a 5' UTR length of 29 nt (SI Appendix, Fig. S1A). This annotation is in line with earlier studies in *Streptococcus pyogenes*, which suggested this internal start codon is used *in vivo* (44) and accordingly used the shorter variant to solve the respective protein crystal structure (PDB ID: 2oze). Similarly, we propose reannotations in eight additional cases, including the *spaFEGRK* operon encoding an antibiotic/multidrug-family ATP-binding cassette (ABC) transport system (Dataset S2).

Promoter Architectures. *C. difficile* encodes 14 sigma factors, among them, consensus sequences have been proposed for the vegetative SigA (28), the general stress response SigB (11–13), the major transition phase SigH (10), and the sporulation-specific SigK (45) sigma factor, mostly based on comparative transcriptome analyses of respective deletion strains. In addition, a recently published strategy for genome-wide identification of promoter sequences, which combined TSS mapping and positional weight matrices based on transcriptome profiles of sigma factor mutants, expanded several existing sigma factor regulons and defined novel regulons, including one for SigL (17).

Here, we applied multiple expectation maximizations for motif elicitation (MEME)-based searches upstream of all TSS leading to the confirmation of published consensus sequences for SigA, SigB, SigH, and SigK (SI Appendix, Fig. S1B and Datasets S1 and S3). In line with recent data (17), the majority of detected TSS (1,188/2,293) were associated with a SigA-type promoter (Dataset S3), including the known TSSs for *sigH* and *clnR* (10, 46, 47). Similarly, we identify many genes previously predicted to be regulated by SigH (10, 17). Furthermore, we identified 52 genes associated with a SigK promoter signature, including known genes of the SigK regulon such as *sleC* and *cdeC* (48). Comparing our promoter predictions with previous studies showed partial overlaps which we attribute to differential computational strategies for promoter assignment as well as growth conditions used (e.g., sigK promoter mutant analyses were performed in sporulation conditions

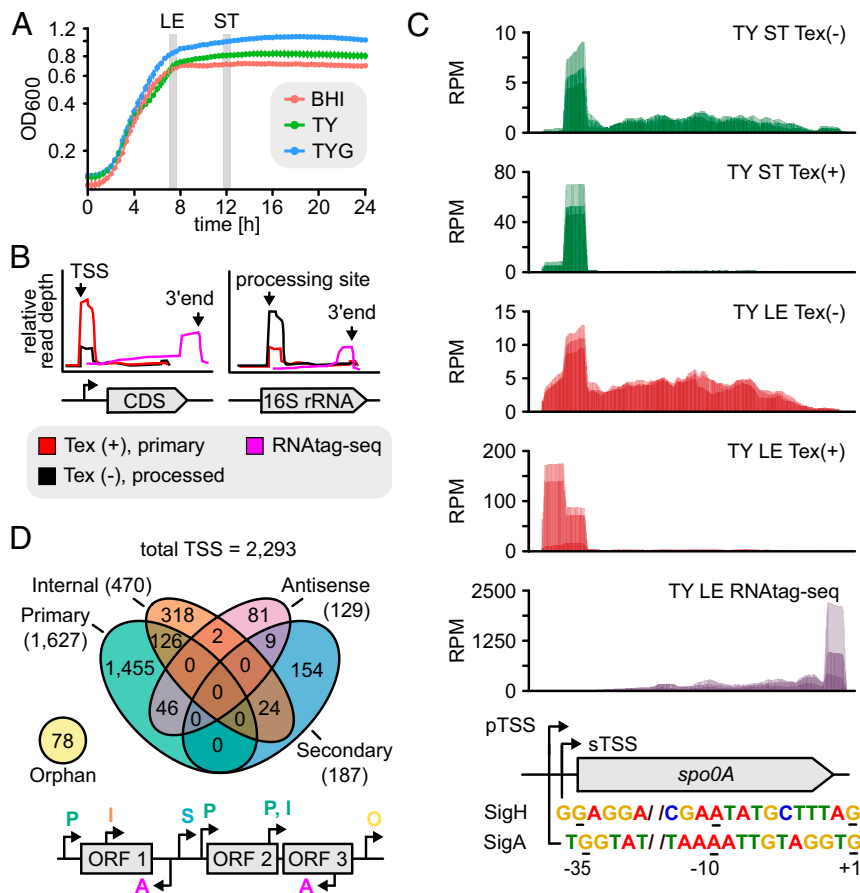


Fig. 1. Global RNA-seq approaches for high-resolution transcriptome mapping. (A) Sequencing samples were generated from strain *C. difficile* 630 grown to late-exponential and stationary phase in tryptone-yeast broth, as well as late-exponential phase in BHI broth. The error bars show SD of three biological replicates. (B) Read profiles generated with dRNA-seq and RNAtag-seq allow the annotation of TSSs and TTSSs. For dRNA-seq, one fraction of total RNA is treated with terminator exonuclease (TEX+), which specifically degrades processed transcripts carrying a 5'-P or 5' OH. The other fraction remains untreated. This differential treatment results in a relative read enrichment for primary transcripts (5'-PPP) in the TEX+ treated libraries allowing the identification of TSSs. For RNAtag-seq, adapters for sequencing are ligated to RNA 3' ends, thereby capturing 3' ends of transcripts. (C) Benchmarking of dRNA-seq approach. Two experimentally determined growth phase-dependent TSSs for *spo0A* are consistent with dRNA-seq-based identification of *spo0A*-associated TSSs. Abbreviations: LE, late-exponential growth phase; ST, stationary growth phase; CDS, coding sequence; TSS, transcriptional start site. (D) Venn diagram showing distribution of TSSs among classes. TSS classification is based on expression strength and genomic location: primary (P), secondary (S), internal (I), antisense (A), and orphan (O).

using specific sporulation medium). Overall, our results add confidence to existing regulon predictions and extend the list of high-confidence candidates that have been repeatedly identified to be associated with a certain sigma factor (indicated in Dataset S3).

We further identified 39 genes with alternative TSSs that were associated with two different promoter sequences. For example, RecA, which controls the DNA damage response by homologous recombinational repair of damaged DNA, is associated with a SigA- and SigB-type promoter supporting its role in stress responses (SI Appendix, Fig. S1C).

