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To cite this version:
Kim Boi Le Huyen, Cintia Daniela Gonzalez, Gaetan Pascreau, Valérie Bordeau, Vincent Cattoir, et al.. A small regulatory RNA alters Staphylococcus aureus virulence by titrating RNAIII activity. Nucleic Acids Research, 2021, 49 (18), pp.10644-10656. 10.1093/nar/gkab782. hal-03369833

HAL Id: hal-03369833
https://hal.science/hal-03369833
Submitted on 30 May 2022

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A small regulatory RNA alters *Staphylococcus aureus* virulence by titrating RNAIII activity

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Received April 06, 2021; Revised August 25, 2021; Editorial Decision August 26, 2021; Accepted September 02, 2021

ABSTRACT

*Staphylococcus aureus* is an opportunistic human and animal pathogen with an arsenal of virulence factors that are tightly regulated during bacterial infection. The latter is achieved through a sophisticated network of regulatory proteins and regulatory RNAs. Here, we describe the involvement of a novel prophage-carried small regulatory *S. aureus* RNA, SprY, in the control of virulence genes. An MS2-affinity purification assay reveals that SprY forms a complex *in vivo* with RNAIII, a major regulator of *S. aureus* virulence genes. SprY binds to the 13th stem-loop of RNAIII, a key functional region involved in the repression of multiple mRNA targets. mRNAs encoding the repressor of toxins Rot and the extracellular complement binding protein Ecb are among the targets whose expression is increased by SprY binding to RNAIII. Moreover, SprY decreases *S. aureus* hemolytic activity and virulence. Our results indicate that SprY titrates RNAIII activity by targeting a specific stem loop. Thus, we demonstrate that a prophage-encoded sRNA reduces the pathogenicity of *S. aureus* through RNA sponge activity.

INTRODUCTION

*Staphylococcus aureus* is an opportunistic human and animal pathogen that can cause a wide range of illnesses, from food poisoning and superficial abscesses to more life-threatening diseases such as pneumonia, osteomyelitis, bacteremia, endocarditis and toxic shock syndrome (1,2). The infection process requires the controlled expression of virulence factors allowing bacteria to escape the host defense system, adapt to changing environmental conditions, and attack and destroy host cells. This process involves a wide range of wall-associated proteins and extracellular factors that are expressed during the different stages of infection. Global regulatory elements including transcription factors, sigma factors, two-component systems and regulatory RNAs assure the coordinated expression of these virulence factors.

Among the regulatory RNAs, small RNAs (sRNAs) are mostly non-coding, relatively short (50–550 nucleotides), and located within core genomes or mobile genetic elements. They are usually conditionally expressed, i.e. depending upon specific stress and growth phase. sRNAs control expression of their target genes by pairing with RNAs or forming complexes with proteins; in this way they usually modulate the stability and translation of mRNAs, and modify the activity of proteins (3). Over the past decade, hundreds of putative sRNAs discovered in *S. aureus* were compiled in the *S. aureus* RNA Database SRD (4). However, a recent analysis proposes that among them, only 50 are ‘bona fide’ sRNAs (5). *Staphylococcus aureus* sRNAs were demonstrated to contribute to regulation of dozens of functions. sRNAs contribute to regulation of bacterial metabolism, such as RsaE (6–8), or to antibiotic resistance, such as SprX (9). Several sRNAs were shown to be involved in the virulence of *S. aureus* such as SprD, which regulates expression of the immune evasion protein Sbi (10).

RNAIII is a paradigm for sRNA regulation of virulence (11,12). Besides encoding the PSM 6-hemolysin (known as Hld), RNAIII also positively regulates expression of *hla*, encoding the α-hemolysin (13), which leads to cell lysis (14–16), and intervenes in the expression switch between surface proteins and secreted toxins. Through direct binding to mRNA targets, RNAIII prevents the translation of major surface proteins, such as protein A (17), Sbi (10) and Ecb (18), which play key roles in adhesion and immune evasion. In addition, RNAII inhibits translation of *rot* mRNA, encoding the repressor of toxins Rot (19–21), which blocks the transcription of exoproteins and toxins (22). By inhibiting Rot, RNAIII indirectly activates transcription of exotoxins.
and indirectly inhibits synthesis of protein A at the transcriptional level.

In this study, we report that SprY, an sRNA expressed from prophage φ12, is involved in the regulation of *S. aureus* virulence. By antisense pairing with RNAIII, SprY prevents RNAIII from regulating its targets and consequently decreases *S. aureus* hemolytic activity and virulence in a murine sepsis infection model. Together, our data reveal an sRNA acting as a sponge for RNAIII and further elucidate the regulation network controlling *S. aureus* virulence.

