






# A multifaceted small RNA modulates gene expression upon glucose limitation in *Staphylococcus aureus*

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## Abstract

Pathogenic bacteria must rapidly adapt to ever-changing environmental signals resulting in metabolism remodeling. The carbon catabolite repression, mediated by the catabolite control protein A (CcpA), is used to express genes involved in utilization and metabolism of the preferred carbon source. Here, we have identified RsaI as a CcpA-repressed small non-coding RNA that is inhibited by high glucose concentrations. When glucose is consumed, RsaI represses translation initiation of mRNAs encoding a permease of glucose uptake and the FN3K enzyme that protects proteins against damage caused by high glucose concentrations. RsaI also binds to the 3' untranslated region of *icaR* mRNA encoding the transcriptional repressor of exopolysaccharide production and to sRNAs induced by the uptake of glucose-6 phosphate or nitric oxide. Furthermore, RsaI expression is accompanied by a decreased transcription of genes involved in carbon catabolism pathway and an activation of genes involved in energy production, fermentation, and nitric oxide detoxification. This multifaceted RNA can be considered as a metabolic signature when glucose becomes scarce and growth is arrested.

**Keywords** carbon metabolism; catabolite control protein A; pathogenic bacteria; regulatory RNAs; sRNA; translational regulation

**Subject Categories** Metabolism; Microbiology, Virology & Host Pathogen Interaction; RNA Biology

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## Introduction

All bacteria require a carbon source, providing energy for their growth and for the synthesis of all macromolecules. Besides, pathogenic bacteria during the host infectious process must cope with hostile conditions such as nutrient deficiency, temperature, and oxidative and osmotic shocks, and must overcome innate immune responses. For instance, *Staphylococcus aureus* uses carbohydrates to grow under high nitric oxide (NO) and anaerobiosis (Vitko *et al.*, 2016). To survive in these complex environments and to counteract the host defense, *S. aureus* produces a plethora of virulence factors. The synthesis of these factors is fine-tuned by intricate interactions between multiple regulators involving both proteins and RNAs (Novick, 2003). Additionally, biosynthetic intermediates, generated from the central metabolism of *S. aureus*, have strong impacts on the synthesis of virulence factors. Besides, several metabolite-sensing regulatory proteins (CcpA, CodY, Rex, and RpiR) act as key regulatory factors to coordinate the synthesis of genes involved in metabolic pathways, in stress responses, and in pathogenesis (Somerville & Proctor, 2009; Richardson *et al.*, 2015). Through the adaptation of the metabolism of the bacteria to specific host microenvironment, these proteins contribute to *S. aureus* pathogenesis (Richardson *et al.*, 2015).

Among these proteins, the catabolite control protein A (CcpA) acts as the main global regulator of carbon catabolite repression, allowing the bacteria to use the preferred carbon source (i.e., glucose) in a hierarchical manner (Seidl *et al.*, 2008a, 2009). Catabolite control protein A belongs to the LacI repressor family and binds to a specific DNA sequence, called the *cre* (catabolite-responsive element) sequence, which is conserved in many Gram-positive bacteria. Transcription of CcpA is constitutive, and the protein is activated through the binding of its co-regulator histidine-containing

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phosphocarrier protein HPr in the presence of glucose. Inactivation of *ccpA* gene decreases transiently the growth rate and yield as long as glucose is present in the medium (Seidl *et al*, 2006) and affects the expression of a large number of metabolic encoding genes in a glucose-dependent and glucose-independent manner (Seidl *et al*, 2008a). Additionally, CcpA has a strong impact on the expression of *S. aureus* virulon. It enhances the yield of the quorum-sensing-induced RNAlII, which represses protein A and various adhesion factors at the post-transcriptional level, and conversely activates the synthesis of many exotoxins. However, CcpA also modulates the transcription of mRNAs encoding protein A,  $\alpha$ -hemolysin (*hla*), and toxic shock syndrome toxin (TSST; Seidl *et al*, 2006, 2008b), represses capsule formation, and activates biofilm formation in a glucose-rich environment (Seidl *et al*, 2008a). Indeed, a *S. aureus* *ccpA* deletion mutant strain was less pathogenic than the wild-type strain in a murine abscess model (Li *et al*, 2010) and *ccpA* inactivation increased the susceptibility of hyperglycemic animals to acute pneumonia infections (Bischoff *et al*, 2017). Nevertheless, the mechanism by which CcpA affected *S. aureus* pathogenesis cannot be simply resumed as a modulation of the RNAlII-dependent regulatory networks. Therefore, it has been suggested that CcpA can also act indirectly on gene expression through the action of other regulatory proteins or small non-coding RNAs (sRNAs; Somerville & Proctor, 2009; Richardson *et al*, 2015).

In Enterobacteriaceae, several sRNAs have been shown as key actors of the uptake and the metabolism of carbohydrates (reviewed in Bobrovskyy & Vanderpool, 2016). For instance, they participate in the regulation of the galactose operon and carbon catabolite repression, metabolism of amino acids, and contribute to bacterial survival during phospho-sugar stress. The importance of sRNAs in regulatory networks is now well recognized to rapidly adjust cell growth to various stresses and changes in the environment. Thus, they are obvious candidates creating the links between virulence and metabolism. One example is RsaE, a sRNA conserved in *S. aureus* and *Bacillus subtilis*, that controls enzymes involved in the TCA cycle (Geissmann *et al*, 2009; Bohn *et al*, 2010) and in arginine degradation pathway (Rochat *et al*, 2018). In *B. subtilis*, the transcriptional repressor ResD represses RoxS (homologous to *S. aureus* RsaE) to readjust the pool of  $\text{NAD}^+/\text{NADH}$  in responses to various stress and stimuli (Durand *et al*, 2017). Its promoter is also highly conserved among *Staphylococceae* and is recognized by the orthologous response regulator SrrA in *S. aureus*. Responding to reactive oxygen species through SrrAB, *S. aureus* RsaE may also intervene in the survival of cells against host immune reactions (Durand *et al*, 2015, 2017).

Here, we have identified a signaling pathway responding to glucose uptake, which involves a sRNA, called RsaI (or RsaOG). This 144 nucleotides-long sRNA is highly conserved among *Staphylococceae* and carries two conserved regions including two G-track sequences and a long unpaired interhelical region rich in pyrimidines (Geissmann *et al*, 2009; Marchais *et al*, 2010). The expression of RsaI is observed exclusively at the stationary phase of growth in rich medium (Geissmann *et al*, 2009) and is enhanced after vancomycin exposure (Howden *et al*, 2013). In this study, we revealed that CcpA is the main repressor of RsaI expression in the presence of glucose and that this inhibition is alleviated after the utilization of glucose. The identification of the targetome of RsaI using the MS2-affinity purification approach coupled with RNA sequencing

(MAPS) unexpectedly showed two classes of RNA targets, including mRNAs involved in glucose uptake, sugar metabolism, and biofilm formation, as well as various sRNAs. Using site-directed mutagenesis, we identified two distinct and functional regions of RsaI. All in all, our data showed the existence of two sRNAs involved in essential pathways responding to either glucose or glucose-6 phosphate uptake, with RsaI acting as a signature for a metabolic switch when the preferred carbon source is metabolized.

We will discuss the importance of sRNA-mediated regulation in *S. aureus* to fine tune the expression of genes according to essential nutrient availability and their possible consequences on metabolism adaptation and virulence.

## Results

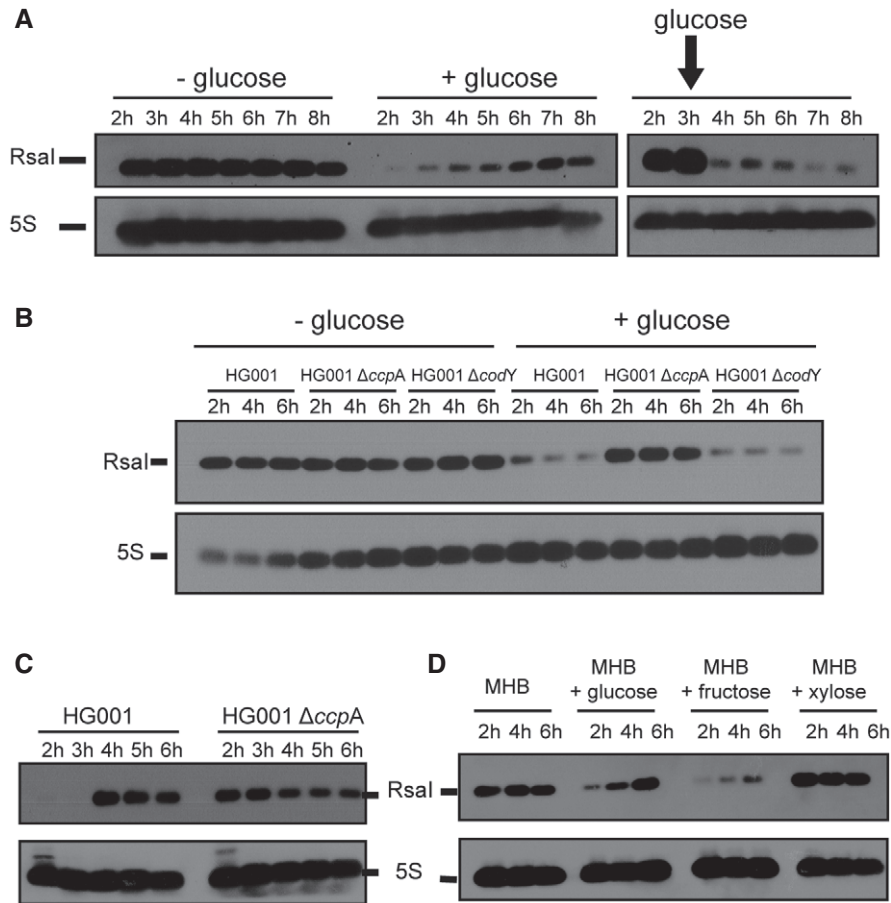
### The expression of RsaI is inhibited by glucose and by the catabolite control protein A

We have previously shown that the synthesis of RsaI is high at the stationary phase of growth in BHI medium while its expression was constitutive in MHB medium (Geissmann *et al*, 2009). These data suggested that RsaI expression is regulated in a manner dependent on nutrient or biosynthetic intermediate availability. A major difference between BHI and MHB composition is their carbon source, glucose, and starch, respectively. We wondered whether the expression of RsaI might be dependent on the available carbon source. For this purpose, Northern blot experiments were performed on total RNAs prepared from HG001 (wild-type) strain grown in MHB medium at various time points (Fig 1A). In MHB, where the glucose is not immediately available as the carbon source, RsaI was constitutively and highly expressed (Fig 1A). Conversely, when glucose was added, either at the beginning of the culture or after 3 h of growth, the steady-state level of RsaI was immediately dropped, indicating the repressing effect of this sugar (Fig 1A).

As CcpA sensed the intracellular concentration of glucose (Seidl *et al*, 2006), we analyzed whether the expression of RsaI was CcpA-dependent. Northern blot analysis was performed on total RNA extracts prepared from HG001 strain and an isogenic mutant deleted of *ccpA* gene ( $\Delta\text{ccpA}$ ). In parallel, we have also tested CodY, another global regulator of metabolism and virulence, which was shown as a direct regulator of amino acid biosynthesis and transport of macromolecules (Pohl *et al*, 2009; Majerczyk *et al*, 2010). The data showed that in the absence of glucose, the yield of RsaI was similar in all strains. However, in MHB medium supplemented with glucose, the yield of RsaI declined dramatically in the WT and  $\Delta\text{codY}$  strains, whereas it was still high in  $\Delta\text{ccpA}$  strain (Fig 1B). In BHI medium, RsaI was expressed after 4 h of growth in the WT strain, while its expression became constitutive in the mutant  $\Delta\text{ccpA}$  strain (Figs 1C and EV1A).

We also analyzed the expression of RsaI in MHB medium supplemented with various sugars such as fructose and xylose (Fig 1D). The data showed that expression of RsaI was very low at the beginning of growth in MHB supplemented with either glucose or fructose but not with xylose. These data show that CcpA senses both glucose and fructose (Fig 1).