Global Mapping of Transcript Ends. To map TTSSs, we adopted the RNAtag-seq protocol (49), a technique utilizing initial adapter ligation to exposed RNA 3' ends. This approach yielded 2,042 experimentally determined TSSs, which were assigned to the following classes: 3' UTR (downstream of CDS or noncoding RNA), 5' UTR (between the TSS and the start codon of CDS), CDS (within a coding sequence), orphan (downstream of an orphan TSS), and CRISPR (associated with a CRISPR array) (Fig. 2A).

The majority of TTSSs mapped to the region downstream of CDSs. Analysis of primary sequences and secondary structures around

these TTSSs reveals a downstream poly-U tail motif for the majority (75%) of them (Fig. 2B and Dataset S4). In addition, a decrease in folding energy (SI Appendix, Fig. S2A) and a spike in guanine-cytosine (GC) content at preceding TSS positions (Fig. 2C) indicate stable stem-loop structures, pointing to an important role for intrinsic transcription termination in *C. difficile*.

Our genome-wide map of 3' UTRs (Dataset S4) reveals that only ~7% of 3' UTR TTSSs are in close distance to a preceding stop codon (<20 nt), whereas 42% of them are >100 nt away (Fig. 2D). This contrasts a recent study in *B. subtilis* that revealed a large fraction of intrinsic terminators overlapping with coding sequences (50). The ability of these terminators to retain efficient transcription termination activity was assigned to runaway transcription of the RNA polymerase, leading to insensitivity to translation-based transcriptional regulation. We explored the original publication, in which a variety of genomes were analyzed in silico, with respect to the presence and location of intrinsic terminators. Reanalysis of the original dataset for *C. difficile* confirms our experimental results that intrinsic terminators rarely overlap with stop codons (SI Appendix, Fig. S2B), suggesting that runaway transcription is not a determining feature of gene expression in *C. difficile*.

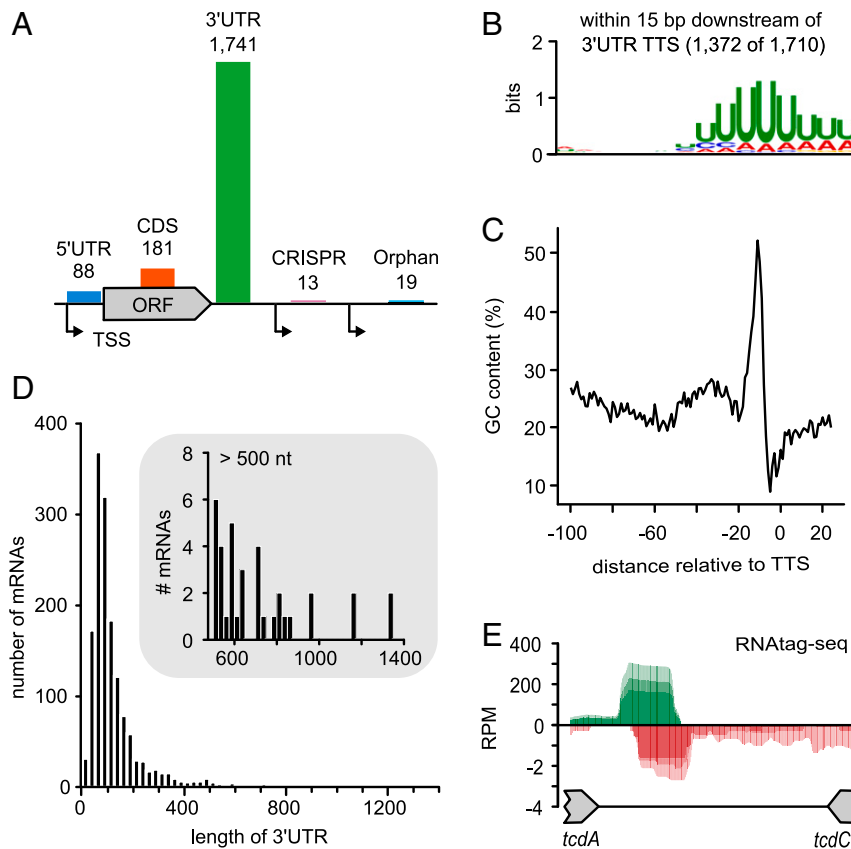


Fig. 2. Global features associated with transcription termination sites (TTSs). (A) Classification of TTS based on their genomic location: 3' UTR (downstream of CDS or noncoding gene), 5' UTR (within the 5' UTR of a coding sequence), CDS (within coding sequence), orphan (downstream of orphan TSS), and CRISPR (associated with CRISPR array). The numbers indicate the amount of annotated TTSs for each class. (B) Consensus motif associated with 3' UTR-located TTSs downstream of coding sequences was identified using the MEME suite. The identified motif is located within 15 bp downstream of detected 3' UTR TTS positions. (C) Calculation of average GC content (%) at each indicated position relative to 3' UTR-located TTSs reveals a spike in GC content immediately upstream of the TTS, indicating a stable stem structure. (D) Frequencies of 3' UTR lengths based on 1,741 detected 3' UTR TTSs. (E) Reads in the RNAtag-seq libraries reveal overlapping TTSs for the convergently transcribed genes *tcdA* and *tcdC*.

We further detected many overlapping termination events between convergently transcribed genes, such as for the toxin gene *tcdA* (*toxA* in CP010905.2 which was used in this study) and the negative regulator of toxin gene expression *tcdC* (Fig. 2E), which is also a pervasive transcriptome feature in *Escherichia coli* (51). Overall, our UTR annotations revealed an abundance of overlapping transcripts between convergently transcribed genes (246 gene pairs or 32.06% of transcripts with TTSs), whereas overlaps between divergently transcribed genes were rare (8 or 1.17% of transcripts with TSSs) (Dataset S8).