**MATERIALS AND METHODS**

**Bacterial strains and growth condition**

All bacterial strains and plasmids used in this work are listed in Supplementary Table S1. The DH5-α *Escherichia coli* strain was grown at 37°C in Luria-Bertani (LB) broth or LB agar plates supplemented with 50 μg/ml ampicillin if necessary. The *S. aureus* RN4220 strains was used to prepare phage-containing vectors expressing sRNA or target-gfp fusions. In this study, *S. aureus* HG003 strain was used to co-transform the target-gfp fusions with the sRNA expressing plasmid. Cultures of these co-transformed *S. aureus* strains were grown at 37°C either in brain heart infusion broth (BHI, Oxoid) or on BHI agar plates. When necessary, media were supplemented with 10 μg/ml of chloramphenicol and/or erythromycin. HG003 Δ*rnaIII*:tag004 (SaPhB618) and HG003 Δ*sprY*:tag112 (SaPhB980) were constructed using pMADΔ*rnaIII*:tag004 and pIMsprY2::tag112, respectively, as described (23).

**Plasmids constructions**

Supplementary Table S2 lists all the primers used. To construct the sRNA-expressing vectors, we used pICS3 (24). To construct the pICS3-sprY, sprY with its endogenous promoter was amplified by PCR with primers 13–14 and cloned in pICS3 vector digested by PstI and NarI. To introduce mutations in sprY, primers 15–16 were used.

To construct the pCN33-*PtuA*-rot-gfp vector, which expresses rot under control of the *PtuA* promoter, we amplified 373 nucleotides rot with primers set 18–19. HG003 strains carrying each of the target-gfp fusions and the sRNA plasmids were grown on BHI agar plates supplemented with 10 μg/ml chloramphenicol and erythromycin. The fluorescence measurements of the co-transduced HG003 strains were performed as previously described (24).

PCR products corresponding to the sequence of ms2 tag, or ms2 fused with sprY were cloned into pRMC2 digested by KpnI / SacI (see Supplementary Table S2 for primers containing MS2 tag sequence). All cloning experiments were performed with Gibson Assembly Master Mix (New England Biolabs). The reactions were then transformed into *E. coli* DH5-α by heat shock at 42°C. The plasmids were purified from overnight cultures in LB broth supplemented with Ampicillin 100 μg/ml, extracted (Miniprep Extraction Kit, Qiagen) and Sanger sequenced by using BigDye Terminator v3.1 Cycle sequencing Kit, using a 5130 × 1 capillary electrophoresis genetic analyzer (Applied Biosystems). The purified plasmids were used for the transformation in *S. aureus* RN4220 strain by electroporation shock. The φ80 phages prepared from RN4220 were then used to transduce plasmids in HG003 strains. pMADΔ*rnaIII*:tag004 is a pMAD derivative containing the PCR-amplified rnaIII upstream, tag004 and rnaIII downstream sequences cloned by Gibson assembly (using primers RNAIII UpF, RNAIII UpR, RNAIII DwF, RNAIII DwR for rnaIII adjacent sequences) as described. pIMsprY2::tag112 is a pIMAY derivative containing the PCR-amplified sprY upstream, tag004 and sprY downstream sequences cloned by Gibson assembly (using primers pIMAY-Up-SprY2-F, pIMAY-Up-SprY2-R, pIMAY-Down-SprY2-F and pIMAY-Down-SprY2-R for sprY adjacent sequences) as described (23).

**Proteins extraction and Western blots**

*Staphylococcus aureus* strains were grown until exponential phase (OD<sub>600</sub> = 0.8) or stationary phase (OD<sub>600</sub> = 10.1) in BHI at 37°C, with agitation at 160 rpm, and the cells were then pelleted for 10 min at 4°C (8000 × g). The total proteins extractions were prepared according to (25). Rot expression was visualized by anti-Rot antibodies (Benson, 2012, JB) and anti-rabbit IgG secondary antibodies (Jackson). Western blots were revealed using the Amersham ECL Plus detection Kit. Signals were visualized using LAS 4000 (GE Healthcare).

**In vitro transcription, RNA labeling and Gel-shift assays**

All RNAs were transcribed from PCR-generated DNA using MEGAscript T7 kit (Ambion). The template for transcription was amplified using HG003 genomic DNA and forward primers containing T7 promoter sequences (Supplementary Table S1). RNAs were labeled at 5'-end using [γ-<sup>32</sup>P] ATP (Amersham Biosciences) and T4 polynucleotide kinase (Invitrogen). Labeled and unlabeled RNAs were purified on a 5% acrylamide urea gel, eluted in Elution buffer (20 mM Tris-HCl pH 7.5, 250 mM NaCl, 1 mM EDTA, 1% SDS) at 37°C, eluted, ethanol precipitated, quantified by Qubit (Thermo Fisher Scientific) and stored at -80°C.

Gel-shift assays were performed as described in (26). RNAs were denatured in 50 mM Tris/HEPES pH 7–7.5, 50 mM NaCl for 2 min at 80°C, followed by refolding for 10 min at 25°C after adding MgCl₂ at final concentration of 5 mM. The binding reactions were performed in 50 mM Tris-HCl (pH 7.5), 50 mM NaCl, 5 mM MgCl₂ for 20 min at 25°C. About 0.025 pmoles of labeled SprY or SprY mutants were incubated with various concentrations of RNAIII. The samples were supplemented with 10% glycerol and were loaded on a native 4% polyacrylamide gel containing 5% glycerol. The gels were dried and visualized by Typhoon FLA 9500 scanner (GE Healthcare).