Overall, we showed that CcpA represses RsaI expression in the presence of glucose. In accordance with this observation, a



**Figure 1. RsaI responds to glucose through the transcriptional factor CcpA.**

- A Northern blot experiments show the expression of RsaI during growth phase in the HG001 strain in MHB medium with or without the addition of 1.5 g/l of *D*-glucose. Glucose was added either at the beginning of growth (+ glucose, left panel) or after 3 h of growth (right panel).
- B Northern blot experiment shows the expression of RsaI during growth phase in the HG001,  $\Delta$ ccpA mutant strain, and  $\Delta$ codY mutant strain, in MHB medium with (+) or without (–) the addition of 1.5 g/l of *D*-glucose.
- C Northern blot experiment shows the expression of RsaI in the HG001 and  $\Delta$ ccpA mutant strains. Total RNA was prepared after 2, 3, 4, 5, and 6 h of culture in BHI medium at 37°C.
- D Northern blot analysis of RsaI in the HG001 strain grown in MHB medium with or without the addition of 1 g/l of glucose, fructose, or xylose. For all the experiments, loading controls were done using the expression of 5S rRNA (5S) as revealed after hybridization of the membranes with a specific probe. However for these controls, we used aliquots of the same RNA preparations, but the migration of the samples was performed in parallel to the experiments on a separate agarose gel because RsaI and 5S rRNA have very similar sizes.

Source data are available online for this figure.

conserved *cre* (GGAAAcGcTTACAT) sequence was found at position –30 upstream the transcriptional start site of RsaI (Fig EV1B). This region was sufficient to confer repression by glucose in the complemented strain containing pCN51::PrsaI (Fig EV1C).

### The targetome of RsaI as revealed by the MAPS approach

The MAPS approach (“MS2 affinity purification coupled with RNA sequencing”) was used to purify *in vivo* regulatory complexes involving RsaI. MAPS has been successful to identify the RNA targets of any sRNAs in *Escherichia coli* (Lalaouna et al, 2015) and more recently of RsaA sRNA in *S. aureus* (Tomasini et al, 2017). Briefly, the MS2-tagged version of RsaI was expressed from a plasmid under the control of the *agr*-dependent P3 promoter, allowing an accumulation of RsaI at the stationary phase of growth in the

$\Delta$ rsaI mutant strain. RsaI was detected by Northern blot using total RNAs extracted at 2, 4, and 6 h of growth in BHI medium. Using a DIG-labeled RsaI probe, we showed that the steady-state levels of MS2-RsaI were very similar to the wild-type (WT) RsaI and that MS2-RsaI was specifically and strongly retained by the MS2-MBP fusion protein after the affinity chromatography (Fig EV1D). The RNAs were then extracted from the eluted fraction to be sequenced. The data were analyzed using Galaxy (Afgan et al, 2016), and the sequencing reads were mapped, counted per feature, and normalized using the HG001 genome as reference (Herbert et al, 2010; Caldeleri et al, 2017). The enrichment of putative targets was measured by comparing the number of reads obtained from the MS2-RsaI purification and the MS2 alone as control. Since the MS2 tag alone was produced in the WT HG001 background, the untagged RsaI was also expressed under the conditions of growth but was

poorly retained on the affinity chromatography (Fig EV1D). The data were sorted by a decreasing fold change. In the following study, we have considered as RsaI targets, the RNAs that were enriched at least fourfold and were reproducibly detected in two independent experiments (Tables EV1 and EV2).

Two classes of RNAs were co-purified with RsaI, including mRNAs and sRNAs. On the one hand, the most enriched mRNA encodes IcaR, the repressor of the *icaADBC* operon, which encodes the enzymes required for the synthesis of the poly- $\beta$ -1,6-N-acetylglucosamine polymer (PIA-PNAG), the main staphylococcal exopolysaccharide constituting biofilms. In addition, several mRNAs encoding proteins linked to sugar utilization and transport, such as *glcU\_2* encoding a major transporter of glucose, and *fn3K* encoding fructosamine 3-kinase were significantly enriched. It is noteworthy that other less enriched mRNAs (< 4-fold) encode proteins related to sugar metabolism such as the trehalose-specific PTS transporter (TreB), a sugar phosphatase (YidA), and a maltose transport system permease (Table EV2). Finally, in addition to *icaR*, several mRNAs express transcriptional regulators (Xre type, the maltose regulatory protein GlvR, SlyA, SigS). On the other hand, we found the RsaD, RsaE, and RsaG sRNAs as putative RsaI targets (Table EV1), which all contained at least one conserved C-rich motif (Geissmann *et al*, 2009). RsaH could be also included in this group though it was less enriched (< 4-fold; Table EV2).

The MAPS data suggested that RsaI is involved in networks of RNA pairings.

### RsaI contains two distinct regulatory domains

Following the MAPS experiments, we searched for intermolecular base-pairing interactions between RsaI and its potential RNA targets using IntaRNA. Stable interactions were predicted for most of the enriched RNAs (Table EV1). The CU-rich unpaired region of RsaI was predicted to form base-pairings with most of the mRNAs. They were located close or at the ribosome-binding site of most of the mRNAs except for *icaR* and *isaA*, which involves nucleotides in their 3' untranslated regions (Table EV1). A second domain of interaction corresponds to the G-track sequences located in the first hairpin domain of RsaI and the C-rich sequences of the sRNAs RsaD, RsaE, RsaG, and RsaH (Fig 2A).

These results suggested that RsaI contains at least two functional domains, the unpaired CU-rich region dedicated to mRNA pairing and the G-track used to interact with other sRNAs. To validate whether RsaI effectively binds to the RNA candidates identified by MAPS through these motifs, we performed gel retardation assays

using WT and mutated RsaI molecules (Figs 2B and C, and EV2). For that, *in vitro* 5' end-labeled RsaI was first incubated with increasing concentrations of various mRNAs encoding proteins involved in biofilm formation (IcaR), sugar uptake and metabolism (*glcU\_2* and *fn3K*), and several sRNAs (RsaG, RsaE, and RsaD). For these experiments, we used the full-length mRNAs and sRNAs (Table EV3). Complex formation was performed with RNAs, which were renatured separately in a buffer containing magnesium and salt. As expected, the data showed that RsaI formed complexes with high affinity (between 20 and 100 nM, Table EV1) with many RNAs such as *icaR*, *glcU\_2*, and *fn3K* mRNAs, and RsaG sRNA (Fig 2B). The stability of other complexes (e.g., *treB* mRNA and RsaD sRNA) was significantly lowered (> 250 nM) (Fig EV2).

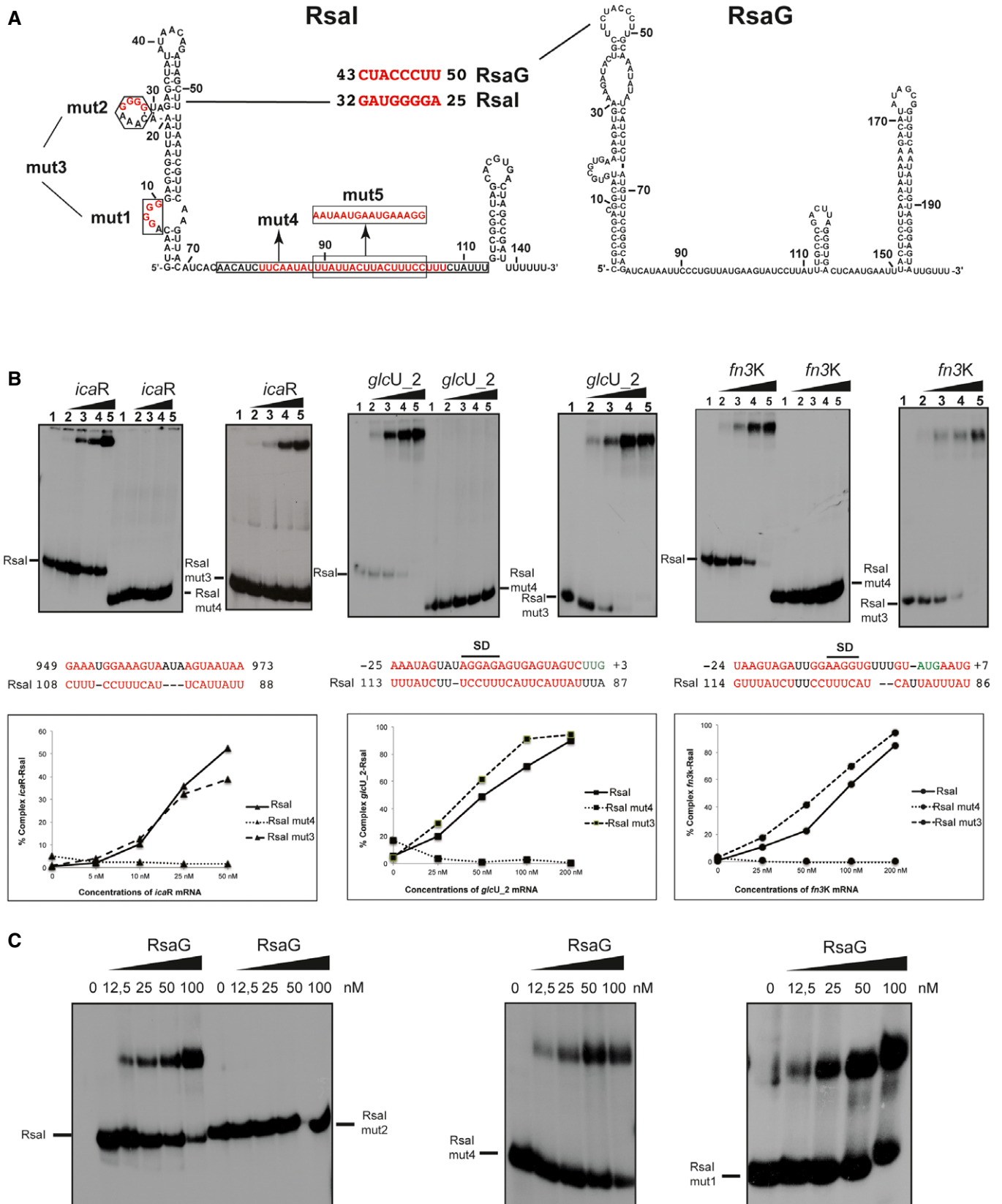
Based on the base-pairing predictions, mutations have been introduced into *rsaI* to map its functional regulatory regions. The two conserved G-track sequences were mutated separately (RsaI mut1:  $\Delta$ G7-G10, RsaI mut2:  $\Delta$ G26-G29) or together (RsaI mut3:  $\Delta$ G7-G10/ $\Delta$ G26-G29), while several nucleotides (RsaI mut4:  $\Delta$ U81-U107) were deleted in the conserved interhelical unpaired sequence (Fig 2A). We analyzed the ability of mutated RsaI derivatives to form complexes with *glcU\_2*, *fn3K*, and *icaR* mRNAs, and RsaG sRNA (Fig 2B). Quantification of the data showed that RsaI mut3 binds to all three mRNAs similarly to the WT RsaI, while complex formation was completely abolished with RsaI mut4. Only the mutation in the second G-track sequences (RsaI mut2) strongly altered the binding to RsaG, while the two other mutated RsaI (mut1 and mut4) recognized RsaG with an equivalent binding affinity as the WT RsaI (Fig 2C).