Annotation of Small ORFs and Operons. High-resolution transcriptome maps allow the identification of ORFs and operon structures that have been overlooked in automated genome annotations. This includes so called small ORFs (sORFs) of usually 50 amino acids or less, an emerging class of bacterial genes with an unfolding spectrum of biological functions (52, 53). In many cases, they are predicted to be membrane proteins, containing an alpha-helical transmembrane domain (54). Focusing on oTSS in particular, we searched for novel ORFs based on the following criteria: 1) presence of a start and stop codon, 2) presence of an RBS within 15 base pairs (bp) upstream of the start codon, and 3) sequence conservation in other *Clostridioides* strains. Based on this approach, we identified 12 sORF candidates, seven of which are predicted to contain a transmembrane helix (Dataset S2). Among the identified candidates, six are toxins and part of previously identified *C. difficile* type-I toxin-antitoxin systems (55, 56).

Among the remaining sORFs, four are high-confidence candidates; one being a conjugal transfer protein with annotation in other *C. difficile* strains; two candidates that each have a Shine-Dalgarno (SD) sequence and a predicted α -helical transmembrane domain; and one sORF associated with a 160-nt long 5' UTR region harboring a c-di-GMP-I riboswitch.

Transcriptome studies have revealed operon structures more complex than previously anticipated (51, 57, 58), but predicted operons in *C. difficile* are still largely based on computational inference from other bacterial species (59). Here, our transcriptome-based annotation predicts 440 operons (Dataset S1), including 64 previously unknown operons such as the dicistronic transcript comprising spermine/spermidine acetyltransferase *bltD* and CDIF630_01360, a putative N-acetyltransferase. Furthermore, we identify 109 suboperon structures which, in 29 cases, uncouple the last or the last two genes from upstream genes within an operon. One example is *acpS-ndoA*, in which the last two genes encoding a type-II toxin-antitoxin system (*ndoA* and *ndoA1*) are uncoupled from the full operon.

5' UTRs and Associated Regulatory Elements. Bacterial 5' UTRs can influence gene expression, usually through embedded *cis*-regulatory elements such as riboswitches, RNA thermometers, and genetic switches. Similar to *B. subtilis* and *Listeria monocytogenes*, the majority of experimentally mapped 5' UTR lengths ranged from 20 to 60 nt (Fig. 3A and Dataset S1) (14, 40, 60). A likely RBS (aGGAGg) was detected in ~90% of these 5' UTRs (Fig. 3A, Inset).

Only six mRNAs appeared leaderless, having a 5' UTR of <10 nt, including the *spoVAE* gene within the tricistronic *spoVACDE* operon (61).

Many genes (561) were associated with surprisingly long 5' UTRs of >100 nt. Among those having a very long 5' UTR of >300 nt, we identified 77 Rfam-predicted riboswitch candidates (62) (Fig. 3B). Premature transcription termination (PTT) was evident for 57 of them, indicating their ON state in which transcription of the parental gene is repressed (Dataset S5). Interestingly, we detected a *speF* riboswitch previously only documented in gram-negative alphaproteobacteria associated with CDIF630_01955, encoding a methyltransferase domain protein. As before, the associated termination site suggests that the riboswitch is functional, which is further supported by a 220-nt Northern blot signal (SI Appendix, Fig. S3).

We further detected 17 putative PTT events in 5' UTRs lacking similarities to conserved riboregulators. A potential generation from mRNA processing is unlikely in these cases, since the characteristic read enrichment in the untreated (TEX-) cDNA library, usually associated with processing sites, is missing. One such PTT event is located in the 5' UTR of the *infC-rpmI-rplT* operon whose expression is controlled through an L20 leader via transcription attenuation in *B. subtilis* (63, 64). This autoregulatory structure is also found in other low-GC gram-positive bacteria (Rfam family RF00558) but does not seem to be conserved on the primary sequence level in *C. difficile*. Nevertheless, secondary structure

prediction reveals extensive interactions between distant bases that is reminiscent of the antiterminator conformation of the L20 leader described in *B. subtilis* (Fig. 3C). Furthermore, PTT-associated genes encoded phosphotransferase systems (*bglF*, *bglF1*, *bglG3*, and *bglG4*) known to be regulated by anti-terminator proteins in *B. subtilis* (65), transcriptional regulators (CDIF630_00097, CDIF630_02384, and CDIF630_02922), multidrug resistance (MDR)-type ABC transporters (CDIF630_03083, CDIF630_02847, and CDIF630_03664), and *aroF* (Dataset S5). Northern blot validation for a selection of candidates confirmed the RNA-seq-predicted transcript sizes and revealed larger bands in several cases that are likely corresponding to the full-length parental gene or its degradation products (SI Appendix, Fig. S3).

The sRNA Landscape of *C. difficile* in Changing Environmental Conditions and during Infection.

Our primary transcriptome analysis identified 42 transcripts that lacked an internal ORF, qualifying them as potential sRNAs (Dataset S5). A classification based on their genomic location (Fig. 4A) revealed the largest group to be 3' UTR-derived sRNAs (18 3' UTR), followed by those encoded cis-antisense either to a gene, a 5'/3' UTR, or another sRNA (13 AS). In comparison, only few sRNAs were located in intergenic regions (8 IGR) or derived from the 5' UTR of mRNAs (3 5' UTR). Sequence conservation beyond *C. difficile* was extremely rare (SI Appendix, Fig. S4A and Dataset S7), whereas most are highly conserved among clinical strains of *C. difficile* covering five major genomic clades and