**RNA extractions, northern blots, RNA half-life determination and qPCR assay**

The cells were collected at exponential and at stationary phases of growth, pelleted for 10 min at 4°C (4500 × g) and resuspended in RNA lysis buffer (0.5% SDS, 20 mM acetate of sodium, 1 mM EDTA, pH 5.5). Total RNA

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was extracted as previously described (7). The Northern blot assays were carried out as previously described (9). The membranes were hybridized with specific 32P-labeled probes (Supplementary Table S2) in ExpressHyb solution (Clontech) and were washed according manufacturer recommendations. The membranes were then exposed and scanned with Typhoon FLA 9500 scanner (GE Healthcare). The images quantifications were realized with ImageQuant Tool 7.0.

For sRNA half-life determination, rifampicin was used as the most common treatment to stop the transcription (27). Staphylococcus aureus HG003 strain and its derivatives were cultured overnight, diluted to 1/100, grown for 5 h at 37°C, and incubated with 20 mg/ml rifampicin. About 8 ml of each strain was collected before and at 2, 5, 10, 20, 30, 40, 60 and 90 min after adding rifampicin. These samples were centrifuged, the pellets were frozen in liquid nitrogen and then stored at -80°C. Total RNA was extracted. For the quantitative real-time PCR (qRT-PCR), total RNA extraction samples were treated with DNaseI Amplification grade Kit (Invitrogen). cDNAs preparations and qRT-PCR experiments were performed as previously described (24). The gyrB gene was used for normalization. As for absolute quantification by qPCR, instead of using gyrB, we prepared a PCR reaction with respective PCR products at concentrations of 10^6/8/6/4/2/0 copies/μl. Identification of TSS of sprY was performed as described in (28). For this reverse transcription (RT) was done on 5 μg of total RNA of HG003 strain with labeled primer 2.

**Preparation of the MS2-affinity column**

To prepare the 6His-MBP-MS2 protein, we used the pHMN plasmid (29) containing the 6His tag at the N-terminal and the MS2 tag at the C-terminal. The induction of protein production and bacterial lysis was performed as described by (30). The 6His tag allows a first purification on nickel resin, using an AKTA (GE healthcare). After being washed with water, the pumps and the column were equilibrated with lysis buffer (50 mM NaH2PO4, 300 mM NaCl, 0.5% Tween 20, 10 mM Imidazole 10% glycerol). The bacterial lysate was added into the column and various fractions were recovered using the elution buffer (lysis buffer + 250 mM of imidazole). The fractions of interest were passed through a desalting column to remove traces of the elution buffer. A second purification was carried out using an amylose column washed and then equilibrated with buffer 2 (20 mM Tris-HCl, 0.2 mM NaCl, 0.5 mM EDTA). The fractions of interest were passed through the column and then eluted using a second elution buffer (buffer 2 + 10 mM maltose buffer). The fractions were also passed through a desalting column to remove all traces of maltose. For the preparation of the MS2-affinity column, 100 μl of amylose resin (NEB # E8021S) was added in a Bio-spin disposable chromatography column (Biorad # 732-6008). The column was then washed three times with 1 ml of buffer A (20 mM Tris-HCl pH 8, 150 mM KCl, 1 mM MgCl2, 1 mM DTT, 1 mM PSMF). A solution of 100 pmoles of 6His-MBP-MS2 protein was added to the column and was incubated for 5 min, and washed twice with 1 ml of buffer A.

**Purification of bacterial lysates, RNA sequencing and Bioinformatic analysis**

Staphylococcus aureus strains were grown until exponential phase (OD600 = 0.8) or stationary phase (OD600 = 10.1) in 50 ml of BHI (Oxoid). Then expressions of ms2-sprY and ms2 were induced with 1 μM of anhydrotetracycline for 10 min. The bacteria were plated in ice for 10 min and centrifuged for 5 min at 4000 × g. The pellets were then washed with 1 ml of buffer A, centrifuged for 1 min at 16 000 × g and stored at -80°C. Frozen pellets were thawed on ice and resuspended in 2 ml of buffer A. Mechanical lysis with Fastprep (Fastprep, MP Biomedicals) in the presence of 250 μl of glass beads was carried out for 3 × 30 s at 6500 × g. The samples were centrifuged for 10 min at 4°C at 16 000 × g. The bacterial lysates obtained were passed through an affinity column. Columns were washed 5 times with 1 ml of buffer A and then eluted with 1 ml of buffer A + 15 mM maltose. In order to carry out checks, we extracted the RNAs at different points in the experiment. We obtained four RNA extracts corresponding to the total RNAs taken before passage through the amylose column (input), RNA passed through a column (Flow-Through), RNA recovered after the last washing of the column (W) (Supplementary data). The RNAs from the fractions eluted from the columns are extracted with chloroform phenol as previously described (10). For MS2 and MS2-SprY, the northern blots were carried out using 5 μg of RNA according to the protocol as previously described (9). The primers used are listed in Supplementary Table S2. A bio-analyzer (2100-Agilent bio-analyzer) was used to quantify and verify the purity of the samples before being sequenced (following the manufacturer’s instructions [Agilent]). RNA sequencing was carried out as previously described (31). The cDNA libraries were prepared with the NEBNext Ultra™ II Directional RNA Library Prep Kit for Illumina® and then sequenced as paired-end reads (2 × 75 bp) using an Illumina MiSeq platform and the MiSeq reagent kit version 3. The reads were mapped against the genomic sequence of S. aureus HG003 strain (Genbank accession no. CP000253) and then counted using the CLC Genomics Workbench software v8.1 (Qiagen). Statistical analysis was performed using the DESeq2 R package (32). Raw and processed data generated in this study have been submitted to the Gene Expression Omnibus (GEO) repository at the National Center for Biotechnology Information (NCBI) and are available under accession no. GSE166499.