To demonstrate that the RNA-RNA interactions were also occurring *in vivo*, MAPS approach was used to monitor the effect of the RsaI mutations on its target RNAs. The RsaI mut2 and mut4, which independently affected *in vitro* binding of sRNAs and mRNAs, respectively, were tagged with MS2 and expressed in the HC001 $\Delta$ *rsaI* mutant strain. As described above, the enrichment of putative targets was calculated using a similar procedure (Table EV4). For this analysis, we have used the data coming from three independent experiments performed with the WT MS2-RsaI, the two MS2 controls and the MS2-RsaI mutants (mut2 and mut4) taking into account that the MAPS was done in two distinct sets of experiments (see Supplementary Material and Methods). This allowed us to directly compare the fold changes between the WT and the mutant RsaI versions (Table EV5). In the fraction containing MS2-RsaI mut4, most of the mRNA targets including *icaR*, *fn3K*, and *glcU\_2* were strongly reduced while RsaG was still significantly enriched. Conversely, we observed that the three mRNAs were still

**Figure 2. RsaI binds to *icaR*, *glcU\_2*, *fn3K* mRNA, and the sRNA RsaG.**

- A Secondary structure model of RsaI. In red, are the nucleotides deleted in the RsaI mutants (mut1 to mut4), and the nucleotides, which were substituted (mut5). The potential base-pairings between RsaI and RsaG are shown. Squared nucleotides are conserved sequences in RsaI.
- B Gel retardation assays show the formation of the complex between RsaI and *icaR*, *glcU\_2* and *fn3K* mRNAs. The 5' end-labeled wild-type RsaI (RsaI), RsaI mutant 3 (RsaI mut3, deletion of the two G-track motifs), and RsaI mutant 4 (RsaI mut4, deletion of the C/U-rich unpaired interhelical region) were incubated with increasing concentrations of mRNAs: lane 1, 0 nM; lane 2, 25 nM; lane 3, 50 nM; lane 4, 100 nM; and lane 5, 200 nM. Below the gels, the predicted interactions between RsaI and its targets are shown. Translation start codons are in green, and SD is for Shine and Dalgarno sequence. Graphs represented the % of complex formed between either RsaI or its two mutant forms (RsaI mut3 and RsaI mut4) and the target mRNA (*icaR*, *glcU\_2*, *fn3K*) as the function of mRNA concentrations.
- C Gel retardation assays show the formation of the complex between RsaI and RsaG. The 5' end-labeled wild-type RsaI (RsaI), RsaI mutant 2 (RsaI mut2), RsaI mutant 4 (RsaI mut4), and RsaI mutant 1 (RsaI mut1) were incubated with increasing concentrations of RsaG given in nM on the top of the autoradiographies.

Source data are available online for this figure.



co-purified together with the MS2-RsaI mut2 at a level close to the WT RsaI while RsaG was strongly reduced in the fraction containing MS2-RsaI mut2 (Table EV5). In addition, with the two MS2-RsaI mutants, RsaE was almost lost while the enrichment of the transcriptional regulator XRE was strongly reduced. Finally, *dck* mRNA remained enriched at the same level as for the WT MS2-RsaI (Table EV5). These data strongly suggested that mutations at the G-track motif of RsaI had a major effect on RsaG/RsaE sRNAs binding, whereas the deletion of the C/U-rich sequence in RsaI mut4 had a major impact on the recognition of many mRNAs. The loss of RsaE in the MAPS performed with MS2-RsaI mut4 suggested that the sRNA might also be associated with one of the RsaI-dependent mRNA target. In support to the hypothesis, base-pairings have been recently predicted between RsaE and the 5'UTR of *icaR* mRNA (Rochat et al, 2018).

Taken together, our data reveal that RsaI has at least two distinct regulatory domains that directly interact either with mRNAs or with sRNAs.

### RsaI inhibits ribosome binding by masking the Shine and Dalgarno sequence of mRNAs

Because the C/U unpaired region of RsaI was predicted to form base-pairings with the Shine and Dalgarno (SD) sequence of several mRNAs (Figs 2 and 3A), we analyzed whether RsaI would have the ability to compete with the ribosome for binding to *glcU\_2* and *fn3K* mRNAs. Using toe-printing assays, we analyzed the effect of RsaI on the formation of the ternary initiation complex constituting of mRNA, the initiator tRNA, and the 30S subunit. Binding of the 30S on the two mRNAs is illustrated by the presence of a toe-print signal at position +16 (+1 being the initiation codon). For *fn3K* mRNA, two toe-print signals were observed at +16 and at +20, most probably corresponding to the presence of two AUG codons distant of 5 nucleotides. However, only the first AUG is used *in vivo* (Gemayel et al, 2007). The addition of increasing concentrations of RsaI together with the 30S strongly decreased the toe-print signals for both mRNAs showing that RsaI is able to form a stable complex with the mRNA sufficient to prevent the binding of the 30S subunit (Fig 3A).

The *in vivo* relevance of RsaI-dependent repression of several mRNAs (*glcU\_2*, *fn3K*, *treB*, *HG001\_01242*, *HG001\_02520*) was then analyzed using gene reporter assays. The whole leader regions of the mRNAs (54 nts in *glcU\_2*, 34 nts in *fn3K*, 24 nts in *treB*, 35 nts

in *HG001\_01242*, and 72 nts in *HG001\_02520*) including 100 nucleotides of their coding sequences were cloned in frame with *lacZ* under the control of the strong promoter *PrpoB*. The synthesis of  $\beta$ -galactosidase was analyzed in the  $\Delta$ *rsaI* mutant strain and in the same strain transformed with the plasmid carrying the *lacZ* reporter and RsaI under its own promoter. For the *fn3K-lacZ* fusion, we also analyzed its expression in the mutant strain expressing RsaI mut4. The  $\beta$ -galactosidase activity was reduced almost eight times in cells expressing wild-type RsaI for all reporter constructs (Fig 3B). In addition, the expression of RsaI mut4 did not cause any repression of the  $\beta$ -galactosidase expressed from *fn3K-lacZ* (Fig 3B). Therefore, disrupting the interaction between RsaI mut4 and *fn3K* alleviated the repression of the reporter gene suggesting that the regulation of *fn3K* is occurring at the translational level through specific RNA-RNA interactions.

These data showed that RsaI hinders ribosome binding on *glcU\_2* and *fn3K* mRNAs in agreement with the fact that the C/U-rich region of RsaI would bind to the SD sequence of the mRNAs. We propose a similar mechanism for *treB*, *HG001\_01242*, and *HG001\_02520* mRNAs considering that the predicted base-pairings include the RBS of these mRNAs (Tables EV1 and EV2).

### RsaI interacts with the 3'UTR of *icaR* mRNA and affects PIA-PNAG synthesis

The MAPS approach revealed that *icaR* mRNA, which encodes the repressor of the main exopolysaccharidic compound of *S. aureus* biofilm matrix, is the most enriched mRNA co-purified with MS2-RsaI. This mRNA is of particular interest because it contains a large 3'UTR that is able to bind to its own SD sequence through the anti-SD UCCCCUG motif (Ruiz de los Mozos et al, 2013). Consequently, the long-range interaction provokes an inhibitory effect on translation and generates a cleavage site for RNase III (Ruiz de los Mozos et al, 2013). RsaI is predicted to form base-pairings with the 3'UTR of *icaR* downstream of the anti-SD sequence (Table EV1, Fig 4A). We first monitored whether the long-range interaction might be critical for RsaI binding. Previous work showed that the substitution of the anti-SD UCCCCUG sequence by AGGGGAC significantly destabilized the long-range interaction to enhance *icaR* translation (Ruiz de los Mozos et al, 2013). However, gel retardation assays showed that the WT and the mutant *icaR* mRNAs bind to RsaI with an equivalent binding affinity (Fig 4B), suggesting that the anti-SD sequence is not

#### Figure 3. RsaI inhibits *glcU\_2* and *fn3K* mRNA translation.

- A Toe-print assays showing the effect of RsaI on the formation of the ribosomal initiation complex of *glcU\_2* and *fn3K* mRNAs, respectively. Lane 1: incubation control of mRNA alone; lane 2: incubation control of mRNA with 30S subunits; lane 3: incubation control of mRNA with RsaI; lane 4: formation of the ribosomal initiation complex containing mRNA, 30S, and the initiator tRNA<sup>Met</sup> (tRNAi); lanes 5–9: formation of the initiation complex in the presence of increasing concentrations of RsaI, respectively: 50 nM (lane 5), 100 nM (lane 6), 150 nM (lane 7), 300 nM (lane 8), and 400 nM (lane 9). Lanes T, A, C, G: sequencing ladders. The Shine and Dalgarno (SD) sequence, the start site of translation (START), and the toe-printing signals (+16) are indicated. At the bottom of the gels are shown the predicted interactions between RsaI and its targets. Translation start codons are in green, and the Shine and Dalgarno (SD) sequence is underlined, and the arrowheads depict the toe-printing signals.
- B The  $\beta$ -galactosidase activity (Miller Units) has been measured from various fusions expressed from a plasmid which also carries *rsaI* gene under its own promoter: *PrpoB::glcU::lacZ::rsaI*, *PrpoB::fn3K::lacZ::rsaI*, *PrpoB::treB::lacZ::rsaI*, *PrpoB::HG001\_1242::lacZ::rsaI*, and *PrpoB::HG001\_2520::lacZ::rsaI* expressed in the mutant strain *HG001-ΔrsaI*. The same constructs were made in the absence of *rsaI* gene. For the *fn3K-lacZ* fusion, we also used an additional construct *PrpoB::fn3K::lacZ::rsalmut4* expressing both the fusion FN3K-LacZ protein and RsaI mut4. The  $\beta$ -galactosidase activity was normalized for bacterial density, and the results represented the mean of four independent experiments. The error bars are standard deviations, and the statistical significance was determined using the Student's *t*-test. \**P* < 0.05, \*\**P* < 0.005, \*\*\**P* < 0.0005, \*\*\*\**P* < 0.0001, ns is for not significant.

Source data are available online for this figure.