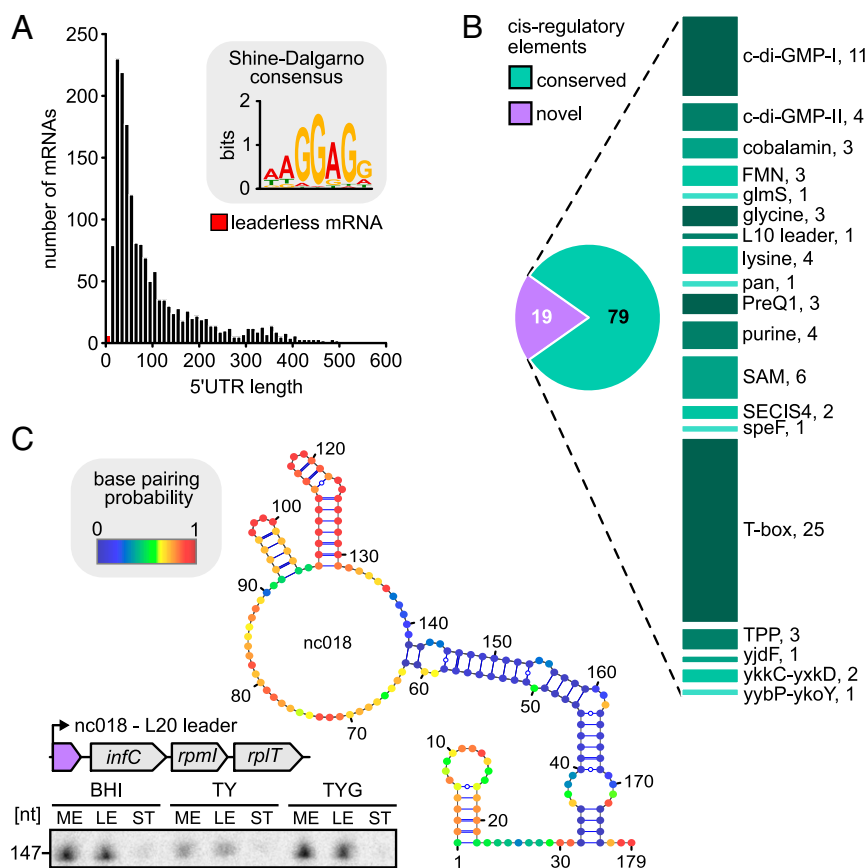


Fig. 3. Discovery of *cis*-regulatory elements in 5' UTR regions. (A) Frequencies of 5' UTR lengths based on 1,646 primary and secondary TSSs. The red bars indicate six leaderless mRNAs with a 5' UTR length of <10 nt. The *Inset* shows the predicted SD sequence motif of *C. difficile* 630. (B) Classification of PTT events in 5' UTRs of mRNAs. The majority is associated with conserved RNA families. The remaining 19 PTT events lack homology to known riboregulators and are classified putative regulators. (C) Putative L20 leader in the 5' UTR of the *infC* operon. (Top) Predicted secondary structure for CDIF630nc_018 using RNAfold. Nucleotides are colored according to base-pairing probabilities. (Bottom) Schematic overview of the *infC* operon organization including 5' UTR region and expression analysis of CDIF630nc_018 by Northern blot using a radioactively labeled DNA probe. Total RNA was extracted at midexponential (ME), late-exponential (LE), and stationary (ST) growth phases from *C. difficile* 630 grown in BHI, TY, and TY supplemented with 0.5% glucose (TYG) media.

diverse toxin repertoires (*SI Appendix, Fig. S4B*) (66). Some of the sRNAs that were absent in a subset of strains were either transposon-associated sRNAs (CDIF630nc_004 and CDIF630nc_069) or located on a prophage (CDIF630nc_095). Of note, this high level of sequence conservation within the species generally extended to noncoding 5' and 3' UTR regions (92% of 5' UTRs with >90% identity and 90% of 3' UTRs with >90% identity, *Dataset S9*).

Northern blot validation revealed expression throughout exponential growth for most sRNA candidates (*Fig. 4B*). About one-half of them are down-regulated after entry into stationary phase, while three candidates (CDIF630nc_028, CDIF630nc_089, and

CDIF630nc_105) showed a marked accumulation. In agreement with the expression profile, CDIF630nc_028 has a predicted promoter sequence for SigB, while CDIF630nc_089 is associated with a SigH promoter. Moreover, Northern blot validation of sRNA expression in the hypervirulent RT027 isolate R20291 revealed identical expression profiles in both strains for the selected candidates (*SI Appendix, Fig. S5*).

To gain insight into sRNA activities during infection, we took advantage of published *in vivo* transcriptome data from *C. difficile*-infected ex-germ-free mice as well as conventional mice pretreated with different antibiotics (67). Reanalysis of the data