**Hemolysis assays**

Staphylococcus aureus strains grown until exponential phase diluted to OD600 = 0.1 in BHI. The supernatants were taken at 2 h of bacterial growth, filtered with 0.45-μm filter and stored at -20°C. For blood sample preparation, 1 ml of human or mouse blood was centrifuged for 10 min at 4000 × g at 25°C, followed by multiple washes with PBS 1X (qsp 1 ml) to eliminate the plasma and lysed red blood cells and re-suspended in 10 ml of PBS 1×. The prepared blood sample and the supernatants (diluted in 1/10) were added at a ratio 50:50 in 96 pointy wells plates to a final volume of 150 μl per well. After an incubation of 1 h at 37°C, the plate was centrifuged for 10 min at 4000 × g at 25°C and 100 μl of the supernatants
were collected into a new 96 flat well plates and read at OD 540 nm. The mix of blood sample with PBS 1× or with 0.1% Triton were used respectively as negative and positive controls of hemolytic activity.

Animal infection model
HG003, HG003ΔsprY and complemented strains were used to study the virulence level in a murine intravenous sepsis model. All experimental protocols were approved by the Adaptive Therapeutics Animal Care and Use Committee (APAFIS#2123–2015100214568502v4). For the sepsis model, we used female Swiss mice (Janvier Labs), 6–8 weeks old and weighing ~30 g. Groups of five female mice were inoculated i.v. with 200 μl of bacterial suspensions containing 2 × 10⁸ S. aureus cells in 0.9% NaCl. The survival of the mice was monitored for 12 days, and the statistical significance of differences between groups was evaluated by comparing Kaplan–Meier survival curves with the Mantel-Cox test. A P-value of 0.05 was considered significant.

RESULTS
SprY, a novel sRNA expressed from S. aureus srna cluster
A small staphylococcal RNA expressed from prophage φ12 (33) in NCTC8325 derivatives was initially described as a putative 5′UTR of a small open reading frame SAOUHSC_A014155 of 60 amino acids, and called S629 (34). However, based on in-depth analysis of the HG003 strain genome, we designated S629 as a bona fide sRNA gene (7,9). Because the gene S629 is adjacent to the sprX2 sRNA gene (7, 9), we renamed this sRNA SprY (Figure 1A and Supplementary Figure S1.A). SprY is an acronym coined for small pathogenicity island sRNA (35). The SprY 5′-end determined by reverse transcription (RT) (Supplementary Figure S1.B) corresponds to nucleotide position 1464380 of the NCTC8325 genome sequence (Supplementary Figure S1.A). Detection of SprY by northern blot (Figure 1B) and in silico identification of a Rho-independent transcriptional terminator upstream of SAOUHSC_A014155 (using ARNold; (36)) indicate that SprY is a bona fide sRNA of about 125 nucleotides. It is found in several S. aureus isolates with 95% sequence identity but is not present in other Staphylococcaceae (5). SprY is present only in strains containing the φ12 prophage; accordingly, it was detected by northern blot in Newman and HG003 strains but not in USA300 and N315 (Figure 1B).

SprY amounts are highest in pre-stationary phase and slowly decrease during stationary phase (Figure 2A). SprY half-life is 23.3 ± 1.45 min in pre-stationary phase (Figure 2B), implying that SprY is stable RNA, compared to most mRNAs with half-life of 2–4 min (37,38). Since sprY and sprX2 are adjacent sRNA genes, we considered that their expression could be inter-dependent. For this purpose, we analyzed the amounts of both, SprY and SprX2 sRNAs in HG003 (39) and its derivatives deleted for either sprY (ΔsprY) or sprX2 (ΔsprX2). Despite their proximity, deletion of one sRNA gene did not affect the expression of the other (Figure 2C). In addition, overexpression of sprX2 (pICS3-sprX2) or sprY (pICS3-sprY) did not affect the expression level of the other sRNA (Figure 2C). We concluded that sprY and sprX2 are expressed independently from each other. Despite the close proximity of these genes, their expression patterns are different, since SprX2 accumulates during exponential phase while SprY accumulates in pre-stationary phase (Supplementary Figure S2. (9)).

The SprY structure predicted by RNAfold (40,41) and LocaRNA softwares (42–44) revealed three stem loops, the last one being a Rho-independent transcriptional terminator (Supplementary Figure S3). The activity of several staphylococcal sRNAs involves single-stranded C-rich regions (8,45). However, this feature is not present in SprY, suggesting that it may not target G-rich regions associated with Shine-Dalgarno sequences but other sequences.