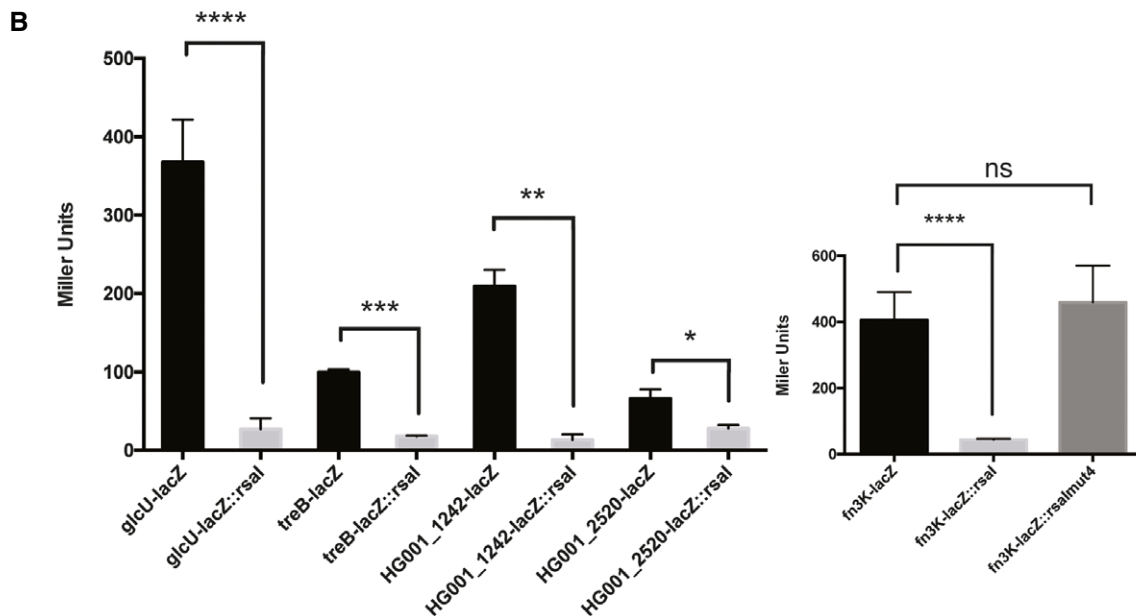
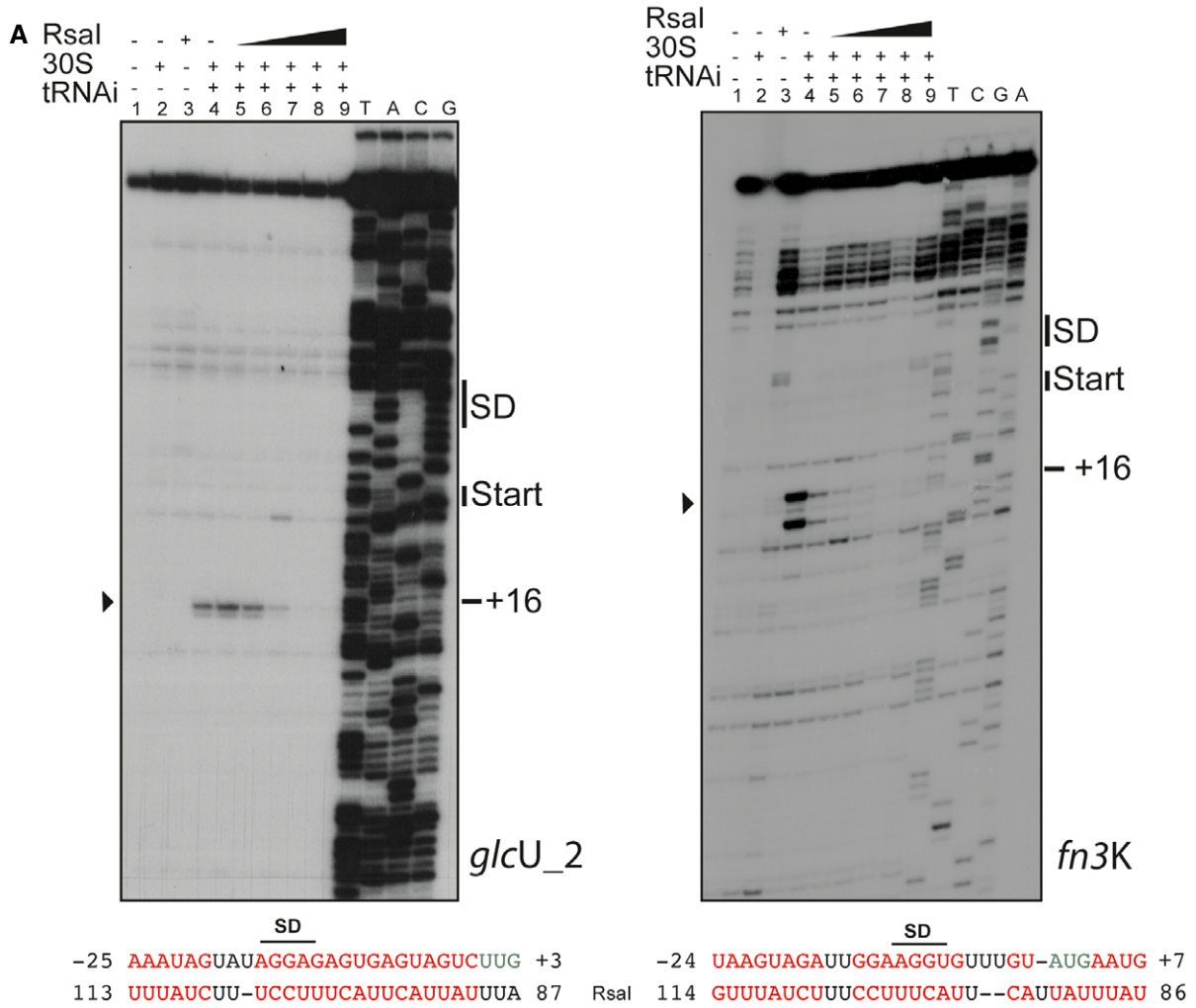


Figure 3.

required for RsaI binding. Instead, the 3'UTR of *icaR* retained the full capacity to efficiently bind RsaI (Fig 4B, Table EV1). We then analyzed whether RsaI binding to the 3'UTR would alter the circularization of *icaR* mRNA. Gel retardation assay was performed with radiolabelled 3'UTR in the presence of the 5'UTR at a concentration of 500 nM leading to almost 50% of binding (Fig EV3A, lane 2). The addition of increasing concentrations of RsaI led to the formation of a 3'UTR-RsaI binary complex but also of a ternary complex involving the three RNA species (Fig EV3A, lanes 4–8). Hence, RsaI does not prevent the formation of the interaction between the two UTRs of *icaR* mRNA *in vitro*.

Then, to analyze the *in vivo* effect of the RsaI interaction with the *icaR* mRNA, we monitored the production of PIA-PNAG as a natural reporter, assuming that variations on IcaR expression would be directly reflected on the production of this exopolysaccharide. For that, dot-blot assays were performed with anti-PIA-PNAG-specific antibodies to monitor the levels of PIA-PNAG in the strain 132, which expresses *rsaI* and produces high levels of this exopolysaccharide compared to HG001 in our experimental conditions. We measure the PIA-PNAG levels in WT 132 strain and the isogenic mutant strains  $\Delta$ *rsaI* and  $\Delta$ 3'UTR *icaR*, which carried a deletion of the 3'UTR of *icaR*. Bacteria were grown for 6 h in TSB containing NaCl, which is required to enhance PIA-PNAG synthesis in 132 strain (Vergara-Irigaray *et al*, 2009) (Fig 4C). The results showed that both RsaI and the 3'UTR of *icaR* are required for efficient production of PIA-PNAG because only the WT strain produces significant levels of exopolysaccharides. In addition, the three strains were transformed with a plasmid overexpressing RsaI or RsaI mut5 carrying a substitution of nucleotides 88–103 (UUUUUACUUA CUUUC to AAUAAUGAAUGAAAGG) under the control of a strong and constitutive promoter. This mutation decreased the stability of RsaI-*icaR* duplex (Fig EV3B). Northern blots confirmed RsaI overexpression in these strains (Fig 4C). Dot-blot results revealed that production of PIA-PNAG could be restored in  $\Delta$ *rsaI* strain when RsaI is constitutively expressed. However, complementation is observed neither in the  $\Delta$ *rsaI* expressing the RsaI mut5 nor in the  $\Delta$ 3'UTR mutant strains overexpressing RsaI or RsaI mut5. These results suggested that the interaction of RsaI with 3'UTR is required to promote PIA-PNAG synthesis (Fig 4C). Interestingly, overexpression of RsaI in the WT strain considerably increased PIA-PNAG production. Although this effect is less pronounced in the WT expressing RsaI mut5, it suggested the presence of additional regulatory pathways.

Taken together, these data suggest that RsaI would contribute to PIA-PNAG synthesis by at least reducing the IcaR repressor protein levels through a specific interaction with the 3'UTR of *icaR* mRNA.

## Two sRNAs responded to sugar uptake

The MAPS experiments revealed that RsaG was the most highly co-purified sRNA together with RsaI (Table EV1), and the complex formed between RsaI and RsaG is very stable (Fig 2C). Interestingly, *rsaG* is localized just downstream *uhpT* encoding the hexose phosphate transporter (Fig EV4A), whose transcription is activated by the two-component system HptRS in response to extracellular glucose-6 phosphate (G-6P), another major carbon source produced by host cells (Park *et al*, 2015). We therefore analyzed whether RsaG expression would also respond to the cellular concentration of G-6P. Northern blot experiments were performed on total RNA extracts produced from the HG001 strain grown in BHI medium supplemented with G-6P. Under these conditions, the synthesis of RsaG is strongly enhanced (Fig EV4B, left panel), in contrast to RsaI, which is completely inhibited under the same conditions (Fig EV4B, right panel). The deletion of *hptRS* considerably reduced the levels of RsaG (Fig EV4C). Therefore, these data indicated that RsaG is activated by HptRS upon G-6P signaling together with *uhpT* and that RsaG sRNA might be derived from the *uhpT* 3'UTR.

We then addressed the consequences of RsaI-RsaG pairings on target recognition. Using gel retardation assays, we analyzed whether RsaG is able to form a ternary complex with RsaI and one of its target mRNA or whether RsaG competes with the mRNA for RsaI binding. These experiments were conducted for three mRNA targets (*glcU\_2*, *HG001\_1242*, and *HG001\_0942*), which all formed stable complexes with the 5' end-labeled RsaI (Figs 2, 5A and EV2, Table EV1). On the opposite, RsaG is not able to interact with *glcU\_2* and *HG001\_1242* as shown the gel retardation assays performed with 5' end-labeled RsaG (Fig 5B). We then performed competition experiments between RsaI and RsaG for mRNA binding. The experiments were performed in the presence of a single concentration of unlabeled RsaG (50 nM) sufficient to bind most of 5' end-labeled RsaI in the presence of increasing concentrations of mRNA (Fig 5A). For the three mRNAs, the addition of RsaG causes the formation of a high molecular weight complex most likely formed by RsaG, RsaI, and the mRNA (Fig 5A). Hence, RsaG does not interfere with the mRNA binding to RsaI.

Furthermore, *in vivo*, the synthesis of the  $\beta$ -galactosidase expressed from the *fn3K-lacZ* fusion was measured in the WT and

### Figure 4. Interaction of RsaI to *icaR* mRNA induces biofilm production.

- A Secondary structure model of *icaR* mRNA. The nucleotides in red are involved in the long-range interaction between the 3' and 5'UTRs or with RsaI. SD and anti-SD are, respectively, for the Shine and Dalgarno sequence and for the sequence complementary to the SD sequence.
- B Gel retardation assays show the formation of the complex between RsaI and *icaR* full-length, *icaR* SUBST, *icaR*-5'UTR, and *icaR*-3'UTR. The 5' end-labeled of RsaI was incubated with increasing concentrations of the various mRNAs. UTR is for untranslated region, and SUBST stands for the substitution of UCCCCUG sequence by AGGGGAC. This mutation, which is located in the 3'UTR of *icaR*, significantly destabilizes the long-range interaction and enhances *icaR* translation. Lane C represents binding between radiolabelled RsaI and full-length *icaR* mRNA (50 nM). Due to the rather short migration of the gel, the full-length mRNA is still observed in the pocket.
- C *In vivo* effect of RsaI on PIA-PNAG synthesis in the *S. aureus* wild-type (WT) 132 strain, the  $\Delta$ *rsaI* mutant, and the strain carrying a deletion of *icaR* 3'UTR ( $\Delta$ 3'UTR). This last mutant strain has been transformed with the pES plasmids expressing *rsaI* or *rsaI* mut5. The PIA-PNAG exopolysaccharide biosynthesis was quantified using dot-blot assays. Serial dilutions (1/5) of the samples were spotted onto nitrocellulose membranes, and PIA-PNAG production was detected with specific anti-PIA-PNAG antibodies. RsaI was detected in the same samples by Northern blot using a probe directed against RsaI. Ethidium bromide staining of rRNA was used as loading controls of the same gel.

Source data are available online for this figure.



