

Involvement of *Enterococcus faecalis* Small RNAs in Stress Response and Virulence

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Candidate small RNAs (sRNAs) have recently been identified in *Enterococcus faecalis*, a Gram-positive opportunistic pathogen, and six of these candidate sRNAs with unknown functions were selected for a functional study. Deletion mutants and complemented strains were constructed, and their virulence was tested. We were unable to obtain the *ef0869-0870* mutant, likely due to an essential role, and the *ef0820-0821* sRNA seemed not to be involved in virulence. In contrast, the mutant lacking *ef0408-0409* sRNA, homologous to the RNAII component of the toxin-antitoxin system, appeared more virulent and more able to colonize mouse organs. The three other mutants showed reduced virulence. In addition, we checked the responses of these mutant strains to several stresses encountered in the gastrointestinal tract or during the infection process. In parallel, the activities of the sRNA promoters were measured using transcriptional fusion constructions. To attempt to identify the regulons of these candidate sRNAs, proteomics profiles of the mutant strains were compared with that of the wild type. This showed that the selected sRNAs controlled the expression of proteins involved in diverse cellular processes and the stress response. The combined data highlight the roles of certain candidate sRNAs in the adaptation of *E. faecalis* to environmental changes and in the complex transition process from a commensal to a pathogen.

It has now become evident that RNAs are not just involved in transmitting genetic information or have only structural and catalytic roles, like those of mRNA, rRNA, or tRNA, but also act as central regulatory molecules in numerous biological processes. Members of the bacterial small RNA (sRNA) family, including functional *cis*- and *trans*-acting sRNAs, have begun to be well characterized (1–7). Some of these molecules play important roles in bacterial physiology, affecting iron homeostasis; sugar metabolism; responses to stress, such as oxidative stress and conditions induced by DNA damage; stationary phase; cell surface composition; and pathogenesis (8–15). While mechanisms of action, messenger targets, and functions are now partially understood for species like *Escherichia coli*, similar investigations are needed for numerous other bacteria, including *Enterococcus faecalis* (16). Enterococci are normal inhabitants of the gastrointestinal (GI) tracts of humans and animals. While they are typically harmless to healthy individuals, one species, *E. faecalis*, is one of the leading microorganisms involved in hospital-acquired infections, such as catheter-associated urinary tract infections, endocarditis, and surgical and burn wound infections (17–19). Notably, the risk of death for patients infected with multidrug-resistant strains, (i.e., vancomycin-resistant enterococci [VRE]), is considerably higher than for those infected with antibiotic-susceptible strains (20, 21). Unlike Gram-positive pathogens, like *Streptococcus pyogenes* and *Staphylococcus aureus*, *E. faecalis* lacks classical virulence factors, such as proinflammatory toxins or immune modulators, but instead has “opportunism factors.” These factors are directly associated with the organism’s ability to survive hostile conditions, including intrinsic resistance to antibiotics, host immune responses, antagonistic molecules produced by competing microbial flora, and the stresses imposed by its environment (17–19). Several transcriptional regulators in *E. faecalis* that are involved in the stress response, as well as in the virulence of the bacterium,

have been characterized (22–27). However, no investigation of the role of sRNAs has been undertaken. Some studies have been conducted on the roles of sRNAs in the expression of the plasmid-encoded toxin-antitoxin (TA) system and of the CRISPR systems and on the riboswitch regulation of the expression of ethanolamine operon regulation (28–32). In 2008, Livny and coworkers identified 17 putative sRNAs in *E. faecalis* using a bioinformatics approach, followed in 2011 by a study that combined selected *in silico* sRNAs and the 5′ tag rapid amplification of cDNA ends (RACE) method (33, 34). In the same year, the first experimental identification of sRNA in *E. faecalis*, using tiling microarray experiments with probes covering intergenic regions, 5′ and 3′ RACE-PCR, and Northern blot analysis, was published (35). In the last study, around 100 potential sRNA candidates were identified, 11 of which were analyzed further.

In the present study, we focused on 6 of these 11 candidate sRNAs (5 members of the core genome and 1 located in the pathogenicity island [PAI]) by constructing mutants and comple-

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TABLE 1 Bacterial strains and plasmids used