SprY interacts directly with RNAIII, a major regulator of virulence factors of S. aureus
Most sRNAs act by base-pairing to RNA targets. To identify potential direct targets of SprY, we performed in vitro MS2-RNA affinity purification coupled with RNA sequencing (MAPS) (46). For this purpose, an MS2 tagged version of SprY was expressed under the control of an inducible promoter in HG003 (Supplementary Figure S4.A and B). After 2 and 6 h of growth, each followed by 5 min of induction, RNAs in complex with MS2-SprY were isolated by MS2 affinity chromatography, eluted, and identified by RNAseq. RNAIII, rpmG1 mRNA and SAOUHSC_J342a mRNA were enriched 7.24-, 11.11- and 7.59-fold, respectively, with MS2-SprY compared to MS2 alone (Supplementary Figure S4.C and Supplementary Table S3). This significant enrichment suggests that these three RNAs are SprY targets. However, no base-pairing with a significant energy between SprY and rpmG1 mRNA was predicted by in silico analysis suggesting that rpmG1 mRNA enrichment with MS2 tagged SprY could be due to an indirect interaction. In contrast, IntaRNA software (47,48) indicated potential base-pairings of SprY with the 5′ UTR of SAOUHSC_J342a (Supplementary Figure S4.D) and with RNAIII (Figure 3A), supporting the MAPS results. As RNAIII is the major virulence regulator and effector of quorum sensing in S. aureus, we focus our study on the interaction between SprY and RNAIII.

IntaRNA analysis indicates potential base-pairing between RNAIII and SprY involving the 1st hairpin of SprY (4th to 46th nucleotide) and the 13rd hairpin of RNAIII (Figure 3A). Expectedly, SprYMa lost the ability to bind RNAIII (Figure 3C), implying that the 1st SprY hairpin is required for pairing. To bring further evidence to support the predicted pairing, an RNAIII derivative with compensatory mutations (RNAIIIΔm) restoring the pairing with SprYMa (Supplementary Figure S5.C) was synthesized and tested by EMSA. A gel retardation was observed between SprYMa and RNAIIIΔm but not between SprYMa and RNAIII (Figure 3C). In addition, RNAIII mutated in the 13th loop, which corresponds to a part of the
Figure 1. *sprY* and *sprX2* genomic localization and expression in *Staphylococcus aureus* HG003 strain. (A) The genomic localization of *sprY* and *sprX2* genes in *S. aureus* HG003 strain. *sprY* gene is located around 50 bp upstream of *sprX2* gene (7,9) in the staphylococcal phage ϕ12. (B) Expression of SprY in different strains of *S. aureus*. RNA extraction was performed in HG003, USA300, Newman and N315 strains at exponential phase (E) and stationary phase (S). As loading controls, the blots were also probing for tmRNA.

Figure 2. SprY expression in HG003 strain. Expression of *sprY* was determined by northern blot analysis using labeled DNA probes for SprY. As loading controls, the blots were also probed for tmRNA. (A) SprY expression profile during a 24 h growth of *Staphylococcus aureus* HG003 strain (WT). The growth curve of WT strain is presented by line, with the quantification of SprY expression level relative to the amount of tmRNA from the same RNA extraction in black chart (a.u., arbitrary units). (B) Stability of SprY. HG003 (WT) was grown until 6 h (t = 0) and the RNA extraction was performed at 2, 5, 10, 20, 40, 60 and 90 min after adding Rifampicin. The quantification of SprY stability, in semi-log plot, was performed by ImageQuant Tools 7.0. The data represent the mean of three different experiments ± standard error. The maximum value of rnaIII expression at time 0 is normalized to 100% and the time at which the sRNA reaches 50% of its original level is indicated with red dotted line. (C) The expression of *sprX2* and *sprY* by northern blots, using labeled DNA probes for SprY and SprX2. As loading control, the blots were also probing for tmRNA.

predicted SprY binding region, noticeably lost the capacity to bind SprY (Supplementary Figure S5_A and B). Altogether, our in vivo (MAPS), in silico and in vitro (EMSA) experiments revealed an interaction between the 1st SprY hairpin and the 13th RNAIII loop.

In many cases, base-pairing with sRNAs affects RNA target stability. We therefore tested if binding of SprY would affect the stability of RNAIII. The stability of RNAIII was compared in HG003 harboring the plasmids pICS3-sprY, pICS3-sprYmA and the control vector pICS3 (Figure 3D). No significant difference was observed in RNAIII stability, whether SprY or SprYmA were overexpressed (Figure 3E). This result prompted us to further analyze the role of SprY toward RNAIII.