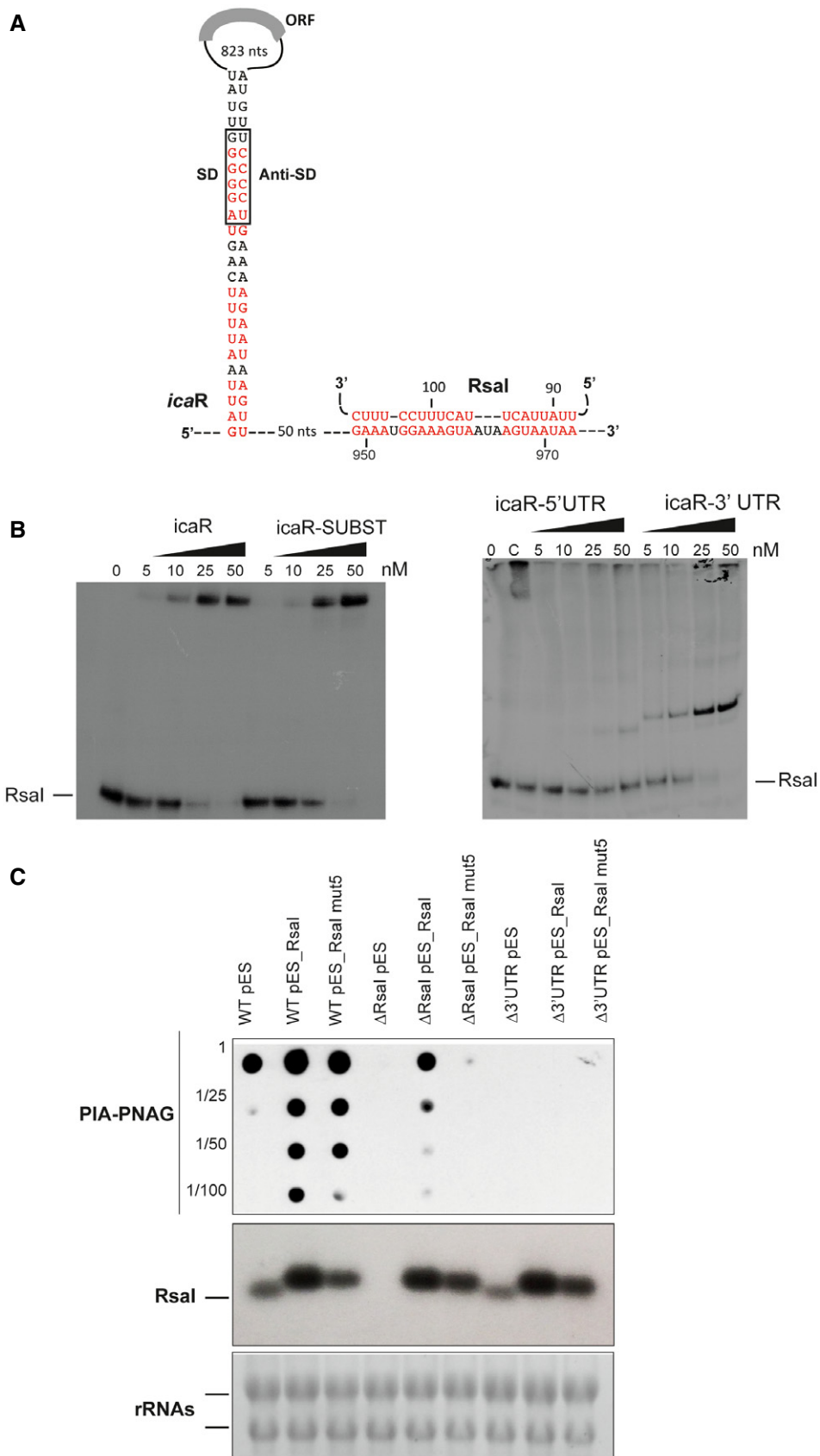
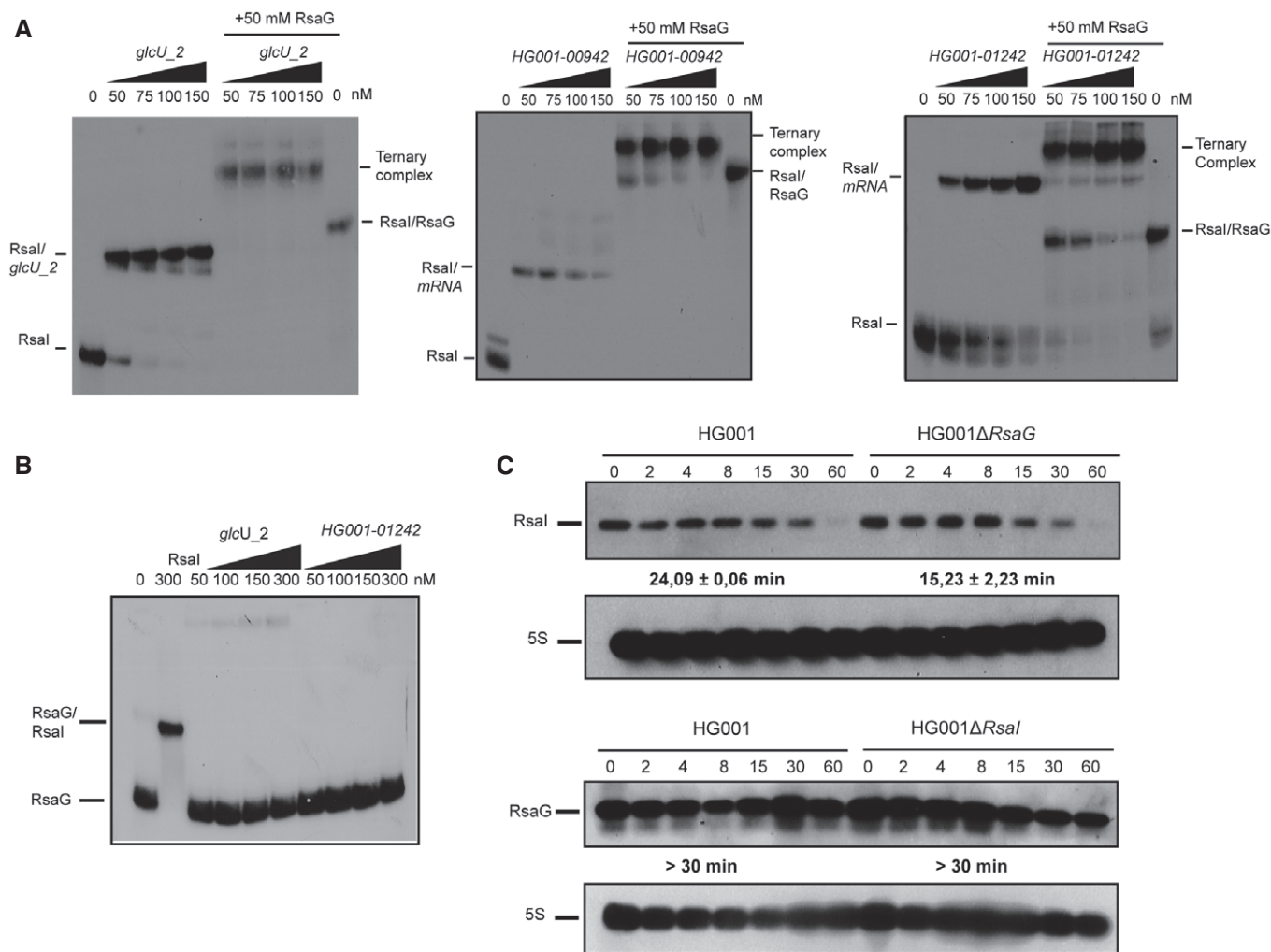


Figure 4.



**Figure 5. Regulatory function of RsaI is not impaired by binding of RsaG.**

**A** Ternary complex formation between RsaI, various mRNAs (*glcU\_2*, *HG001\_00942* and *HG001\_1242*), and RsaG. The 5' end-labeled RsaI was incubated with increasing concentrations of the target mRNA alone or in the presence of 50 nM of RsaG. The various complexes are notified on the sides of the autoradiographies.

**B** RsaG does not form stable complexes with *glcU\_2* and *HG001\_01242*. Binding assays were done in the presence of 5' end-labeled RsaG and either cold RsaI (300 nM) or increasing concentrations of *glcU\_2* and *HG001\_01242*.

**C** Measurements of the half-lives of RsaI and RsaG in *HG001-ΔRsaG* or *HG001-ΔRsaI* mutant strains using Northern blot experiments. The cells were treated with rifampicin at 4 h of growth, and total RNAs were extracted after 2, 4, 8, 15, 30, and 60 min at 37°C in BHI. 5S rRNA was probed to quantify the yield of RNAs in each lane using the same samples, which were however run on two different gels. Calculated half-lives are shown beneath the autoradiographies and are the average of 2 experiments. The data were normalized to 5S rRNA.

Source data are available online for this figure.

*ΔRsaG* mutant strains. The expression of the fusion was strictly identical in both strains. The overexpression of RsaI WT from a plasmid causes a weak but specific inhibition of β-galactosidase synthesis because the RsaI-dependent repression was alleviated in the strain transformed with a plasmid expressing RsaI mut4 (Fig EV4D). This experiment showed that the expression of RsaG does not significantly impact the synthesis of one of the RsaI target.

Using rifampicin treatment, the half-lives of RsaI and RsaG were measured in the WT and the mutant strains (Fig 5C). Quantification of the autoradiographies showed that the two RNAs are highly stable as it was previously shown (Geissmann *et al*, 2009). However, the stability of RsaI was reproducibly higher in the WT strain (24 min) than in the *ΔRsaG* mutant strain (15 min; Fig 5C).

Taken together, the data show that the synthesis of RsaG and RsaI is regulated in response to different but related carbon sources, G-6P and glucose, respectively. In addition, their binding has no major consequences on the RsaI target recognition and regulation, although RsaG has a slight effect on RsaI stability.

#### RsaD, the other sRNA target candidate of RsaI, responds to nitric oxide

The signaling pathway of RsaD, another potential sRNA partner of RsaI, has been studied. RsaD binds to RsaI *in vitro* albeit with a lower affinity than RsaG. Although RsaD was significantly enriched in the MAPS experiments using the wild-type MS2-RsaI (Table EV2),

the data were unfortunately not reliable with the mutant MS2-RsaI mut2 and mut4. This is most probably due to the fact that RsaD is weakly expressed at the stationary phase of growth in BHI medium as revealed by the transcriptomic analysis (Table EV6).

Upstream the *rsaD* gene, we identified a conserved motif AGTGACAA that has been described as a binding site for the response regulator SrrA (Pragman *et al*, 2004; Kinkel *et al*, 2013). SrrAB is a major two-component system affecting the temporal expression of the virulence factors (Pragman *et al*, 2004) while it promotes resistance to nitrosative stress and hypoxia (Richardson *et al*, 2006; Kinkel *et al*, 2013). We demonstrated that the expression of RsaD drops considerably in a  $\Delta$ srrAB mutant strain (Fig EV4E) while its expression is not under the control of the two-component system SaerS, another global regulator of virulence (Fig EV4E; Liu *et al*, 2016). Because SrrAB is able to sense and respond to nitric oxide (NO) and hypoxia (Kinkel *et al*, 2013), we tested the effect of NO on RsaD synthesis by adding diethylamine-NONOate, as it was previously described for RsaE (Durand *et al*, 2015). We observed a significant and reproducible enhanced expression of RsaD in the WT HG001 strain about 10 min after the addition of diethylamine-NONOate to BHI medium (Fig EV4E). RsaD expression decreased after 20 min due to the short half-life of diethylamine-NONOate. Noteworthy, RsaE, which was significantly enriched together with MS2-RsaI, was also shown to be under the control of SrrAB (Durand *et al*, 2015).

These data suggested that through the binding of sRNAs, a network of sRNAs would coordinate sugar metabolism pathways, energy production, and NO stress responses.

### The effect of RsaI on global gene expression in *Staphylococcus aureus*

Before to perform the comparative transcriptomic analysis, we first analyzed whether the deletion or the overexpression of RsaI might cause growth phenotype. Unexpectedly, we did not succeed to overexpress RsaI from a strong promoter in the WT HG001 strain. This result was not due to technical problems as we managed to express RsaI under the control of its own promoter on the same plasmid. Hence, the growth phenotypic experiment was performed in the 132 strain. The WT 132 and the  $\Delta$ rsaI mutant strains harbor a similar growth curve in BHI medium (Fig EV5A). However, when RsaI was expressed under the control of a strong promoter, a slight but reproducible delay in growth was observed only in the WT 132 strain (Fig EV5A). Northern blot analysis was performed to measure the steady-state levels of RsaI during growth in the various strains (Fig EV5B). Quantification of the autoradiographies and normalization with 5S rRNA showed that the highest yields of RsaI were observed in the WT 132 strain expressing RsaI from the strong promoter, and only in this context, a significant amount of RsaI was detected at the beginning of growth (Fig EV5B).