Strain or plasmid	Relevant properties	Reference or source
Strains		
<i>E. faecalis</i>		
EryS	Strain V583 (clinical isolate); erythromycin sensitive	37
EryS derivatives harboring		
Δ0408-0409	EryS isogenic derivative sRNA <i>ef0408-ef0409</i> mutant	This study
Δ0605-0606	EryS isogenic derivative sRNA <i>ef0605-ef0606</i> mutant	This study
Δ0820-0821	EryS isogenic derivative sRNA <i>ef0820-ef0821</i> mutant	This study
Δ1368-1369	EryS isogenic derivative sRNA <i>ef1368-ef1369</i> mutant	This study
Δ3314-3315	EryS isogenic derivative sRNA <i>ef3314-ef3315</i> mutant	This study
Δ0408-0409C	Δ0408-0409 mutant strain derivative complemented	This study
Δ0605-0606C	Δ0605-0606 mutant strain derivative complemented	This study
Δ0820-0821C	Δ0820-0821 mutant strain derivative complemented	This study
Δ1368-1369C _{gene}	Δ1368-1369 mutant strain with pMSP3535 carrying <i>ef1368-ef1369</i> gene; Em ^r	This study
Δ1368-1369C _{sRNA}	Δ1368-1369 mutant strain with pMSP3535 carrying <i>ef1368-ef1369</i> sRNA; Em ^r	This study
Δ3314-3315C	Δ3314-3315 mutant strain derivative complemented	This study
EryS pVEPhoZ	EryS strain with <i>pVEPhoZ</i> integrated in the <i>phoZ</i> locus; Tet ^r	43
EryS pMSP3535	EryS strain with <i>pMSP3535</i> ; Em ^r	42
pVEPhoZ- <i>Pef0408-0409</i>	pVEPhoZ carrying sRNA <i>ef0408-0409</i> promoter; Tet ^r	This study
pVEPhoZ- <i>Pef0605-0606</i>	pVEPhoZ carrying sRNA <i>ef0605-0606</i> promoter; Tet ^r	This study
pVEPhoZ- <i>Pef1368-1369</i>	pVEPhoZ carrying sRNA <i>ef1368-1369</i> promoter; Tet ^r	This study
pVEPhoZ- <i>Pef3314-3315</i>	pVEPhoZ carrying sRNA <i>ef3314-3315</i> promoter; Tet ^r	This study
<i>E. coli</i>		
DH5α	<i>recA endA1 gyrA96 thi-1 hsdR17</i> (r _K ⁻ m _K ⁺) <i>supE44 relA1 lacΔM15</i>	Invitrogen
Plasmids		
pLT06	<i>lacZ</i> P- <i>pheS</i> from pCJK47 <i>cat</i> from pGB354, <i>orfB orfC repA</i> (Ts) <i>orfD</i> from pCASPER	
pLT06-Δ0408-0409	pLT06 derivative carrying a 2-kb DNA fragment from <i>E. faecalis</i> EryS without sRNA <i>ef0408-0409</i>	This study
pLT06wtef0408-0409	pLT06 derivative carrying a 2.1-kb DNA fragment from <i>E. faecalis</i> EryS with sRNA <i>ef0408-0409</i>	This study
pLT06-Δ0605-0606	pLT06 derivative carrying a 2-kb DNA fragment from <i>E. faecalis</i> EryS without sRNA <i>ef0605-0606</i>	This study
pLT06wtef0605-0606	pLT06 derivative carrying a 2.3-kb DNA fragment from <i>E. faecalis</i> EryS with sRNA <i>ef0605-0606</i>	This study
pLT06-Δ0820-0821	pLT06 derivative carrying a 2-kb DNA fragment from <i>E. faecalis</i> EryS without sRNA <i>ef0820-0821</i>	This study
pLT06wtef0820-0821	pLT06 derivative carrying a 2.1-kb DNA fragment from <i>E. faecalis</i> EryS with sRNA <i>ef0820-0821</i>	This study
pLT06-Δ1368-1369	pLT06 derivative carrying a 1.8-kb DNA fragment from <i>E. faecalis</i> EryS without sRNA <i>ef1368-1369</i>	This study
pLT06-Δ3314-3315	pLT06 derivative carrying a 1.7-kb DNA fragment from <i>E. faecalis</i> EryS without sRNA <i>ef3314-3315</i>	This study
pLT06wtef3314-3315	pLT06 derivative carrying a 2.1-kb DNA fragment from <i>E. faecalis</i> EryS with sRNA <i>ef3314-3315</i>	This study
pVEPhoZ	pVEPhoZ carrying the 5' part of the <i>phoZ</i> gene; Tet ^r	43
pVEPhoZ- <i>Pef0408-0409</i>	pVEPhoZ carrying the sRNA <i>ef0408-0409</i> promoter; Tet ^r	This study
pVEPhoZ- <i>Pef0605-0606</i>	pVEPhoZ carrying the sRNA <i>ef0605-0606</i> promoter; Tet ^r	This study
pVEPhoZ- <i>Pef1368-1369</i>	pVEPhoZ carrying the sRNA <i>ef1368-1369</i> promoter; Tet ^r	This study
pVEPhoZ- <i>Pef3314-3315</i>	pVEPhoZ carrying the sRNA <i>ef3314-3315</i> promoter; Tet ^r	This study
pMSP3535	8.35 kb; <i>nisR nisK PrnA</i> (nisin-inducible promoter); Em ^r	42
pMSP3535- <i>ef1368-ef1369</i> gene	pMSP3535 carrying <i>ef1368-ef1369</i> gene; Em ^r	This study

^a Em, erythromycin; Tet, tetracycline; Cm, chloramphenicol.

mented strains in order to determine whether they were implicated in pathogenesis and the stress response in *E. faecalis*. It appeared that the *ef0869-0870* sRNA might be essential, since construction of the corresponding mutant strain always failed. The five other candidate sRNA mutant strains (Δ*ef0408-0409*, Δ*ef0605-0606*, Δ*ef0820-0821*, Δ*ef1368-1369*, and Δ*ef3314-3315*) were tested in the *Galleria mellonella*, urinary tract infection, and intracellular-macrophage survival models. For those that were involved in the virulence of *E. faecalis*, we looked at their phenotypes in response to several stress conditions potentially encountered by the bacteria in their niches or during the infection process. Lastly, to identify the regulons, we performed a 2-dimensional (2D) polyacrylamide gel electrophoresis (PAGE) technique in combination with mass spectrometry (MS) to compare the cytoplasmic proteomes of sRNA mutants with those of their corresponding wild-

type strains. This study highlights for the first time the implication of certain candidate sRNAs in the virulence and the stress response of the opportunistic pathogen *E. faecalis*.

MATERIALS AND METHODS

Bacterial strains, plasmids, and growth culture conditions. The bacterial strains and plasmids used in this study are listed in Table 1. *E. coli* strains were cultured with shaking at 37°C in LB medium with chloramphenicol (10 μg/ml), erythromycin (100 μg/ml), or tetracycline (10 μg/ml) when required. *E. coli* DH5α (36) was used as the recipient for cloning. Mutants were constructed from the parental *E. faecalis* strain EryS, an erythromycin-sensitive strain derived from the vancomycin-resistant clinical isolate V583 (37). *E. faecalis* EryS and its derivatives were grown without shaking at 37°C in M17 medium supplemented with 0.5% glucose (M17glu). Growth of the *E. faecalis* strains was also examined in horse serum under oxidative-stress conditions (with shaking in M17glu supple-

mented with 1.5 mM H₂O₂ or in CCM17 MOPS [morpholinepropane-sulfonic acid] medium without carbon source supplemented with 0.5% glycerol), under osmotic-stress conditions (growth in M17glu supplemented with 8% sodium chloride), or under detergent conditions (growth in M17glu supplemented with 0.08% bile salts). For the acid challenge assays, overnight cultures of the wild type, the sRNA mutants, and the complemented strains were diluted to a final optical density at 600 nm (OD₆₀₀) of 0.01 in fresh medium and allowed to grow to an OD₆₀₀ of 0.5. Then, cells were harvested by centrifugation and resuspended in the same volume of M17/7, pH 2.8. Determination of the number of CFU was performed by plating on M17glu after serial dilution and counting colonies after incubation at 37°C for 24 h. The survival rates (the number of CFU after exposure to the different conditions divided by the number of CFU under control conditions) were determined. Each survival experiment was performed at least three times.