**SprY affects expression of RNAIII targets**

RNAIII is the effector of quorum sensing and a major regulator of a plethora of virulence factors. Interestingly, the 13th RNAIII stem-loop is involved in the repression of several among its numerous mRNA targets. By binding to the 5′ leader region of *rot* mRNA (for repressor of toxins), RNAIII, via the 13th stem-loop, negatively regulates the expression of this target at the translational level (49) and in consequence obstructs the expression of toxins (20,21). Another well-studied target of the 13th RNAIII stem-loop is *ecb* (for extracellular complement binding protein) (19,24), which is involved in blocking bacterial recognition by the host immune system (18).
Figure 3. SprY interacts with RNAIII. (A) Interaction between SprY and RNAIII was predicted by IntaRNA software (47). Mutant of SprY and compensatory mutant RNAIII were constructed (SprYmA and RNAIIImA). The nucleotides underlined and bolded correspond to the mutations in the sprY and rnaIII sequences. (B) Schematic presentation of RNAIII secondary structure and the region of RNAIII that interacts with SprY (shown by line). (C) Complex formations between SprY, SprYmA with RNAIII and RNAIIImA were analysed by native gel retardation assays. Shift assays of purified labeled SprY and SprYmA (SprY* and SprYmA*) were performed with increasing concentrations of RNAIII and RNAIIImA (0.25, 0.5 and 1.25 μM). (D) Verification of sprY expression in HG003 strain containing empty plasmid (pICS3) or over-expressing sprY (pICS3-sprY) or muted sprY (pCIS3-sprYmA) by northern blot using labeled DNA probes for SprY and tmRNA as control. (E) RNAIII stability was studied by northern blot using RNA total extraction in HG003 pICS3, HG003 pCIS3-sprY and HG003 pCIS3-sprYmA, at exponential growth phase (OD = 0.8). Time (min) corresponds to the time after adding Rifampicin. The RNAIII expression levels of in each sample were is normalized with the control gene (tmRNA). The data represent the mean of three different experiments ± standard error for RNAIII stability. The maximum value of rnaIII expression at time 0 is normalized to 100% and the time at which the sRNA reaches 50% of its original level is indicated with dotted line. All statistical analyses were performed using two-way ANOVA test and Bonferroni comparison test.

We therefore questioned if the SprY binding to the 13th RNAIII stem-loop could affect the activity of RNAIII against specific targets. Surprisingly, sprY overexpression showed no significant impact on rot mRNA levels (Figure 4A). We hypothesized that SprY could affect rot translational initiation rather than rot mRNA quantity. We constructed a rot-gfp translational gene fusion under the control of the constitutive P_{tufA} promoter in pCN33 resulting in pCN33-P_{tufA}-rot-gfp. HG003 and HG003ΔrnanIII containing empty pICS3 or pICS3-sprY were transformed with pCN33-P_{tufA}-rot-gfp. In addition, a strain overexpressing rnaIII under constitutive promoter amiA (pICS3-P_{amiA}-rnaIII) was used as a control. As expected, SprY overproduction significantly increased rot translation only in the presence of RNAIII (Figure 4B). Furthermore, the overexpression of SprYmA had no effect over the fluorescence of Rot-GFP fusion (Figure 4B), which implies that the regulation of rot expression by SprY involved the interaction between SprY 1st hairpin and RNAIII. In addition, deletion of sprY resulted in a small but significant decrease in fluorescence intensity of the Rot-GFP fusion, while strains overexpressing sprY showed increased fluorescence (Supplementary Figure S6). We also confirmed the effect of SprY on endogenous rot expression in an rnaIII dependent manner by Western blot (Figure 4C). Overexpression of SprY increases Rot level in the HG003 wild-type strain but not in the absence of RNAIII.

The effect of SprY was also tested on the expression of ecb, another target regulated by the 13th RNAIII stem-loop. Overexpression of SprY induced an increase in ecb mRNA levels (Figure 4D). As in the case for rot, SprY overproduction significantly increased ecb translation only in the presence of RNAIII (Figure 4E). Moreover, like for rot regulation, the overexpression of sprYmA had no effect on the fluorescence of Ecb-GFP (Figure 4E), which implies that the regulation of ecb expression by SprY requires the interaction between SprY and RNAIII. In addition, as in the case for rot, the absence of sprY resulted in decreased fluorescence of the Ecb-GFP fusion (Supplementary Figure S6). Bacterial growth of all strains used for this experiment was essentially equivalent (Supplementary Figure S7). As with Rot, we observed a significant decrease in fluorescence intensity of Ecb-GFP in the absence of SprY, and an increase of fluorescence of strains overexpressing sprY (Figure 4). Taken together, our results indicate that SprY affects the expression of RNAIII targets likely by titrating the regulation mediated by its 13th loop.
Figure 4. SprY affects the expression of RNAIII targets. Analysis of \( \text{rot} \) (A) and \( \text{ecb} \) (D) transcript levels in HG003 strain containing empty plasmid (pICS3) or overexpressing \( \text{sprY} \) (pICS3-sprY) by qPCR, using RNA extraction of those strains at early exponential growth phase. The mRNA expression level of \( \text{rot} \) and \( \text{ecb} \) in qPCR is normalized with the control gene (gyrB) and is calculated with \( 2^{-\Delta\Delta CT} \) relative quantification. (C) Analysis of Rot protein level in HG003 strains (WT and \( \Delta \text{rnaIII} \)) containing empty plasmid (pICS3) or over-expressing \( \text{sprY} \) (pICS3-sprY) by western blot using polyclonal antibodies against Rot. The translational initiation level of \( \text{rot} \) (B) and \( \text{ecb} \) (E) under SprY regulation were studied by using gfp gene reporter. *Staphylococcus aureus* strains (WT and \( \Delta \text{rnaIII} \)) containing the pCN33-\( \text{PtufA-rot-gfp} \) or pCN33-\( \text{PtufA-ecb-gfp} \) fusion plasmids co-transformed with pICS3, pICS3-sprY, pICS3-sprYmA or pICS3-PamiA-rnaIII. The fluorescent intensity was measured every 10 min over 20 h in a Biotek microplate reader. BacteriagrowthindifferentHG003strainsandGFPfluorescenceintensityareshownrespectivelyintheleftpanelatOD600andinrightpanelquantifiedbyBiotek. Allstatistical analyses were performed using Student’s \( t \) test. The error bars correspond to the average values from three independent experiments. Statistical significance is indicated by bars and asterisks as follows: **, \( P < 0.01 \); ***, \( P < 0.005 \); ****, \( P < 0.001 \).