In order to get a global overview of RsaI impact on gene regulation, a comparative transcriptomic analysis was performed on total RNAs extracted from the WT HG001 strain, the isogenic HG001 $\Delta$ rsaI mutant strain, and the same mutant strain complemented with a plasmid expressing RsaI under the control of its own promoter in order to avoid side effects of a too strong expression of RsaI on growth (Tables EV6 and EV7, and Fig EV5C). The cultures were done in triplicates with high reproducibility in BHI medium at 37°C until 6 h, under the conditions allowing the

expression of RsaI (Fig 1). We have considered a gene to be regulated by RsaI if the ratio between two strains was at least twofold. Changes in gene expression were more pronounced between the mutant  $\Delta$ rsaI versus the same strain expressing RsaI from a plasmid rather than between the mutant  $\Delta$ rsaI versus the WT strain (Table EV7). This effect is most likely due to the higher expression of RsaI in the complemented strain in comparison with the WT strain (Fig EV5C). The levels of 26 and 50 mRNAs were significantly decreased and enhanced, respectively, when the complemented strain was compared to the mutant  $\Delta$ rsaI (Table EV7). The complete data set is presented in Table EV6. Beside genes encoding phage-related proteins, most of the RsaI-dependent activation was observed for genes involved in fermentation processes, in NO resistance, and in energy-generating processing (in red in Table EV7). In addition, weaker effects were also observed for mRNAs encoding proteins involved in iron-sulfur cluster repair (ScdA), in NO detoxification (qoxABCD, hmp), and in various metabolic pathways. The level of RsaG was also enhanced in the complemented strain. Conversely, the overexpression of RsaI caused a reduced expression of genes that are functionally related. Several of them are involved in glycolysis (*fba1*) and pentose phosphate pathway (*gnd*, *tkt*), in thiamine co-factor synthesis (*thiW*, *tenA*), and in arginine catabolism (*arcABCD*, *arcR*) (in blue in Table EV7). Additionally, significant repression was also observed for *miaB* encoding tRNA-specific modification enzyme, *tyrS* encoding tryptophanyl-tRNA synthetase, *ebpS* encoding the cell surface elastin binding protein, and to a weaker extent for *rex* encoding redox-sensing transcriptional regulator. Interestingly, the MAPS approach revealed that several of these mRNAs (i.e., *qoxABCD* operon, *tyrS*, and *plfA-plfB*) were also enriched together with RsaI (Table EV2). Surprisingly, the most enriched RsaI targets identified by MAPS (i.e., *icaR*, *glcU\_2*, and *fn3K*) did not show significant mRNA level variations when RsaI was deleted or overexpressed suggesting that RsaI would primarily regulate their translation (Table EV1).

These data showed that high concentrations of RsaI affected the mRNA levels of several enzymes involved in sugar metabolism, in arginine catabolism, in the pentose phosphate pathway, and in various processes linked to NO detoxification, energy production, and fermentation. However, the small overlap between the two approaches suggested that many of the RsaI-dependent effects as revealed in the transcriptomic data resulted from indirect effects.

## Discussion

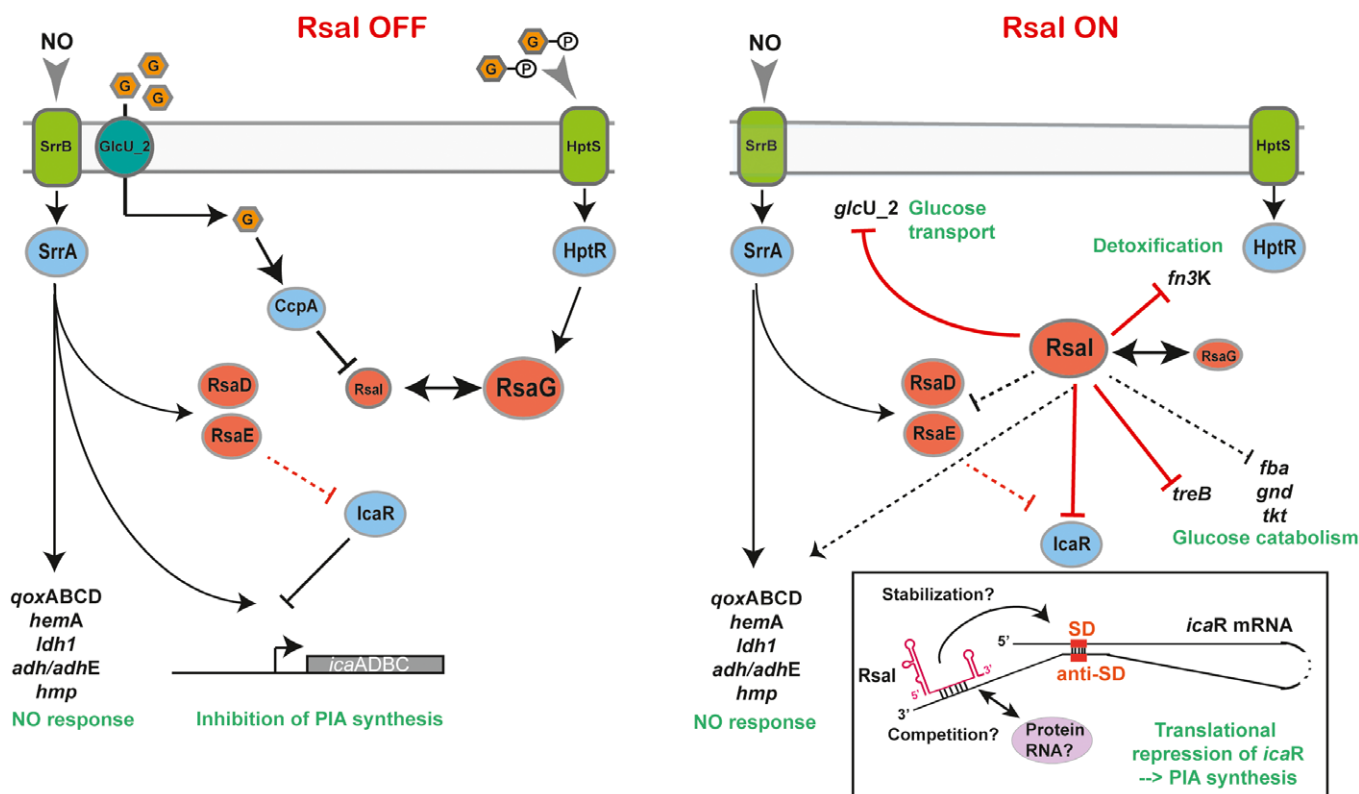
In this work, we have investigated the cellular functions of one abundant sRNA, called RsaI (or RsaOG), which is highly conserved among *Staphylococcaeae* (Geissmann *et al*, 2009; Marchais *et al*, 2010). In contrast to many sRNAs that contained C-track motifs, this RNA has the particularity to carry two conserved G-rich sequences and a large unpaired CU-rich interhelical region (Fig 2). This RNA was proposed to fold into a pseudoknot structure involving the two conserved regions limiting the access of regulatory regions (Marchais *et al*, 2010). Combining several global approaches (MAPS, transcriptomic analysis), our data show that RsaI is linked to a large regulon involved in sugar uptake and metabolism,

biosynthetic and co-factor synthesis, cytochrome biosynthesis, anaerobic metabolism, as well as iron-sulfur cluster repair, and NO detoxification (Fig 6).

**The expression of RsaI is derepressed when glucose concentration decreases**

The cellular level of RsaI is tightly controlled during growth in rich BHI medium. In this study, we showed that RsaI expression is strongly and rapidly repressed at the transcriptional level through the activity of CcpA in response to glucose availability. Carbon catabolite repression is a universal regulatory phenomenon that allows the cells to use the preferred carbon source to produce energy and to provide the building blocks for macromolecules. Concomitantly, it represses genes that are involved in the metabolism of less-preferred carbon sources. To do so, CcpA has to be activated through a cascade of events involving its co-regulator histidine-containing phosphocarrier protein (HPr), which has been phosphorylated by its cognate kinase/phosphorylase HprK/P activated in the presence of glycolytic intermediates. It is thought that binding of the phosphorylated HPr to CcpA enhances its

DNA-binding affinity to the *cre*-binding site to repress or activate the target genes. RsaI repression requires the presence of CcpA, most probably through the binding at a *cre* motif located at -45/-30 (GGAAAACGCTTACAT) from the *rsaI* transcriptional start site (Figs 1 and EV1B). Interestingly, deletion of *ccpA* gene affected vancomycin resistance (Seidl et al, 2009) and recent observations showed that sub-inhibitory treatment of cells with vancomycin led to an enhanced expression of RsaI (Howden et al, 2013). We showed that the repression of RsaI is alleviated as soon as the concentration of glucose is strongly reduced. Therefore, RsaI might be a signature of a metabolic switch of the bacterial population. Using the MAPS approach and gel retardation assays, we could identify several mRNAs that strongly bind to RsaI with its long unpaired and conserved CU-rich region. Binding of RsaI to *glcU\_2* encoding a glucose permease and to *fn3K* encoding the fructosamine 3 kinase hindered the formation of the initiation ribosomal complex. Because their steady-state yields are not affected, we suggest that both mRNAs are regulated at the translational level. We propose that this mechanism is most likely common to all mRNAs that could form base-pairings between their ribosome-binding sites and the CU-rich region of RsaI. This is the case for mRNAs encoding a



**Figure 6. Model of RsaI regulation networks in *Staphylococcus aureus*.**

In the presence of glucose or glucose 6-phosphate (left panel), the expression of RsaI is inhibited, in contrast to RsaG. In the absence of glucose or when glucose is metabolized (right panel), repression of RsaI is alleviated to regulate its RNA targets. In blue, are the transcriptional protein regulators, and in red, the regulatory sRNAs. Gray arrows represent functional links, black arrows are for activation, and bars for repression. Red lines corresponded to post-transcriptional regulation and black lines to transcriptional regulation. Dotted lines represented the regulatory events for which the regulation is not yet demonstrated. A schematic view of the post-transcriptional *icaR* mRNA regulation is represented in the insert. The 3' UTR of *icaR* contains an anti-Shine and Dalgarno (anti-SD) sequence able to bind the SD sequence in the 5' UTR (Ruiz de los Mozos et al, 2013). The CU-rich unpaired sequence of RsaI (in red) binds to the 3' UTR of *icaR* mRNA downstream of the anti-SD and represses the translation indirectly either by stabilizing the interaction between the two UTRs or by preventing the action of trans-acting activators (protein or RNA).

transcriptional regulator of the XRE family, the PTS system trehalose-specific EIIBC component TreB, a peptidase, a cell wall-binding lipoprotein, and DegV containing protein (Table EV2). Noteworthy, *in vivo*, most of these mRNAs were no more found enriched using MS2-RsaI mut4 in which the CU-rich region has been deleted (Table EV5). Among these proteins, two of them are directly involved in sugar uptake and metabolism (GlcU\_2, TreB). In *S. aureus*, the PTS (phosphotransferase system)-dependent and PTS-independent transport systems ensure efficient glucose transport (reviewed in Götz *et al*, 2006). If a rapidly metabolizable sugar (such as glucose) is used during growth at a rather low concentration, the transport will occur via the PTS system, and concomitantly, the carbon catabolite repression system will be activated through CcpA protein. At high concentration of glucose, it is assumed that the sugar will be transported by the PTS system and in addition by the permease GlcU\_2. The glucose transported by the permease will be phosphorylated within the cell by the glucose kinase GlkA. Hence, it is reasonable to propose that RsaI would repress the synthesis of GlcU\_2 when the glucose concentration dropped. In addition, RsaI represses the expression of *treB* mRNA most probably through base-pairings with its SD sequence. TreB is the enzyme specific for trehalose, a diholoside which is transported exclusively by PTS (Bassias & Brückner, 1998). Transcriptomic analysis also revealed that RsaI strongly repressed various key enzymes involved in glucose catabolism pathway such as fructose-biphosphate aldolase (*fbal1*), 6-phosphogluconate dehydrogenase 2C decarboxylase (*gnd*), and the thiamine-dependent enzyme transketolase (*tkl*). Additionally, we showed that RsaI represses the synthesis of fructosamine 3-kinase (Fn3K), which deglycated products of glycation formed from ribose 5-phosphate or erythrose 4-phosphate produced by the pentose phosphate pathway (Gemayel *et al*, 2007). This enzyme is part of a repair machinery to protect the cells from damages caused by glycation as the results of high glucose concentrations (Deppe *et al*, 2011). The pentose phosphate pathway is also an alternative route for glucose metabolism and provides the source of pentose phosphates necessary for nucleotide synthesis. Although it is not known whether RsaI regulated their expression in a direct or indirect manner, base-pairings were predicted between RsaI and the ribosome-binding site of *fbal1* and *tkl* mRNAs. The transketolase is a key enzyme of the pentose phosphate pathway, which requires thiamine diphosphate as a co-factor. Interestingly, the thiamine pathway is also repressed in strain expressing RsaI (Table EV6). Although, we have observed very similar pathways that are deregulated by RsaI using the MAPS and RNA-seq approaches, not so many overlaps were identified. It is possible that the conditions of the MAPS approach performed on the WT strain expressing all the ribonucleases have preferentially enriched the mRNAs that are regulated at the translational level. Therefore, it is tempting to deduct that RsaI would inhibit the synthesis of the major permease of glucose uptake, of enzymes involved in the glycolysis, of unnecessary enzymes involved in detoxification of high glucose concentration, and of the pentose phosphate pathway when glucose concentration decreases (Fig 6).