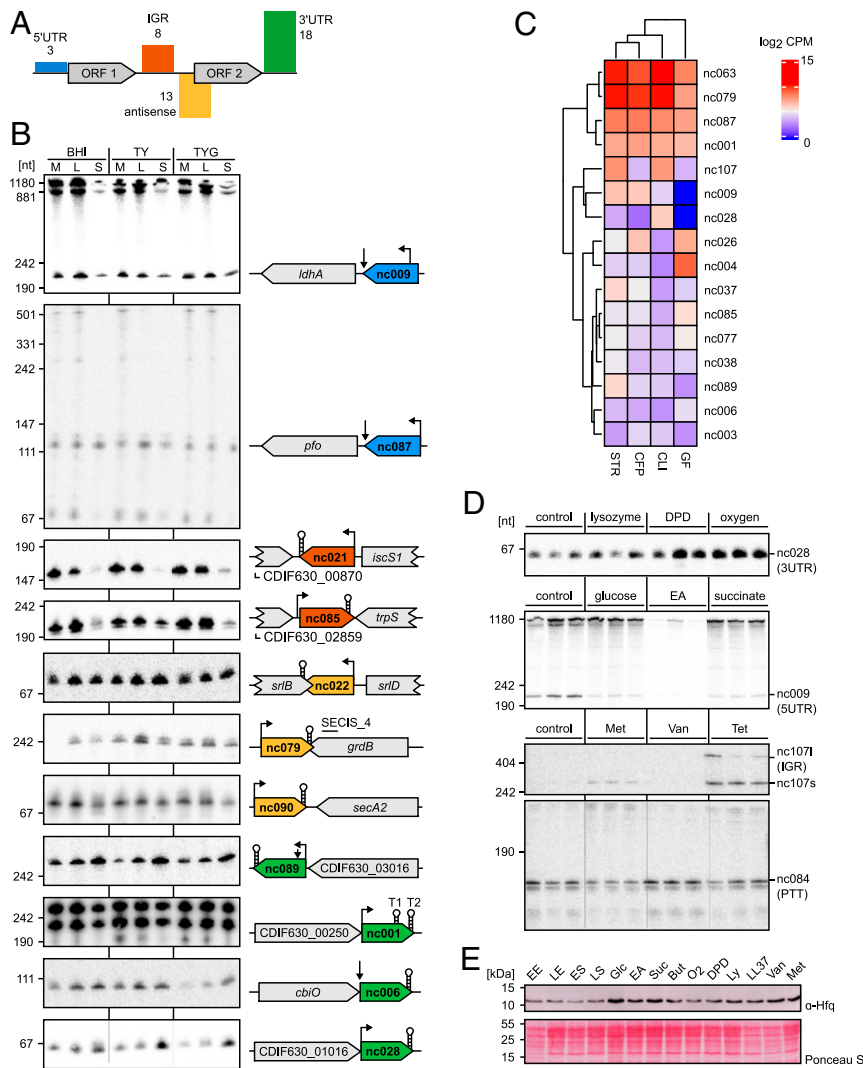


Fig. 4. *C. difficile* 630 sRNA landscape in changing environmental conditions. (A) Classification of annotated sRNA candidates based on their genomic location. The numbers indicate amount of annotated sRNAs for each class. (B) Expression profiles of representative candidates for each sRNA class. Total RNA was extracted at midexponential (ME), late-exponential (LE), and stationary (ST) growth phases from *C. difficile* 630 grown in TY, TY supplemented with 0.5% glucose (TYG), and BHI medium and analyzed by Northern blot using radioactively labeled DNA probes. (C) Heatmap of log₂ CPM values for newly annotated sRNAs during *C. difficile* 630 intestinal colonization of different antibiotic-treated mouse models (Streptomycin, STR; Cefoperazone, CFP; Clindamycin, CLI) and exGerm-free mice (GF). Published data were taken from ref. 67 and reanalyzed using the updated genome annotation. Only sRNAs with a CPM value ≥50 were considered to be expressed. (D) Expression profiles of sRNAs that are differentially expressed in response to various nutritional and stress conditions. Total RNA was extracted from midexponential TY cultures of *C. difficile* 630 WT exposed to TY (control), lysozyme, iron limitation (2,2'-Dipyridine, DPD), oxygen, glucose, ethanolamine (EA), succinate, metronidazole (Met), vancomycin (Van), and tetracycline (Tet) for 20 min and analyzed by Northern blot using radioactively labeled DNA probes. (E) Western blot analysis of Hfq protein levels across different growth phases and growth media. Equal optical density (OD) units of total cell lysates of *C. difficile* 630 WT were loaded. Western blot membranes were incubated with anti-Hfq antibody. Ponceau S staining of the blotting membrane served as loading control. Total cell extracts of equal OD units were prepared from *C. difficile* 630 WT grown in TY medium to early exponential (EE), late-exponential (LE), early stationary (ES), and late stationary (LS) phase and from bacteria grown to late-exponential phase in TY medium supplemented with glucose (Glc), ethanolamine (EA), succinate (Suc), butyrate (But), 2,2'-Dipyridine (DPD), lysozyme (Ly), LL-37, vancomycin (Van), metronidazole (Met), or in 1% oxygen atmosphere (O₂). Precise concentrations used are listed under *SI Appendix, Materials and Methods*.

revealed expression of 16 sRNAs (cutoff = 50 counts per million mapped reads [CPM]), including the most broadly conserved sRNAs CDIF630nc_001, CDIF630nc_026, and CDIF630nc_079 as well as the SigB- and SigH-associated CDIF630nc_028 and CDIF630nc_089, respectively (Fig. 4C and Dataset S10). To identify infection-related signals regulating sRNA expression, we exposed *C. difficile* to a panel of nutritional and stress conditions including intestinal metabolites (glucose, ethanolamine, succinate, and butyrate), oxygen stress, iron limitation, as well as antimicrobial compounds (lysozyme, LL-37, vancomycin, metronidazole, and tetracycline). Within this panel, we observed regulation for several sRNAs or PTTs (Fig. 4D), e.g., SigB-regulated CDIF630nc_028 responded to iron limitation and oxygen stress, suggesting a potential role during infection. To complement the picture, we also quantified native protein levels of the sRNA chaperone Hfq during growth in these conditions (Fig. 4E). We observed similar Hfq levels not only in different growth phases but also under a variety of nutritional and stress conditions, demonstrating that Hfq is constitutively produced in changing environmental conditions.

C. difficile Hfq Is a Global RNA Chaperone. In many gram-negative model organisms, sRNAs require the RNA chaperone Hfq to facilitate their interaction with target mRNAs (68). However, whether Hfq functions as a central RNA-binding protein (RBP) in *C. difficile* remains unknown. Therefore, we performed Hfq immunoprecipitation followed by sequencing of bound RNA species (RIP-seq) in a strain expressing C-terminally 3×FLAG-tagged *hfq* (Hfq-FLAG) under its native promoter to draft the spectrum of Hfq-bound RNAs in vivo. Bacteria were grown to midexponential, late-exponential, and stationary phase (SI Appendix, Fig. S6A) and subjected to the RIP-seq protocol (SI Appendix, Fig. S6B). Western blot validation using monoclonal FLAG antibody confirmed expression of Hfq throughout all growth phases and specific enrichment of Hfq-FLAG from *C. difficile* 630 lysate (SI Appendix, Fig. S6C). In addition, copurification of Hfq-bound sRNAs was confirmed by Northern blot analysis of lysates and eluate fractions from Hfq-FLAG and Hfq-ctrl cultures (SI Appendix, Fig. S6C).

The majority of RNA species coimmunoprecipitating with Hfq-3×FLAG were mRNAs (Fig. 5A and Dataset S6). Hfq bound mostly to the 5' and 3' UTRs of target mRNAs, and of note, many enriched 5' UTRs harbored a riboswitch or PTT (66/124 5' UTRs) (Fig. 5A) as recently observed in an independent Hfq RIP-seq analysis (20). Many mRNAs showed growth phase-dependent association with Hfq (SI Appendix, Fig. S7B and C) and mainly included genes involved in the biosynthesis of amino acids, membrane transport, signal transduction, and translation. Most interestingly, we also found several mRNAs encoding proteins involved in sporulation (*spo0A*, *spoVB*, *oppB*, *sigG*, and *sspA2*), toxin production (*tcdE*), motility (*flgB*, *fliJ*, *flhB*, and *cheW1*), quorum sensing (*luxS* and *agrB*), the cell wall protein-encoding *cwpV*, as well as several adhesins potentially involved in biofilm formation (CDIF630_03429, CDIF630_01459, and CDIF630_03096) (SI Appendix, Fig. S7C and Dataset S6).