General molecular methods. PCR was performed with Phusion High-fidelity DNA polymerase (Finzymes, Vantaa, Finland). The primers used for this work are listed in Table S1 in the supplemental material. PCR products and plasmids were purified using a Nucleospin plasmid kit (Macherey-Nagel, Düren, Germany). Restriction endonucleases and T4 DNA ligase were purchased from Promega (Madison, WI, USA) and used according to the manufacturers' instructions. Genomic-DNA extraction and other standard methods were carried out as described by Sambrook (38).

Construction of sRNA mutants and complemented strains and transcriptional fusion. For the construction of the sRNA mutant strain, allelic replacements were carried out as previously described (39, 40). Briefly, DNA fragments (obtained using PCR with chromosomal DNA of *E. faecalis* EryS as the template) containing ligated upstream and downstream sequences of the desired deletion were cloned into the plasmid pLT06 (41) (Table 1), and 1 µg of recombinant plasmid was used to transform *E. faecalis* competent cells. For the construction of the $\Delta ef0408-0409$, $\Delta ef0605-0606$, $\Delta ef0820-0821$, $\Delta ef0869-0870$, $\Delta ef1368-1369$, and $\Delta ef3314-3315$ mutant strains, fusion PCR products of the respective upstream (955, 989, 1,056, 1,020, 765, and 836 bp, respectively) and downstream (1,066, 1,030, 967, 958, 996, and 869 bp, respectively) fragments were constructed. Single-crossover transformants (dark-blue and chloramphenicol-resistant colonies) were used for temperature shifts in order to release the plasmid. Candidate clones resulting from a double-crossover event were isolated on M17glu agar with or without chloramphenicol. In antibiotic-susceptible clones, the loss of the plasmid and the deletion of the respective sRNA were verified by PCR and sequencing.

For construction of complemented mutant strains, the pLT06 plasmid containing the intact sRNA was introduced into the corresponding sRNA mutant. Double-recombination events allowed the restoration of intact sRNA. For the complementation of the antisense sRNA *ef1368-1369*, a DNA fragment containing the entire *ef1369* gene or the sRNA antisense to the *ef1369* gene of *E. faecalis* EryS was obtained by PCR using primers listed in Table S1 in the supplemental material. The PCR product was cloned into the pMSP3535 plasmid downstream of the nisin-inducible promoter (*Pnis*) (42). The recombinant plasmid was first transformed into *E. coli* DH5 α cells, amplified, and finally used to transform competent cells of the *E. faecalis* $\Delta ef1368-1369$ mutant. After electroporation, the colonies were analyzed by sequencing for the presence of the plasmid containing an intact *ef1369* gene or corresponding antisense sRNA.

For the transcriptional fusion constructions, the 201-bp, 135-bp, 98-bp, 194-bp, and 163-bp putative promoter regions of sRNAs corresponding to *ef0408-0409*, *ef0605-0606*, *ef0820-0821*, *ef1368-1369*, and *ef3314-3315*, respectively, were cloned into the pVEPhoZ plasmid (43). This integrative plasmid was then introduced into the *E. faecalis* EryS chromosome by single crossover in the *phoZ* locus, as described by Le Jeune et al. (43).

Infection experiments and survival in macrophages. Infection of *G. mellonella* larvae with *E. faecalis* was accomplished as previously described by Lebreton et al. (21). Briefly, using a syringe pump (KD Scientific, Hol-

iston, MA, USA), larvae (about 0.3 g and 3 cm in length) were infected subcutaneously with washed *E. faecalis* wild-type sRNA mutants and the complemented strains from an overnight culture in M17glu, with $1.2 \times 10^5 \pm 0.5 \times 10^5$ CFU per larva administered in 10 µl of sterile saline buffer. In each test, 15 insects were infected, and the experiments were repeated at least five times. Larval killing was then monitored at 20 h postinfection. The results were analyzed by an unpaired *t* test, and all comparisons with a *P* value of less than 0.05 were considered significant.

The virulences of the wild-type strain and the sRNA deletion mutants were also assessed in a urinary tract infection (UTI), essentially as previously described by Lebreton et al. (21). Aliquots of 100 µl from each strain ($\sim 10^8$ cells) were used to inject each of 10 female BALB/c mice (10 weeks old; Harlan Italy S.r.l., San Pietro al Natisone, Udine, Italy) transurethrally. The mouse experiments were performed under a protocol approved by the Institutional Animal Use and Care Committee at the Catholic University of the Sacred Heart, Rome, Italy, and authorized by the Italian Ministry of Health (permit number Z21, 13 March 2012). Infection experiments were repeated three times. Mice were monitored twice daily and were euthanized by CO₂ asphyxiation 7 days after infection. For determination of the numbers of CFU present in organs, serial homogenate dilutions were plated onto *Enterococcus* selective agar (Fluka Analytical, Switzerland). CFU counts were analyzed by an unpaired *t* test.