### RsaI interacts with sRNA containing C-rich motifs

The MAPS approach revealed that several sRNAs (RsaG, RsaD, RsaE) were enriched significantly with RsaI. We demonstrated here

that the second G-track sequence located in the first hairpin domain of RsaI is able to form a highly stable complex with RsaG and to a lesser extent with RsaD. Additionally, mRNA targets interacting with the CU-rich domain of RsaI did not disturb the binding of RsaG suggesting that the two functional domains of RsaI are independent. Conversely, preliminary data suggested that the apical loop of the first hairpin of RsaG contains the C-rich motif that is recognized by RsaI, but which is also used to regulate the expression of several mRNAs (Desgranges E, Marzi S, Romby P, Caldelari I, unpublished data). RsaG is part of the 3'UTR of *uhpT* mRNA, and its expression was strongly enhanced by the extracellular concentration of G-6P. Under these conditions, the levels of RsaI are much lower than for RsaG (Fig EV4B). These two sRNAs are thus involved in pathways related to the use of the preferred carbon sources. Indeed, RsaI negatively controls glucose uptake when glucose is consumed or absent from the medium while RsaG responds to the extracellular concentration of G-6P. Although the functions of RsaG remain to be addressed, we hypothesized that the sRNA might regulate either the expression of unnecessary genes, or of genes involved in sugar metabolism, or of genes required to protect cells against damages linked to sugar-phosphate uptake and metabolism (Bobrovskyy & Vanderpool, 2016). Noteworthy, RsaI sequence is conserved in the *Staphylococcus* genus while RsaG is only conserved in the *S. aureus* species. Our data suggested that RsaG has no major impact on the regulation of RsaI targets. Its slight stabilizing effect on RsaI strongly supports the MAPS data showing that the two sRNAs interact *in vivo* and potentially would form ternary complexes in the presence of the RsaI targets. Whether RsaI might regulate the functions of RsaG remained to be addressed.

In addition, the overproduction of RsaI induced several changes into the transcriptome of *S. aureus* that resembled the regulon of the two-component system SrrAB, which was demonstrated as the essential system responding to both NO and hypoxia (Kinkel *et al*, 2013). The SrrAB regulon has also been shown to confer to the cells the ability to maintain energy production, to promote repair damages, and NO detoxification (Richardson *et al*, 2006; Kinkel *et al*, 2013). We observed here that RsaI enhances the expression of genes encoding cytochrome biosynthesis (*qoxABCD*), as well as genes involved in anaerobic metabolism (*pflAB*, *ldh1*, *focA*, *adh*), in iron-sulfur cluster repair (*scdA*), and most importantly in NO detoxification (*qoxABCD*, *hmp*) and NO resistance (*ldh1*). These effects are most likely indirect and might be due to the interaction between RsaI and the SrrAB-dependent sRNAs RsaD and RsaE, which both contained a typical *srrA* site upstream their genes. These two sRNAs present a C-rich sequence that can potentially form base-pairings with the G-track sequences of RsaI (Table EV1), but the formation of complexes with RsaI is less efficient than with RsaG. It cannot be excluded that an RNA-binding protein might be required in these cases to stabilize the base-pairings. We also do not exclude that the two sRNAs were pooled down because they might share similar mRNA targets with RsaI. RsaD and RsaE are both upregulated by the presence of NO in the cellular medium (Durand *et al*, 2015; Fig EV4D). Although the functions of RsaD require additional studies, *S. aureus* RsaE was previously shown to coordinate the down-regulation of numerous metabolic enzymes involved in the TCA cycle and the folate-dependent one-carbon metabolism (Geissmann *et al*, 2009; Bohn *et al*, 2010) and of enzymes involved in arginine degradation pathway (Rochat *et al*, 2018). Additionally, in

*B. subtilis*, the homologue of RsaE called RoxS is under the control of the NADH-sensitive transcription factor Rex, and the Rex binding site is also conserved in *S. aureus* *rsaE* gene (Durand *et al.*, 2017). Hence, it was proposed that RsaE downregulated several enzymes of the central metabolism under non-favorable conditions and in addition would contribute to readjust the cellular NAD<sup>+</sup>/NADH balance under stress conditions (Bohn *et al.*, 2010; Durand *et al.*, 2015, 2017).

Many of the RsaI-dependent effects, which have been monitored by the transcriptomic analysis, are indirect, and we propose that some of these effects might result from the interaction of RsaI with sRNAs when the preferred carbon source became scarce.

### Physiological consequences of RsaI regulation

Our data suggested that several sRNAs in *S. aureus* are part of intricate regulatory networks to interconnect in a dynamic manner various metabolic pathways following sugar metabolism and uptake. The concept that sRNAs are key actors to coordinate the regulation of metabolic enzymes has been largely demonstrated for *Enterobacteriaceae* (reviewed in Papenfort & Vogel, 2014). These sRNA-dependent regulations often induced significant growth phenotypes in response to the availability of carbon sources and nutrient. For instance, *E. coli* and *Salmonella* SgrS contribute to stress resistance and growth during glucose–phosphate stress by inhibiting synthesis of sugar transporters or activating dephosphorylation and efflux of sugars (e.g., Vanderpool & Gottesman, 2004, 2007; Kawamoto *et al.*, 2005; Rice *et al.*, 2012; Papenfort *et al.*, 2013). The repertoire of SgrS targets has been recently expanded to mRNAs involved in key metabolic pathways, which allow restoring metabolic homeostasis during sugar–phosphate stress and growth recovery (Richards *et al.*, 2013; Bobrovskyy & Vanderpool, 2016). *Salmonella* GcvB regulon is required for growth if peptides represent the unique carbon source (Miyakoshi *et al.*, 2015), while *E. coli* Spot42 is important for optimal utilization of carbon sources and its overproduction inhibited growth on several non-preferred carbon sources (Møller *et al.*, 2002; Beisel & Storz, 2011). In *S. aureus*, the overproduction of RsaE induces a growth defect, which was partially alleviated by acetate (Bohn *et al.*, 2010). These examples illustrate how the yields of these sRNAs should be tightly controlled during growth in order to optimize the use of the preferred carbon sources, to restore metabolic homeostasis during stress, and to avoid cell damages or metabolites depletion caused by the intracellular production of glucose–phosphate. Interestingly, in *Bacillus subtilis*, the sRNA SR1 was found repressed under glycolytic conditions mainly by CcpN and to a lesser extent by CcpA (Licht *et al.*, 2005; Heidrich *et al.*, 2007; Gimpel *et al.*, 2012). SR1 expression is induced by L-arginine and acts as a main regulator of arginine catabolism through the translational repression of *ahrC* mRNA (Heidrich *et al.*, 2007). In this case, the deletion or the overexpression of SR1 did not affect growth (Licht *et al.*, 2005). The deletion of *rsaI* gene does not cause a growth phenotype, and only high levels of RsaI constitutively expressed from a plasmid in the WT strain cause a weak growth delay (Fig EV5A). Additional works will be required to identify the growth conditions under which the function of RsaI can be studied in a more relevant way.

One possible track resides in the fact that RsaI interconnects various metabolic pathways. By acting as a post-transcriptional

regulator, it could play an important role during the infection process. *S. aureus* has the ability to generate infections through the colonization of many different metabolic host niches. Several studies have shown that both glycolysis and gluconeogenesis systems are mandatory for the infection process, and moreover, *S. aureus* appears to be resistant to NO radicals that are heavily produced by the macrophages. Interestingly, glycolysis is an important process that contributes to persist within the macrophages and to protect the intracellular bacteria against NO (Vitko *et al.*, 2015). However, if the bacteria escape the macrophages or lyse the host cells, *S. aureus* is thought to form aggregates at the center of highly inflamed and hypoxic abscesses. Under these conditions, the host cells consumed a large amount of glucose to fight the inflammation. Hence, glucose will become scarce for *S. aureus* suggesting that lactate and amino acids derived from the host might be used as the major sources of carbon to enter gluconeogenesis (Richardson *et al.*, 2015). These conditions favored the activation of the two-component system SrrAB, which in turn activates genes required for anaerobic metabolism, cytochrome, and heme biosynthesis, and NO radical detoxification should play an essential role in the survival of the bacteria (Kinkel *et al.*, 2013). Because the CcpA-dependent repression of RsaI is alleviated under conditions where the glucose is strongly reduced, and because many SrrAB-dependent genes are also induced when RsaI is expressed at high levels, it is thus expected that RsaI might also contribute to metabolic adaptations of the cells to the dynamic nature of the host immune environment. Besides, it is also tempting to propose that RsaI might be involved in the dormant state of bacterial cells while environmental conditions are unfavorable (Lennon & Jones, 2011). Interestingly enough, we also observed a RsaI-dependent activation of the expression of the mRNA encoding EsxA, a type VII-secreted virulence factor required for the release of the intracellular *S. aureus* during infection (Korea *et al.*, 2014). We also showed here that RsaI activates the synthesis of the PIA-PNAG exopolysaccharide most probably by decreasing the translation of IcaR, which is the transcriptional repressor of *icaADBC* operon (Figs 4 and 6). We demonstrated that RsaI base-pairs *icaR* mRNA without interfering the interaction between *icaR* 3'UTR and 5'UTR (Fig EV3). Although the precise molecular mechanism is not yet defined, we propose that RsaI would inhibit the synthesis of IcaR by stabilizing the circularization of the mRNA and/or by counteracting with the binding of an activation factor of *icaR* mRNA translation (Fig 6). Interestingly, RsaE, which is another RsaI target, also binds *icaR* mRNA at two distinct positions close to the SD sequence and in the coding sequence through C-rich motifs (Rochat *et al.*, 2018). These motifs are almost identical to the anti-SD sequence (UCCCC) comprised in the *icaR* 3'UTR (Ruiz de los Mozos *et al.*, 2013). This opens the possibility of a competition for hindering the *icaR* SD by RsaE or the 3'UTR. Thus, RsaI and RsaE might interact individually or simultaneously with *icaR* mRNA to repress its expression in response to changes in cellular medium of glucose and NO. Though further investigations are needed to decipher the physiological consequences of such complex interactions, it indicates that the regulation of the synthesis of the PIA-PNAG is rather complexed and multifactorial and is tightly controlled according to the metabolic state of the bacterial population (Cue *et al.*, 2012). The signals modifying the yield of these RNAs would be crucial to determine the levels of IcaR repressor and PIA-PNAG synthesis. We also do not exclude that RsaI might regulate another regulator of the synthesis

of PIA-PNAG. For instance, inactivation of the TCA cycle resulted in a massive derepression of the PIA-PNAG biosynthetic enzymes to produce the exopolysaccharide (Sadykov *et al*, 2008), and glucose enhances PIA-PNAG-dependent biofilm formation (You *et al*, 2014), while SrrAB appears as an inducer of PIA-PNAG-dependent biofilm (Ulrich *et al*, 2007). It remains to be elucidated if the biofilm induction mediated by SrrAB occurs through activation of RsaE that might promote interaction with *icaR* mRNA and/or RsaI. Hence, *icaR* can be considered as a hub gene, integrating both transcriptional regulatory networks and sRNA-mediated post-transcriptional regulatory signals to control biofilm formation in *S. aureus*.