The enrichment of many 5' UTRs could suggest that Hfq is recognizing the SD or ATG sequence region, but lack of clear Hfq peaks at SD or start codon sites does not sufficiently support this assumption at this point. In addition, the enrichment of many 3' UTRs and of 5' UTRs harboring a riboswitch or PTT event could point to a potential role of terminator hairpins in Hfq binding. An RNA motif search using CMfinder identified an RNA structure reminiscent of a terminator hairpin that was enriched in Hfq-bound transcripts, albeit nucleotide identity was low (SI Appendix, Fig. S7A).

Importantly, many predicted sRNAs associated with Hfq, including the previously identified Hfq-binding sRNA CDIF630nc_070 (also known as Rcd1, Fig. 5A and Dataset S6) (18, 28). In addition, all but one type-I antitoxin transcript were enriched (Fig. 5A and

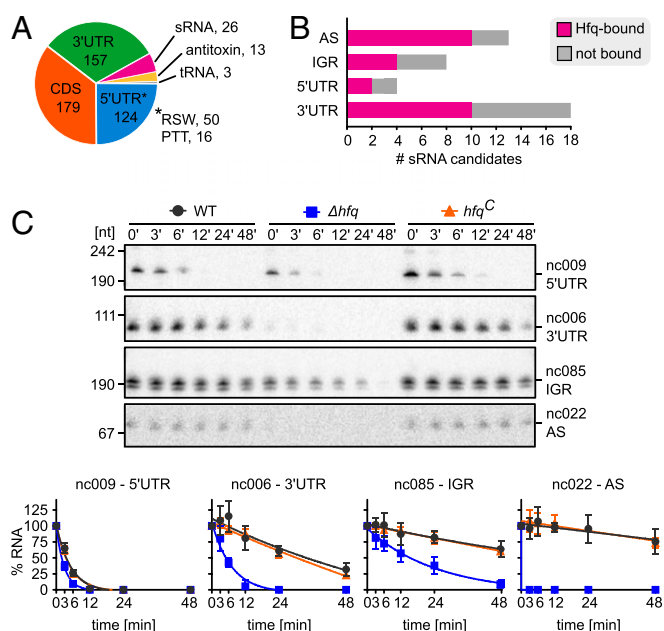


Fig. 5. The spectrum of Hfq-associated RNA ligands and Hfq impact on RNA stability. (A) Pie chart for Hfq RIP-seq showing the relative amount of Hfq-associated sequences mapping to different RNA classes. Hfq RIP-seq was performed on *C. difficile* 630 WT and C-terminally 3×FLAG-tagged *hfq* expressed from a plasmid, grown to midexponential, late-exponential, and early stationary phases in TY medium in two independent experiments. (B) Bar chart showing the fraction of sRNAs for each class that were bound by Hfq in vivo. (C) Hfq acts as a stabilizing factor for its ncRNA ligands in vivo. (Top) Total RNA was extracted from WT, Δhfq , and the complemented *hfq* mutant (*hfq*^C) in late-exponential phase after addition of rifampicin at the indicated time points and analyzed by Northern blotting using radioactively labeled DNA probes. The Hfq-independent sRNA nc009 served as negative control. The representative of three independent Northern blots is shown. Abbreviations: RSW (riboswitch), AS (antisense), IGR (intergenic).

Dataset S6), similar to a recent RIP-seq analysis of Hfq in *C. difficile* (20). Classifying Hfq-associated sRNAs with respect to their genomic location revealed that the majority of cis-antisense-encoded sRNAs in *C. difficile* were bound by Hfq, as well as approximately one-half of all intergenic, 5' UTR-, and 3' UTR-encoded sRNAs (Fig. 5B). Additionally, many sRNAs were bound across all three growth phases, with CDIF630nc_070 and CDIF630nc_079 being the two top-enriched sRNAs (SI Appendix, Fig. S6D and Dataset S6). However, a few sRNAs displayed growth phase-dependent association with Hfq, such as CDIF630nc_090, which was only enriched in stationary phase (SI Appendix, Fig. S6D).

Hfq Acts as a Stabilizing Factor for its noncoding RNA Ligands. One major function of gram-negative Hfq homologs is the stabilization of their sRNA ligands, which appears to be the exception in gram-positive bacteria (69–71). To test if transcript stabilities of Hfq-associating noncoding RNAs (ncRNAs) were affected by Hfq, we generated a *hfq* deletion (Δhfq) in *C. difficile* 630 using a homologous recombination vector we developed for effective counterselection on standard laboratory media using the tightly regulated *E. coli* endoribonuclease *mazF* (for details, see SI Appendix). To complement *hfq* in cis (*hfq*^C), we established an additional vector system which enables efficient insertion of DNA by homologous recombination between *pyrE* (CD630_01870) and CD630_01880.

C. difficile wild type (WT), Δhfq , and *hfq*^C strains were grown to late-exponential phase, and rifampicin was added to arrest transcription. Northern blotting of RNA samples collected at several timepoints after rifampicin addition was performed to

quantify transcript levels for several sRNAs as well as a Hfq-bound riboswitch and type-I antitoxin. Remarkably, half-lives and, in some cases, the steady-state levels of all analyzed Hfq-associated sRNAs (CDIF630nc_006, CDIF630nc_085, and CDIF630nc_022) were decreased, an effect that could be restored in the *hfq*^C strain (Fig. 5C and SI Appendix, Fig. S8 A and B) and that was not observed for the Hfq-independent sRNA CDIF630nc_009. In

addition, half-lives were also reduced for the tested type-I antitoxin as well as the c-di-GMP-II riboswitch, suggesting a potential dual function for the latter *cis*-acting regulator (SI Appendix, Fig. S8B).

A Role of CDIF630nc_085 in the Regulation of Ethanolamine Utilization.