Survival of *E. faecalis* in mouse peritoneal macrophages was tested using an *in vivo-in vitro* infection model, as described previously (44). Bacterial cells were pelleted and resuspended in an adequate volume of phosphate-buffered saline (PBS) for injection. Male BALB/c mice (10 weeks old; Harlan Italy S.r.l., San Pietro al Natisone, Udine, Italy) were infected with $4.8 \times 10^7 \pm 0.3 \times 10^7$ cells of each strain (estimated by CFU determination) by intraperitoneal injection of 200 µl of the PBS-enterococcus suspension. After a 6-h infection period, the peritoneal macrophages were collected by peritoneal lavage, centrifuged, and suspended in Dulbecco's modified Eagle's medium (DMEM) containing 10 mM HEPES, 2 mM glutamine, 10% fetal bovine serum, and 1× nonessential amino acids, supplemented with vancomycin (10 µg/ml) and gentamicin (150 µg/ml). The cell suspension was dispensed into 24-well tissue culture plates and incubated at 37°C under 5% CO₂ for 2 h. After exposure to antibiotics (i.e., 8 h postinfection) to kill extracellular bacteria, the infected macrophages were washed, and triplicate wells of macrophages were lysed with detergent. After dilution with brain heart infusion (BHI) broth, the lysates were plated on BHI agar to quantitate the viable intracellular bacteria. The remaining wells of infected macrophages were maintained in DMEM with the antibiotics for the duration of the experiment. The same procedure was performed at 24, 48, and 72 h postinfection. In the meantime, at 8, 24, 48, and 72 h postinfection, supernatant fluids from each well were removed, and extracellular bacteria were quantitated by counting on BHI agar plates. To assess their viability, macrophages were detached from tissue culture wells with cell scrapers and stained with trypan blue, and viable macrophages were counted with a hemacytometer. The procedure was repeated three times, and the results were analyzed using one-way analysis of variance with a Bonferroni correction posttest with SPSS statistical software (SPSS, Chicago, IL, USA). All statistical analyses were performed using Prism software (version 5.00) for Windows (GraphPad Software, San Diego, CA, USA). For all comparisons, a *P* value of less than 0.05 was considered significant.

Alkaline phosphatase assays. For alkaline phosphatase (AP) assays, overnight cultures grown in M17glu were diluted in fresh medium to an OD₆₀₀ of 0.01. At an OD₆₀₀ of 0.5, the stressing agents mentioned above for growth conditions were added to the cultures, and after 30, 60, and 90 min of incubation at 37°C, 10 ml of culture was harvested. AP activity measurements were performed as described by Le Jeune et al. and were expressed in Miller units (MU) by the following formula: $MU = (OD_{405} \times 1,000) / (OD_{600} \times \text{volume [ml]} \times \text{time [min]})$ (43).

RNA isolation and reverse transcriptase-quantitative PCR (RT-qPCR). In order to assess comparative transcriptional gene expression, we used the EryS wild-type strain and its sRNA derivative mutants cultured

on M17glu at 37°C in exponential phase. Total RNA from three biological replicates was extracted by a method based on the protocol of Toledo-Arana et al. and modified as described by Michaux et al. (40, 44).

For RT-PCR experiments, 2 micrograms of RNA was reverse transcribed with random-hexamer primers and QuantiTect enzyme (Qiagen, Valencia, CA) according to the manufacturer's recommendations. cDNAs were then used as templates for PCRs using primers listed in Table S1 in the supplemental material. Quantification of 23S rRNA and *gyrA* (encoding the A subunit of the DNA gyrase enzyme) mRNA provided internal controls. Amplification (using 5 µl of a 1/100 cDNA dilution), detection (with automatic calculation of the threshold value), and real-time analysis were performed three times for each cDNA sample, using the iCycler iQ detection system (Bio-Rad Laboratories). Relative mRNA levels of each gene in each sample were calculated using comparative cycle time, as described previously by Meijerink et al. (45).

Protein extraction, 2D PAGE technique, protein identification, and Western blotting. Cytoplasmic proteins were prepared as described by Becher et al. (46). In brief, 50 ml of bacterial cell culture was centrifuged (7,000 × g) for 10 min at 4°C. The cell pellets were washed twice with 1 ml of ice-cold Tris-EDTA (TE) buffer and transferred into screw-top tubes containing 500 µl of glass beads. The cells were disrupted using a Fast Prep instrument (MP Biomedical LLC, Santa Ana, CA, USA) for 30 s at 6.5 m/s. The lysate was centrifuged for 25 min at 21,000 × g at 4°C to remove the cell debris, and the supernatant was transferred into a new tube and centrifuged for 45 min at 21,000 × g at 4°C in order to remove insoluble and aggregated proteins. Proteins were prepared from three biological replicates.

2D PAGE was performed using an immobilized pH gradient (IPG) technique as described previously (47). In the first dimension, proteins were separated on IPG strips (GE Healthcare, Uppsala, Sweden) in a linear pH range of 4 to 7. One hundred micrograms of protein extracts was loaded onto the IPG strips. The second dimension was performed as described by Engelmann and Hecker (48). The resulting gels were stained with the fluorescent dye Krypton according to the manufacturer's instructions (Thermo Scientific, Waltham, MA, USA) and scanned using a Typhoon 9400 Variable Mode Imager. For quantitation of proteins, the 2D gel image analysis was performed with the software DELTA 2D version 4.3 (Decodon GmbH, Greifswald, Germany) as described previously (49). Identified protein spots were labeled with the respective protein names or locus numbers on the gel. Multiple spots of the same protein were numbered consecutively. Gel-to-gel variations of spot positions on 2D gel images were first compensated for by warping (exact mode) of the images. Subsequently, they were fused intraexperimentally to a virtual fusion gel (union mode). This recomposed gel was used for protein spot detection. Automatically calculated spot borders were corrected and optimized manually if necessary. The final spot mask was transferred from the fusion gel to each gel image of the corresponding experiment. Spot intensities were normalized using total normalization. For each protein spot, the amounts of protein in the mutant sample and in the corresponding protein spot of the wild-type sample were calculated and presented as the protein ratio. For statistical analyses, standardized data were loaded into the J. Craig Venter Institute Multiexperiment Viewer 4.4.1. Each experiment was analyzed separately. To test whether the amount of protein in a given spot changed significantly in the mutant, we used *t* test analysis (Welch approximation; *P* values based on permutation and based on all possible permutations; adjusted Bonferroni correction). Only protein spots whose amount changed at least 2-fold were considered differentially expressed.