**Abundance of RNAIII and SprY depends on growth conditions**

As the regulatory activity of RNAIII controlled by SprY is based on RNAIII/SprY direct binding, the endogenous ratio of each partner likely controls the biological functions of this interaction. In rich medium, both SprY and RNAIII accumulate during growth and diminish in late stationary phase (Figure 5A). The expression profiles in different media were similar for both sRNAs (Figure 5B), even when bacterial growth rates were different (Supplementary Figure S8_A). Quantities of SprY and RNAIII were determined in different media in exponential (2 h) and stationary (6 h) phases by RT-qPCR (Supplementary Figure S8_B). Compared to RNAIII, the SprY copy number was 2-fold higher in BHI and RPMI, and 4-fold higher in TSB at 2 h of growth; however, it was significantly lower than RNAIII in NZM at 2 h (0.1-fold) (Figure 5C). The RNAIII copy number is far greater than that of SprY in all media.
Growth. The chart represents the ratio of SprY HG003 wild-type strain in BHI, TSB, NZM and RPMI at 2 and 6 h of growth by qPCR absolute. The RNA samples were prepared from the pellet of northern blot. The accumulation of SprY led to a significant decrease in RNAIII over, Rot is known to repress toxins production (20). Since Rot is also the binding site for targets such as ecb, SA2353, and RNAIII sequence pairing with SprY is the mimicry of targeted RNA by other RNAs are essential mechanisms to adjust the action of regulatory sRNAs reviewed in (52). RNAs with sponge-like activity were recently described in different bacterial species such as E. coli, Bacillus subtilis and Pseudomonas aeruginosa (53–55) and reviewed in (52). These sponge-like RNAs use different mechanisms to control sRNA regulators. Their binding can induce the cleavage and/or destruction of sRNA, as the S. aureus hemolytic activity in all strains except for HG003 ΔRNAIII, indicating that RNAIII is required for SprY-mediated hemolysis.

Since SprY restricted hemolysis in mouse blood, we considered mice as a suitable model to test the role of sprY during staphylococcal infection. A murine intra-venous sepsis model was used with an inoculum of 2 × 10⁸ bacteria per mouse. The survival rate of mice infected with either HG003 or HG003 ΔsprY was not statistically different (Figure 6C). We also tested the HG003 ΔsprY containing either the pICS3 (control plasmid) or overexpressing sprY (pICS3-sprY). The virulence of HG003 ΔsprY overexpressing sprY was drastically reduced compared to control strain (pICS3) (Figure 6D). Taken together, our results show that SprY reduces S. aureus virulence likely by decreasing expression of virulence factors including RNAIII-activated hemolysin.

**DISCUSSION**

We report the function of SprY alias S629 (34), a recently identified bona fide sRNA (5), expressed from HG003 prophage φ12 (also known as bacteriophage Sa2 or φsa2) (33). Here, we identified SprY as an sRNA tuning the sophisticated RNAIII regulation network. Co-purification of in vivo RNAs associated to SprY revealed that SprY binds RNAIII. RNAIII is the key effector of the accessory gene regulator (agr) system and the major ribonucleic staphylococcal regulator of virulence, which controls translation and stability of several mRNA targets by antisense base-pairing involving its various stem-loop structures (17,19) (Figure 7A and Supplementary Figure S9). SprY binding does not affect RNAIII stability but alters its function. We propose that SprY sterically prevents the formation of complexes between RNAIII and its targets, therefore affecting their expression. Indeed, SprY overproduction increases rot mRNA translation only when the interaction between SprY and RNAIII takes place. Moreover, SprY stimulates expression of the staphylococcal extracellular complement binding protein, Ecb (50) in an RNAIII-dependent manner. In accordance with positive regulation of rot, whose regulon includes genes encoding proteins with hemolytic activity such as hla, psma and hlgACB (22,51), SprY reduces the hemolytic activity of S. aureus on human and mouse blood dependently from RNAIII presence.