To gain insight into sRNA functions, we chose the bona fide intergenic sRNA CDIF630nc_085 (short nc85) (Fig. 4B) for further

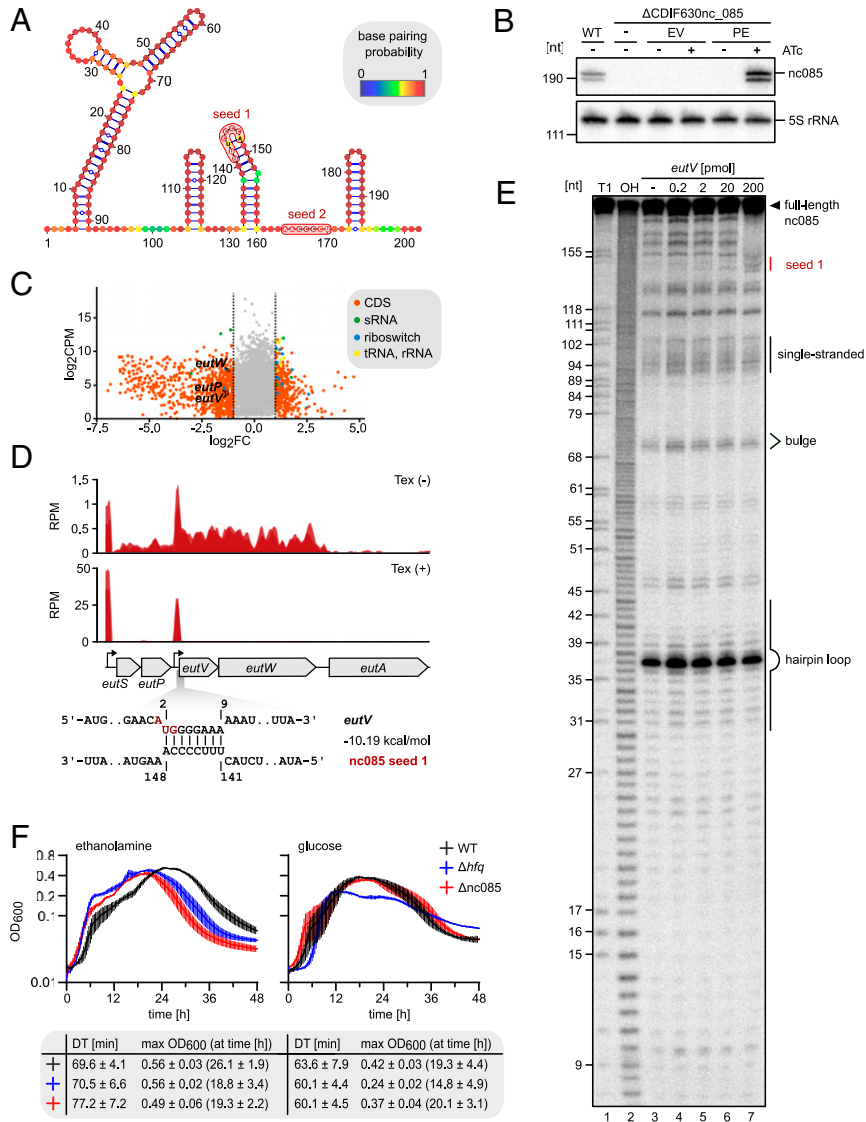


Fig. 6. CDIF630nc_085 regulates ethanolamine utilization in *C. difficile*. (A) Predicted secondary structure for CDIF630nc_085 (nc85) using RNAfold. Nucleotides are colored according to base-pairing probabilities. Two seed regions predicted to interact with differentially expressed mRNA targets are located in an extended small stem loop (seed region 1) and a single-stranded stretch (seed region 2) in the 3' region of the sRNA, respectively. (B) Northern blot validation of nc85 pulse expression in a nc85 deletion strain. sRNA expression was transiently induced for 20 min by addition of 100 ng/mL anhydrotetracycline (ATc) to late-exponential cultures of *C. difficile* WT, Δ nc85 (-), Δ nc85 harboring the empty pRPF185 backbone (EV), and Δ nc85 harboring the pulse-expression vector carrying nc85 (PE) under the control of an ATc-inducible promoter. (C) RNA was extracted from ATc-induced cultures harboring the empty vector (EV) or the pulse-expression plasmid (PE) and analyzed by RNA-seq. The genes that were differentially expressed upon pulse expression of nc85 (false discovery rate ≤ 0.1 , \log_2 fold change [\log_2 FC] ≥ 1 , or \log_2 FC ≤ -1) are colored according to their transcript class. (D) Read profiles from untreated (Tex-), Tex-treated (Tex+), and RNAtag-seq libraries mapping to the *eut* region. TSSs were identified upstream of *eutS* and *eutV*. An RNA duplex formed by nc85 with the start codon region of *eutV* was predicted using IntaRNA. The red letters indicate the start codon. (E) In-line probing of 0.2 pmol of ³²P-labeled nc85 in the absence (lane 3) or presence of either 0.2 pmol (lane 4), 2 pmol (lane 5), 20 pmol (lane 6), or 200 pmol (lane 7) *eutV* target region. Partially RNase T1- (lane 1; T1) or alkali-digested (lane 2; OH) nc85 served as ladders. Seed 1 (red) denotes the seed region that is protected from cleavage changes in the presence of the target. A representative image of two independent experiments is shown. (F) Growth of *C. difficile* 630 WT, Δ hfq, and Δ nc85 in CDMM supplemented with either glucose (5 mM) or ethanolamine (5 mM). Growth experiments were performed in two independent experiments with three biological replicates each ($n = 6$). The error bars show SEM. Bottom table: listed are the calculated doubling times (DT) \pm SD as well as the maximum optical densities (OD₆₀₀) and the timepoint when they were reached during the experiment, for each strain and each growth condition. Due to the complex, diauxic growth behavior of all strains in the analyzed growth conditions and doubling times were calculated only for the first exponential phase.

characterization because its expression during intestinal colonization (Fig. 4C) suggested a potential virulence function. Secondary structure prediction showed a moderately folded structure with an extended stem loop at the 5' end, two hair pins in the center of the sRNA, and a Rho-independent terminator at the 3' end (Fig. 6A). To identify potential mRNA targets of nc085, we pulse expressed the sRNA from an anhydrotetracycline-inducible plasmid in a Δ CDIF630nc_085 background (Fig. 6B) and determined differentially expressed genes using RNA-seq analysis (Fig. 6C and Dataset S11). We observed many sporulation-associated genes to be down-regulated upon nc085 expression (Dataset S11). In addition, gene cluster analysis revealed ABC transporters and two-component systems as the largest group of regulated genes (SI Appendix, Fig. S9B). Searching for base-pairing regions within the 5' UTRs of differentially expressed genes identified two potential seed regions within nc085 (Fig. 6A, SR1 and SR2), both complementarity to several target mRNAs (SI Appendix, Fig. S9A).