For protein identification, protein spots of interest were excised from the gel with an Ettan spot picker (GE Healthcare) with a picker head of 2 mm. In-gel digestion was performed with sequencing grade trypsin (Promega, Madison, WI), and extraction of peptides was carried out with the Ettan Spot Handling Workstation (GE Healthcare) using the protocol described by Eymann et al. (50). Identification of *E. faecalis* proteins by matrix-assisted laser desorption ionization–time of flight (MALDI-TOF)

and MALDI-TOF MS was carried out as described previously (51). MALDI-TOF MS analyses of spotted peptide solutions were carried out on a Proteome Analyzer 4800 (Applied Biosystems, Foster City, CA, USA) (for details, see reference 52). For database searches, the Mascot search engine version 2.1.04 (Matrix Science Ltd., London, United Kingdom) was used, with an *E. faecalis* sequence database extracted from the National Center for Biotechnology Information (NCBI) bacterial genomes. Peptide mixtures that yielded a Mowse (molecular weight search) score of 59 at least twice were regarded as positive identifications. Details of the mass spectrometry results for the identified proteins are presented in Table S2 in the supplemental material.

For Western blot analysis, the proteins were transferred after electrophoresis onto a polyvinylidene difluoride membrane, which was then stained with Coomassie blue in order to verify that equal amounts of protein were present in all lanes. Blocking, incubation with antibodies against *E. faecalis* Ers and *E. coli* GroEL, and enhanced-chemiluminescence detection (GE Healthcare, Little Chalfont, United Kingdom) were carried out as previously described by Riboulet-Bisson et al. (53).

In order to predict target genes for the identified sRNAs, the RNA Predator (http://rna.tbi.univie.ac.at/RNAPredator2/target_search.cgi) (54) and CopraRNA (<http://rna.informatik.uni-freiburg.de/CopraRNA/Input.jsp>) (55) servers were used.

RESULTS AND DISCUSSION

Selection of sRNAs and construction of corresponding mutants.

In previous work performed in our laboratory, we characterized 11 candidate sRNAs highly expressed in *E. faecalis* V583 using tiling microarray, 5' and 3' RACE-PCR, and Northern blot analysis (35). The absence of a start codon and a putative ribosome binding site (RBS) in a consensus distance (8 nucleotides [nt]) from the start codon within the studied transcripts strongly suggests the lack of a putative open reading frame. *ef1097-1098*, which corresponded to *E. faecalis* transfer-messenger RNA (tmRNA) (*ssrA*), a unique bifunctional RNA acting as both a tRNA and an mRNA (56, 57); *ef2205-2206* sRNA, which corresponded to the RNA component of the signal recognition particle (SRP) (58); and *efa0080-A0081* and *efB0062-B0063*, the two plasmidic RNAII components of the TA system already identified in *E. faecalis* V583 (29, 30), were not analyzed further. Note that tmRNA is essential for some species, such as *Neisseria gonorrhoeae*, *Haemophilus influenzae*, and *Shigella flexneri* (56, 59). Similar observations were made here, where all attempts to construct a knockout mutant in *E. faecalis* failed (data not shown). To obtain functional insights into the other previously identified candidates, members of the core genome or located in the PAI with unknown roles, deletions were introduced into the genes corresponding to *ef0408-0409*, *ef0605-0606*, *ef0820-0821*, *ef0869-0870*, *ef1368-1369*, and *ef3314-3315* sRNAs. The deletion of the sRNA region, depending on the candidate, was between 30% and the entire sRNA (see Fig. S1 in the supplemental material). Note that, except for *ef0605-0606*, located on the PAI (not systematically observed *E. faecalis* genomes), the presence of selected candidate sRNAs has been shown in the 54 sequenced strains of *E. faecalis* available (35).

In the case of *ef0869-0870*, we tried to construct a mutant strain using the double-crossover strategy with a thermosensitive replication plasmid. Unfortunately, all antibiotic-sensitive clones obtained after the second recombination (several hundred were analyzed) harbored the wild-type gene. This strongly suggests that the sRNA *ef0869-0870* may be essential for *E. faecalis*.

Because *ef1368-1369* likely acts as an antisense RNA of the *ef1369* gene, two different complemented strains were constructed using the replicative vector pMSP3535: the first contained the

gene *ef1369*, and the second expressed the sRNA. With these two strains, we would be able to verify whether any phenotypes observed for the mutant (see below) were due to lack of the gene or the sRNA. Before virulence assays or stress challenges, growth assays under standard conditions (M17glu at 37°C) were performed for the wild type and strains with sRNAs deleted without any obvious difference between them.