The interactions we uncovered between SprY and RNAIII prompt us to hypothesize that SprY acts as an RNA sponge for RNAIII to prevent RNAIII-dependent regulations. The competition between mRNA targets and the mimicry of targeted RNA by other RNAs are essential mechanisms to adjust the action of regulatory sRNAs (reviewed in (52)). RNAs with sponge-like activity were recently described in different bacterial species such as E. coli, Bacillus subtilis and Pseudomonas aeruginosa (53–55) and reviewed in (52,56). These sponge-like RNAs use different mechanisms to control sRNA regulators. Their binding can induce the cleavage and/or destruction of sRNA, as
Figure 6. Impact of SprY to hemolytic activity in *Staphylococcus aureus* in different models of infection. (A and B) The supernatants of different strains of HG003 (WT, ΔsprY, ΔrnaIII) containing empty plasmid (pICS3) or overexpressing sprY (pICS3-sprY) were collected after 6 h of growth in BHI at 37°C. Hemolysis was performed by incubating cell culture supernatants with mouse and human blood samples at a ratio 50:50 at 37°C for 1 h. The hemolytic activity was observed in a flat-bottom 96-well microtiter plate at OD 540 nm. The blood samples prepared in PBS1× represent as negative control and in Triton 0.1% as 100% hemolysis. (C and D) Survival of mice infected with *S. aureus* wild-type strain (WT), deleted strain for sprY (ΔsprY) and complemented strains for sprY (ΔsprY pICS3-sprY). Groups of 5 eight-week-old Swiss female mice were inoculated i.v. with 2 × 10⁸ bacteria and monitored daily for 3 weeks. All statistical analyses were performed using Student’s *t* test. The error bars correspond to the average values from three independent experiments. Statistical significance is indicated by bars and asterisks as follows: **, *P* < 0.01; ****, *P* < 0.001.

...reported for *chb*-ChiX (57). In addition, since degradation is not an obligate outcome of duplex formation, a sponge RNA can act by titrating a regulatory sRNA and by competing with their true targets, as occurs between the tRNA spacer sequence with RybB or RyhB sRNAs (31). Until now, only few cases of base pairings between two sRNAs were identified in *S. aureus*; however, no ‘sponge-like’ function was shown for these interactions (58,59). Thereby, the findings presented here identify a sRNA with an original sponge activity in pathogenic bacteria.

To detectably adjust RNAIII function, SprY has to accumulate in amounts likely comparable to those of RNAIII. SprY and RNAIII expression varies according to growth conditions. The differences in sRNA stoichiometry could push the balance to different outcomes according to the growth environments. In rich media, SprY abundance is comparable to that of RNAIII during growth at low cell density. However, in late exponential growth phase, the ratio between SprY and RNAIII reverses and amounts of RNAIII vastly dominate those of SprY. This result suggests that SprY actively interferes with RNAIII function at low cell density resulting in ablating RNAIII function and consequently allowing the production of surface proteins and inhibiting the production of extracellular toxins (Figure 7B). Early growth phase presumably represents the early stages of infection (reviewed in 60); therefore, the action of SprY is likely to be important during the beginning of infections, allowing *S. aureus* to colonize various niches without causing severe consequences for the host.

Recently, another sRNA, SprX, was also shown to modulate the pathogenicity of *S. aureus* through binding RNAIII. In contrast to SprY, SprX binds the RNAIII 5′-end near the hemolysin *hld* coding sequence (58). Although the
Figure 7. Schematic view of RNAIII and SprY actions. (A) Schematic view of the secondary structure of RNAIII (12) and base-pairing interaction regions with SprY and other mRNA targets. hla mRNA interacts with RNAIII at 2nd and 3rd hairpins (rRNAIII sequence is colored in red). RNAIII encodes the delta-hemolysin (hld, in green). The interaction zone of RNAIII and SprY is indicated in red line and is overlapped with other RNAIII mRNA target listed here. rot mRNA interacts with RNAIII 7th hairpin (in blue line) and RNAIII 13rd stem-loop. mgrA mRNA interacts at two different sites of RNAIII (in purple line). (B) Model for SprY sponge function in RNAIII/mRNA targets regulation. At low cell density (upper panel), SprY presents in a higher number of copies than RNAIII and binds to all RNAIII molecules produced. This interaction prevents RNAIII from binding to rot and ecb mRNA and favors the translation of Rot and Ecb proteins. At high cell density (bottom panel), RNAIII is produced in higher level than SprY and suppresses the sponge effect of SprY, therefore inhibits the translation of rot and ecb, among other targets.
evolutionary questions regarding regulatory control provided by the prophase.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS
We are grateful to Dr Alexandra Gruss for the critical reading of the manuscript. We also acknowledge the ‘plate-form Génomique Santé’ Biogenouest Génomique Biosit core facility for their technical assistance. We thank the animal core facility ARCHE (ARCHE/SFR BIOSIT, Université Rennes 1, France).

FUNDING
Agence nationale de la recherche (ANR) [ANR-15-CE12-0003-01]: Fondation pour la Recherche Médicale (FRM) [DBF20160635724]; (INSERM) Institut National de la Santé et de la Recherche Médicale, the Région Bretagne (PhD thesis) (to K.B.L.H.); School of Pharmacy and Medical Sciences of the University of Rennes 1. Funding for open access charge: FRM [DBF20160635724].

Conflict of interest statement. None declared.

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