Although our study shed light on the regulatory activities of this multifaceted sRNA, there is still much to be learned on how RsaI and other sRNAs can be integrated into the networks regulating the metabolic pathways that are essential for *S. aureus* biofilm formation, survival, persistence, and invasion within the host.

## Materials and Methods

### Plasmids and strains constructions

All strains and plasmids constructed in this study are described in Table EV8. The oligonucleotides designed for cloning and for mutagenesis are given in Table EV3. *Escherichia coli* strain DC10B was used as a host strain for plasmid amplification before electroporation in *S. aureus*. Plasmids were prepared from transformed *E. coli* pellets following the NucleoSpin Plasmid kit protocol (Macherey-Nagel). Transformation of both *E. coli* and *S. aureus* strains was performed by electroporation (Bio-Rad Gene Pulser).

The *rsaI* deletion mutant was constructed by homologous recombination using plasmid pMAD in *S. aureus* HG001 and 132 (Arnaud *et al*, 2004). The deletion comprises nucleotides 2,376,101–2,376,306 according to HG001 genome (Caldelari *et al*, 2017). Experimental details are given in supplementary materials.

The vector pCN51::PrsaI was constructed by ligating a PCR-amplified fragment (Table EV3) containing *rsaI* with 166 pb of its promoter region and digested by *SphI* and *PstI* into pCN51 vector digested with the same enzymes. The vector pUC::T7rsaI was constructed by ligating a PCR-amplified fragment (Table EV3) containing *rsaI* with the T7 promoter sequence and digested by *EcoRI* and *PstI* into pUC18 vector digested with the same enzymes. Mutagenesis of pUC::T7rsaI was performed with QuikChange XL Site-directed mutagenesis (Stratagene) leading to pUC::T7rsaI mut1, 2, 3, and 4 (Table EV3). To obtain the plasmid pCN51::P3::MS2-*rsaI*, a PCR product containing the MS2 tag fused to the 5' end of *rsaI* was cloned into pCN51::P3 by digestion of both PCR fragments and of the plasmid by *PstI/BamHI* (Table EV3). Plasmids from positive clones were sequenced (GATC Biotech) before being transformed in DC10B, extracted, and electroporated into *S. aureus* strains. The plasmids pCN51::P3::MS2-*rsaI* mut2 and mut4 were generated by QuikChange mutagenesis as above. Construction of plasmids pES::rsaI and pES::rsaI mut5 is described in supplementary materials.

### Growth conditions

*Escherichia coli* strains were cultivated in Luria-Bertani (LB) medium (1% peptone, 0.5% yeast extract, 1% NaCl) supplemented with

ampicillin (100 µg/ml) when necessary. *Staphylococcus aureus* strains were grown in brain-heart infusion (BHI), tryptic soy broth (TSB), or Muller Hinton broth (MHB) media (Sigma-Aldrich) supplemented with erythromycin (10 µg/ml) for plasmid selection. When needed, MHB was complemented with either 5 g/l of glucose 6-phosphate or 1 g/l of glucose, fructose, or xylose (Sigma-Aldrich). NO production was induced by addition of 100 µM Na-diethylamine-NONOate (Sigma-Aldrich) as previously described (Durand *et al*, 2015).

### Preparation of total RNA extracts

Total RNAs were prepared from *S. aureus* cultures taken at different times of growth. After centrifugation, bacterial pellets were resuspended in 1 ml of RNA Pro Solution (MP Biomedicals). Lysis was performed with FastPrep and RNA purification followed strictly the procedure described for the FastRNA Pro Blue Kit (MP Biomedicals). Electrophoresis of either total RNA (10 µg) or MS2-eluted RNA (500 ng) was performed on 1.5% agarose gel containing 25 mM guanidium thiocyanate. After migration, RNAs were vacuum-transferred on nitrocellulose membrane. Hybridization with specific digoxigenin (DIG)-labeled probes complementary to RsaI, RsaG, RsaD, 5S, *ccpA* sequences, followed by luminescent detection, was carried out as previously described (Tomasini *et al*, 2017).

### MAPS experiments, and transcriptomic and RNA-seq analysis

Crude bacterial extract was prepared and purified by affinity chromatography as previously described (Tomasini *et al*, 2017). The eluted RNA samples were either used for Northern blot or treated with DNase I prior to RNA-seq analysis. Isolation of tagged sRNAs and the co-purified RNAs was performed in duplicates. The experiments were carried out with the tagged wild-type RsaI and two mutant forms (mut2 and mut4). RNAs were treated to deplete abundant rRNAs, and the cDNA libraries were performed using the ScriptSeq complete kit (bacteria) from Illumina. The libraries were sequenced using Illumina MiSeq with a V3 Reagent kit (Illumina), which preserves the information about the orientation of the transcripts and produces reads of 150 nts, which map on the complementary strand. Each RNA-seq was performed at least in duplicates. The reads were then processed to remove adapter sequences and poor-quality reads by Trimmomatic (Bolger *et al*, 2014), converted to the FASTQ format with FASTQ Groomer (Blankenberg *et al*, 2010), and aligned on the HG001 genome (Caldelari *et al*, 2017) using BOWTIE2 (Langmead *et al*, 2009). Finally, the number of reads mapping to each annotated feature has been counted with HTSeq (Anders *et al*, 2015) using the interception non-empty protocol. To estimate the enrichment values for the MAPS experiment or the differential expression analysis for the transcriptomic experiment, we used DESeq2 (Varet *et al*, 2016). The statistical analysis process includes data normalization, graphical exploration of raw and normalized data, test for differential expression for each feature between the conditions, raw p-value adjustment, and export of lists of features having a significant differential expression (threshold *P*-value = 0.05; fold change threshold = 2) between the conditions. All processing steps were performed using the Galaxy platform (Afgan *et al*, 2016).

For total RNA extracts and MS2-eluted RNAs, DNase I (0.1 U/µl) treatment was performed 1 h at 37°C. The reactions mixtures were then purified by phenol:chloroform:isoamylalcohol 25:24:1 (v/v) and subsequent ethanol precipitation. RNA pellets were resuspended

in sterile milliQ water. RNA was quantified by Qubit (Life Technologies), and the integrity was assessed with the Bioanalyzer (Agilent Technologies). For transcriptomic, 1 µg of total RNA was ribo-depleted with the bacterial RiboZero kit from Illumina. The TruSeq total RNA-stranded kit from Illumina was used for the library preparation. Library quantity was measured by Qubit, and its quality was assessed with a TapeStation on a DNA High sensitivity chip (Agilent Technologies). Libraries were pooled at equimolarity and loaded at 7 pM for clustering. The 50 bases oriented single-read sequencing was performed using TruSeq SBS HS v3 chemistry on an Illumina HiSeq 2500 sequencer.

### Preparation of RNAs for *in vitro* experiments

Transcription of RsaI, RsaI mutants, and RsaG was performed using linearized pUC18 vectors. PCR fragments containing the 5'UTR of the coding region of selected mRNA targets were directly used as templates for *in vitro* transcription using T7 RNA polymerase. The RNAs were then purified using a 6% or 8% polyacrylamide-8 M urea gel electrophoresis. After elution with 0.5 M ammonium acetate pH 6.5 containing 1 mM EDTA, the RNAs were precipitated in cold absolute ethanol, washed with 85% ethanol, and vacuum-dried. The labeling of the 5' end of dephosphorylated RNAs (RsaI/RsaI mutants) and DNA oligonucleotides was performed with T4 polynucleotide kinase (Fermentas) and [ $\gamma$ <sup>32</sup>P] ATP. Before use, cold or labeled RNAs were renatured by incubation at 90°C for 1 min in 20 mM Tris-HCl pH 7.5, cooled 1 min on ice, and incubated 10 min at 20°C in ToeP<sup>+</sup> buffer (20 mM Tris-HCl pH 7.5, 10 mM MgCl<sub>2</sub>, 60 mM KCl, 1 mM DTT).

### Gel retardation assays

Radiolabeled purified RsaI or RsaI mutants (50,000 cps/sample, concentration < 1 pM) and cold mRNAs were renatured separately as described above. For each experiment, increasing concentrations of cold mRNAs were added to the 5' end-labeled wild-type or RsaI mutants in a total volume of 10 µl containing the ToeP<sup>+</sup> buffer. Complex formation was performed at 37°C for 15 min. After incubation, 10 µl of glycerol blue was added and the samples were loaded on a 6% or 8% polyacrylamide gel under non-denaturing conditions (300 V, 4°C). Under these conditions where the concentration of the labeled RNA is negligible, the K<sub>D</sub> dissociation constant can be estimated as the concentration of the cold RNA that showed 50% of binding.

### Toe-printing assays

The preparation of *E. coli* 30S subunits, the formation of a simplified translational initiation complex with mRNA, and the extension inhibition conditions were performed as previously described (Fechter *et al.*, 2009). Increasing concentrations of RsaI were used to monitor their effects on the formation of the initiation complex with *glcU\_2* and *fn3K* mRNAs.

### *In vivo* β-galactosidase assays

Translation fusions were constructed with plasmid pLUG220, a derivative of pTCV-*lac*, a low-copy-number promoter-less *lacZ* vector, containing the constitutive *rpoB* promoter (Table EV8). The

whole leader regions of *glcU\_2* (−54/+99), *fn3K* (−33/+99), *treB* (−23/+99), HG001\_01242 (−34/+99), and HG001\_02520 (−71/+99) (Table EV3) were cloned downstream the *PrpoB* in frame with *lacZ*. The whole gene encoding RsaI or RsaI mut4 (obtained synthetically from IDT) with 166 bp of its promoter region was digested with *PstI* and ligated at the unique *PstI* of pLUG220::PrpoB vector. The final constructs were transformed into the strain HG001Δ*rfaI*. β-galactosidase activity was measured four times as previously described (Tomasini *et al.*, 2017).

### PIA-PNAG quantification

Cell surface PIA-PNAG exopolysaccharide levels were monitored according to Cramton *et al.* (1999). Overnight cultures were diluted 1:50 in TSB-3% NaCl, and bacteria were grown at 37°C. Samples were extracted at 6 h after inoculation. The same number of cells of each strain was resuspended in 50 µl of 0.5 M EDTA (pH 8.0). Then, cells were incubated for 5 min at 100°C and centrifuged 17,000 g for 5 min. Supernatants (40 µl) were incubated with 10 µl of proteinase K (20 mg/ml) (Sigma) for 30 min at 37°C. After the addition of 10 µl of the buffer (20 mM Tris-HCl pH 7.4, 150 mM NaCl, 0.01% bromophenol blue), serial dilutions 1:25 were performed in the same buffer. Then, 10 µl was spotted on a nitrocellulose membrane using a Bio-Dot microfiltration apparatus (Bio-Rad). The membrane was blocked overnight with 5% skimmed milk in phosphate-buffered saline (PBS) with 0.1% Tween 20 and incubated for 2 h with specific anti-PNAG antibodies diluted 1:20,000 (Maira-Litrán *et al.*, 2005). Bound antibodies were detected with peroxidase-conjugated goat anti-rabbit immunoglobulin G antibodies (Jackson ImmunoResearch Laboratories, Inc., Westgrove, PA) diluted 1:10,000, using the SuperSignal West Pico Chemiluminescent Substrate (Thermo Scientific).

## Data availability

The transcriptomics and MAPS data have been deposited in NCBI's Gene Expression Omnibus (Barrett *et al.*, 2013) and are accessible through the GEO Series accession number GSE122092 (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE122092>).

**Expanded View** for this article is available online.

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### Author contributions

DB, ED, and IC performed the MAPS experiments and the validation of the data. CJC, LP, AT-A, and IL constructed several mutant strains and performed the biofilm assays. AC and PF have done the transcriptomic analysis. SM analyzed all the RNA-Seq analysis and quantification of the data. KM and FV contributed to phenotypic analysis of some of the mutant strains. PR and IC were responsible of the project and wrote the manuscript. All authors contributed to the analysis, interpretation of the experiments, and writing of the manuscript.

### Conflict of interest

The authors declare that they have no conflict of interest.

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