Implication of sRNAs in virulence, colonization, and survival within macrophages. In order to determine the impacts of the sRNA deletions on the virulence of *E. faecalis* and their roles in the process of infection, different infection models and assays were employed. First, we compared the survival of larvae of *G. mellonella* infected with the sRNA mutant and the corresponding complemented strains with those infected with the wild type. Infection of *G. mellonella* worms has emerged as a reliable model system to study the pathogenesis of numerous human pathogens (60–66). Indeed, the growth of the larvae at 37°C and the high degree of structural and functional homology between the innate immune systems of *Galleria* and mammals make the model pertinent (67, 68). The nonpathogenic Gram-positive bacterium *Lactococcus lactis* IL1403 was used as a control, and all larvae infected with the organism survived, using the same infection protocol as for *E. faecalis* (data not shown). This model of virulence was used as the first screen for our functional analysis. The mutant strain showing a phenotype was then used for further studies (urinary tract infection, survival in macrophages, stress response, and proteomic study), whereas in the case of the nonvirulent phenotype, the corresponding strain was not further studied. The figures (and supplemental data) present the results when a difference between the mutant and the wild-type strains was observed.

ef0408-0409 sRNA is homologous to the RNAII component of the TA system (also named the *par* system) described previously (29, 30). These systems, also called postsegregation killing systems, first identified on bacterial plasmids, encode a stable toxin and an unstable antitoxin. After cellular division, the loss of the plasmid leads to degradation of the antitoxin and corresponding activation of the toxin. In most of the systems, both components are proteins, but in some cases, as in the well-studied *par* system in *E. faecalis*, the antitoxin is a small regulatory RNA that represses the translation of the toxin mRNA (69). In *E. faecalis* V583, three of these systems were identified in previous studies, two on plasmids and one on the chromosome (35). It has been shown that the *0408-0409 par* addiction module is located on the bacterial chromosome, which raises the question of its cellular role (69, 70). Of note, two direct-repeat sequences allowing the base pairing of the two components of the toxin-antitoxin have been found in the sRNA and *ef0409* sequences. In addition, the two loci showed the same bidirectional terminator (35). As shown in Fig. 1, the rate of killing of the larvae infected with the $\Delta ef0408-0409$ mutant strain was significantly higher than the rate for those infected with the wild type. After 20 h of infection, 15% of the caterpillars infected with the wild-type strain were still alive, whereas less than 3% of the animals infected with the $\Delta ef0408-0409$ mutant strain had survived. The “hypervirulence” phenotype was attenuated back to wild-type level in the complemented $\Delta ef0408-0409$ mutant strain, demonstrating that the observed phenotype was due to the deletion of the sRNA. It was surprising that we were able to create a mutant lacking the *ef0408-0409* sRNA, which is supposed to act as an antitoxin through interaction with RNAI (*ef0409*), encoding the Fts toxin (30, 69). Nevertheless, it is worth noting

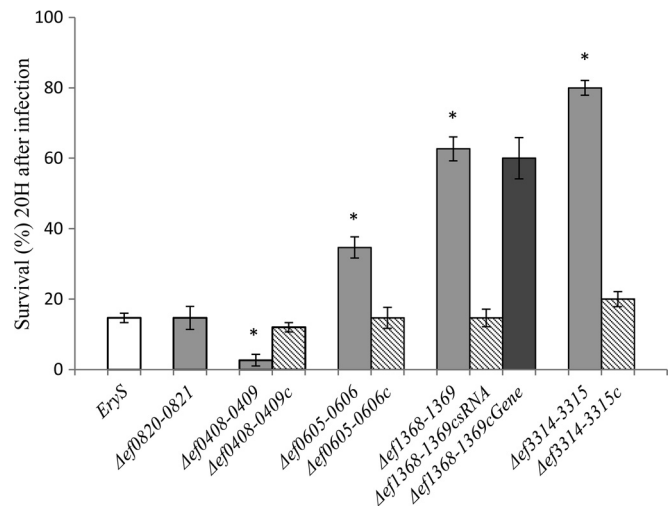


FIG 1 Effects of inactivation of selected sRNAs on virulence. Shown is the percent survival of *G. mellonella* at 20 h after infection with around 1.2×10^5 CFU per larva of *E. faecalis* EryS V583 (wild type); the $\Delta ef0408-0409$, $\Delta ef0605-0606$, $\Delta ef0820-0821$, $\Delta ef1368-1369$, and $\Delta ef3314-3315$ sRNA mutants; and the corresponding complemented strains when required. Note the presence of two complemented strains for the $\Delta ef1368-1369$ mutant, $\Delta ef1368-1369csRNA$, which represents the small-RNA strand complementation, and $\Delta ef1368-1369cGene$, the gene strand complementation. Experiments were repeated at least five times, and the results represent the means \pm standard deviations. *P* values of less than 0.001 (*) for the survival of the sRNA mutant strains compared to the wild type were considered to be significant.

that our transcriptional analysis showed that RNAI (*ef0409*) was transcribed at a very low level (near background) in both the wild-type and $\Delta ef0408-0409$ mutant strains.

The $\Delta ef0605-0606$, $\Delta ef1368-1369$, and $\Delta ef3314-3315$ mutant strains were all less virulent than the wild type. At 20 h postinfection, 35, 63, and 80% of the larvae, respectively, were still alive versus 15% of the larvae infected with the wild type (Fig. 1). Virulence was restored to the wild-type level with the corresponding sRNA-complemented strains. Due to the antisense nature of the $\Delta ef1368-1369$ sRNA, complementation was also tested with the construction containing the *ef1369* gene, encoding a putative transcriptional regulator. In this case, no complementation was observed (Fig. 1). In addition, no difference in caterpillar survival was observed for the wild type with or without the empty pMSP3535 vector (data not shown).

The sRNA *ef0820-0821* seems not to be involved in the virulence of *E. faecalis*, since the survival of larvae infected with the mutant strain was comparable to that of larvae infected with the wild type (Fig. 1). Consequently, we did not further investigate this sRNA.

E. faecalis is one of the leading etiological agents of UTI (71). Therefore, we tested the fate of the $\Delta ef0408-0409$, $\Delta ef0605-0606$, $\Delta ef1368-1369$, and $\Delta ef3314-3315$ mutants, as well as the corresponding complemented and wild-type strains, in kidneys after urinary tract infection of mice. As shown in Fig. 2, the $\Delta ef0408-0409$ and $\Delta ef1368-1369$ mutants were around 0.5 log units more abundant in the kidneys than the wild-type and the complemented strains (with *P* values of 0.0029 and 0.0115, respectively). The UTI experiments performed with the $\Delta ef0605-0606$ sRNA and the $\Delta ef3314-3315$ sRNA mutant strains did not show any difference between the wild type and the mutants (data not shown).