Among them was *eutV*, which is part of a two-component system encoded by the *eutVW* operon, a positive regulator of ethanolamine utilization. Our transcriptome data identified low-level constitutive expression across all growth conditions and a TSS upstream of the *eutVW* operon (Fig. 6D, data only shown for late-exponential TY condition). Moreover, the predicted base-pairing region within the *eutV* 5' UTR comprised the start codon site of *eutV* (Fig. 6D).

To test the prediction experimentally, we performed in-line probing of nc085 with or without the *eutV* 5' UTR. Although high concentrations of the target mRNA were required, changes in the protection from cleavage in the presence of *eutV* were observed (Fig. 6E, lane 7) and provided support for the computationally predicted base-pairing region in nc085. Considering the location of the seed region within a hairpin structure, this interaction may need to be facilitated by Hfq in vivo. To explore the impact of nc085 on ethanolamine utilization, we compared growth of *C. difficile* WT, Δ nc085, and the Δ hfq mutant in *C. difficile* minimal medium (CDMM) supplemented either with glucose or ethanolamine. Cultivation in glucose-supplemented medium resulted in identical growth for WT and Δ nc085, whereas Δ hfq had an extended lag phase and grew to lower maximum optical densities. In contrast, when grown in CDMM supplemented with ethanolamine, both Δ nc085 and Δ hfq exhibited accelerated growth in comparison to the WT (Fig. 6F). Specifically, both strains entered exponential growth faster and reached their maximum optical densities at earlier timepoints (Fig. 6F, Bottom). Together, these data identify a regulatory function of nc085 in the utilization of the abundant intestinal carbon and nitrogen source ethanolamine, which was shown to impact pathogenicity of *C. difficile* (72).

Discussion

In the present study, we have applied global approaches of bacterial RNA biology (35) to capture the transcriptome architecture of *C. difficile* and unravel the scope of posttranscriptional regulation in this important human pathogen.

A Gram-Positive Hfq Homolog that Impacts sRNA Stabilities and Function. It is undisputed that sRNAs are central players in the regulation of physiological and virulence pathways in gram-positive bacteria including *L. monocytogenes*, *B. subtilis*, and *Staphylococcus* species. For example, the sRNA RsaE, which is conserved in the order Bacillales, has been shown to coordinate central carbon and amino acid metabolism in *Staphylococcus aureus* (73, 74) and also to promote biofilm formation in *Staphylococcus epidermidis* by supporting extracellular DNA release and the production of polysaccharide-intercellular adhesin (75). However, the function of Hfq in facilitating the interactions between sRNA regulators and their target mRNAs in gram-positive bacteria remains elusive

since most of the known sRNAs seem to exert their regulatory activity independent of this global RBP (76).

In the present study, our characterization of Hfq-sRNA interactions in *C. difficile* not only revealed a large network of Hfq-binding sRNAs along with potential target mRNAs but also provides clear evidence for the impact of Hfq on sRNA steady-state levels and stabilities in vivo. Our data align with initial studies using a Hfq-depleted *C. difficile* strain that identified pleiotropic effects of Hfq depletion on gene expression using microarray analysis (18). In line with these observations, absence of Hfq impacts growth on various carbon and nitrogen sources, as shown in our study, and results in increased sensitivity to stresses, sporulation rates, and biofilm formation (18). Taken together, the regulatory activities of Hfq in *C. difficile* strongly resemble the situation in gram-negative model organisms such as *E. coli* and *Salmonella enterica* and open avenues for the study of sRNA-based gene regulation in this important pathogen.

Ethanolamine Utilization Is Governed by sRNA Regulation in Gram-Positive Pathogens. We have taken a first step toward the characterization of sRNA-mRNA interactions in *C. difficile* by identifying the ethanolamine utilization pathway to be regulated by the intergenic sRNA nc085. Ethanolamine is an abundant intestinal nutrient that is exploited by both gram-negative and gram-positive enteric pathogens, including *C. difficile*, during intestinal colonization (72, 77–79). Interestingly, the ethanolamine utilization (*eut*) gene cluster is regulated by a small regulatory RNA in both *Enterococcus faecalis* (EutX) and *L. monocytogenes* (Rli55). In both species, the sRNA contains a classic AdoCbl-responsive riboswitch in its 5' portion that allows expression of the full-length sRNA only in the absence of vitamin B₁₂ (80, 81). This full-length transcript harbors a dual-hairpin structure in its 3' part that serves as a binding platform for the ethanolamine-specific, RNA-binding response regulator EutV (82), thereby sequestering EutV from binding to its target sequences in the 5' UTRs of *eut* genes. An in silico search for an AdoCbl riboswitch failed to identify such a conserved element within nc085, and secondary structures of the three sRNAs show a less-complex folding pattern for nc085 that is missing the dual hairpin for EutV binding. Overall, this suggests that the mechanism of *eut* gene regulation by nc085 differs from EutX and Rli55. Currently, down-regulation of the *eutVW* operon upon nc085 pulse expression together with a predicted target site comprising the start codon of *eutV* suggests a classical mode-of-action whereby nc085 binding interferes with translation initiation. Consequently, deleting nc085 should relieve the repression and increase the pool of EutVW that can sense ethanolamine and activate the *eut* gene cluster. Aligning with this model, we observed early onset of growth for the nc085 deletion strain in CDMM with ethanolamine as the sole carbon source (Fig. 6F).

Overall, our findings further confirm that ethanolamine utilization is a conserved target of sRNA-based regulation in gram-positive pathogens.

Materials and Methods

Comprehensive descriptions of all materials and methods that were used in this study are provided in SI Appendix, Materials and Methods.

Data Availability. All RNA-sequencing data are available at the National Center for Biotechnology Information Gene Expression Omnibus database (<https://www.ncbi.nlm.nih.gov/geo>) under the accession number GSE155167. Plasmids pKJAK112, pJAK184, and pJAK080 have been deposited with Addgene (167279–167281).

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