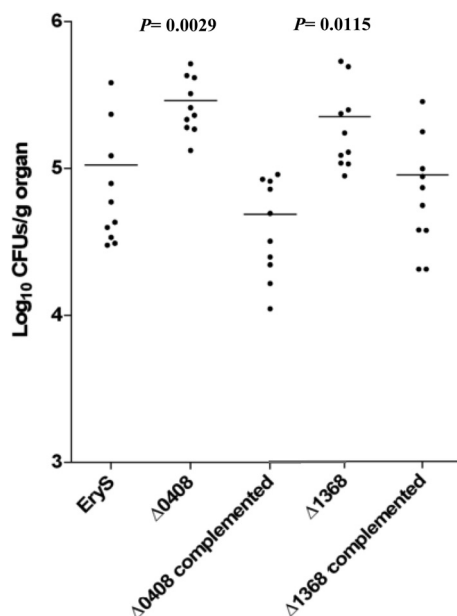


FIG 2 Bacterial persistence within the kidney in urinary tract infection. Enterococcal tissue burdens in kidneys from BALB/c mice infected intravenously with 1×10^9 bacteria/ml of the EryS wild type, the $\Delta ef0408-0409$ and $\Delta ef1368-1369$ mutants, and the corresponding complemented strains are shown. Kidney pair homogenates were obtained from groups of 10 mice sacrificed and necropsied at day 7 postinfection. The results represent values recorded separately for each of the 10 mice. The horizontal bars represent the geometric means. *P* values of less than 0.05 were considered to be significant.

Because of the higher virulence phenotype of the $\Delta ef0408-0409$ mutant observed in *G. mellonella* and its better colonization than the wild type in the urinary tract, these strains were also tested in a systemic infection model. We observed that the tissue burden in the liver and kidneys provoked by $\Delta ef0408-0409$ mutant cells was significantly increased compared to the wild type (see Fig. S2 in the supplemental material).

Phagocytes constitute an important part of the innate immunity against pathogens (72). Thus, we were interested in studying the survival of the sRNA mutant strains in an *in vivo-in vitro* macrophage infection model. As shown in Fig. 3, 3 of the 4 sRNA mutants tested ($\Delta ef0408-0409$, $\Delta ef0605-0606$, and $\Delta ef1368-1369$) survived better within murine peritoneal macrophages than the wild-type or complemented strains over a 72-h period. Of note, no difference was observed between the cell counts in the first hours postinfection, suggesting that phagocytosis levels of the wild-type, mutant, and corresponding complemented strains were similar. No significant difference in survival was observed in the case of the $\Delta ef3314-3315$ mutant (data not shown).

A link between sRNA and virulence has also been observed for other Gram-positive pathogens, like *S. aureus*, *Streptococcus*, and *Listeria*. For example, the RNAlII of *S. aureus* activates the translation of *hla*, encoding hemolysin, and represses the synthesis of major cell surface virulence factors like coagulase and protein A (73–75). In *Listeria monocytogenes*, the *lhrA* sRNA regulates the expression of chitinases ChiA and ChiB, which contribute to pathogenesis (76, 77). This is also the case for *fasX*, *pel*, and *rivX* of streptococci, which affect the expression of several virulence factors (78–80). Taken together, our results show for the first time

that candidate sRNAs play important roles in the virulence of *E. faecalis*.

The role of sRNAs in stress response. To study the functional role of the candidate sRNAs implicated in the virulence process, we selected conditions that might be relevant in the gastrointestinal tract or during the infection process. For that purpose, the wild-type, mutant, and complemented strains were cultured under oxidative-, osmotic-, or detergent stress conditions, as well as in serum (Table 2; see Fig. S3 to S6 in the supplemental material). In the case of acid stress, survival was monitored at a killing pH of 2.8. Under standard growth conditions (in M17glu at 37°C), the growth rates of the sRNA mutants and the parental strain were similar (Table 2). Under all conditions, the phenotypes of the complemented strains were very similar to those of the wild type. This proved that the behavior of the mutants was due only to the deletion of the respective sRNA and not to an additional mutation.

The growth of the $\Delta ef0408-0409$ strain in serum was comparable to that of the parental strain (Table 2). However, the mutant grew significantly better under different stress conditions (Table 2; see Fig. S3 in the supplemental material). In the case of oxidative stress, the growth rate of the mutant ($\mu_{\Delta} = 0.51$) was significantly higher in the presence of 1.5 mM H_2O_2 than that of the wild type ($\mu_{wt} = 0.17$) or the complemented strain ($\mu_c = 0.21$). It has recently been shown for *E. faecalis* that aerobic growth with glycerol as the energy source leads to the intracellular accumulation of H_2O_2 (81). Under these conditions, the wild-type and mutant strains showed similar growth rates during the first 6 h of growth, but the mutant reached a higher final OD₆₀₀ after 24 h of incubation (0.344 and 0.867, respectively) (see Fig. S3 in the supplemental material). The mutant also showed an increased growth rate in the presence of 0.08% bile salts ($\mu_{\Delta} = 0.41$, $\mu_{wt} = 0.24$, and $\mu_c = 0.21$) or 8% NaCl ($\mu_{\Delta} = 0.38$, $\mu_{wt} = 0.2$, and $\mu_c = 0.22$) (Table 2; see Fig. S3 in the supplemental material). Finally, as shown in Fig. 4A, the $ef0408-0409$ sRNA mutant was significantly more resistant to acidity (M17, pH 2.8) than the wild-type strain (from 13-fold at 30 min to 23-fold at 120 min after the challenge). The $ef0408-0409$ deletion mutant showed a strong effect in the insect and mouse models, where it appeared more virulent and more able to colonize organs and survived better inside macrophages. These observations might be correlated with the ability of the mutant to better cope with stresses, especially oxidative stress. This showed that the chromosomal TA system was not cryptic and was involved in the virulence of *E. faecalis*, likely because of its ability to affect the cellular stress response, as well as its impact on the expression of metabolic enzymes (see below). It is tempting, then, to speculate that this system, likely from a plasmid, has been selected to promote colonization rather than virulence.

It has recently been demonstrated that several sRNA deletion mutants of the haloarchaeon *Haloferax volcanii* showed a gain-of-function phenotype (82). Taken together, it may be suggested that the biological role of some sRNAs can be to stabilize the homeostasis of cells, especially when they are confronted with the multiple environmental changes that occur in real life.

The $\Delta ef0605-0606$ mutant was exposed to the same stresses, and the results are summarized in Table 2. These cells grew better in the presence of 1.5 mM H_2O_2 ($\mu_{\Delta} = 0.37$, $\mu_{wt} = 0.18$, and $\mu_c = 0.19$) or during growth with glycerol than the wild-type strain (see Fig. S4 in the supplemental material). Under the latter condition, the mutant strain reached a much higher final OD₆₀₀ (4.2) than the parental strain (0.344) (see Fig. S4 in the supplemental mate-

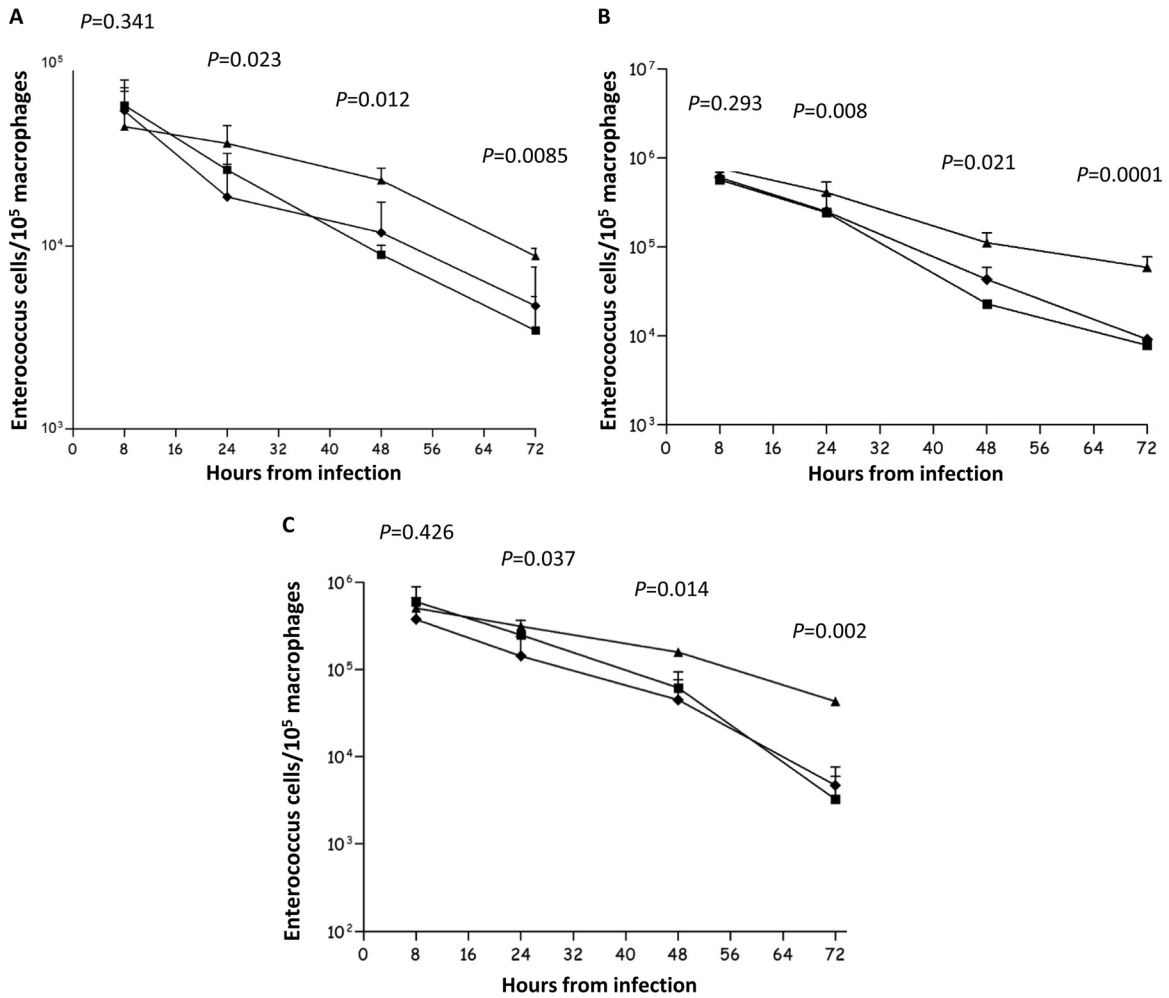


FIG 3 Time course of intracellular macrophage survival of the $\Delta ef0408-0409$ (A), $\Delta ef0605-0606$ (B), and $\Delta ef1368-1369$ (C) mutant strains. Diamonds, *E. faecalis* EryS; triangles, sRNA mutants; squares, complemented strains. The results represent the mean numbers and standard deviations of viable intracellular bacteria per 10^5 macrophages from three independent experiments.

rial). The $\Delta ef0605-0606$ mutant also grew somewhat better in the presence of 0.08% bile salts (see Fig. S4 in the supplemental material), which is mostly due to a reduced lag phase compared to the wild-type and complemented strains. Lastly, no difference compared to the wild-type strain has been observed under acid and osmotic stress or in serum (see Fig. S4 in the supplemental mate-

rial). The *ef0605-0606* sRNA is present in only 9 strains of the 54 *E. faecalis* genomes analyzed, due to its localization in the PAI (35). It has been recently found, using 5' RACE, that it is also induced under aerobic growth conditions (33, 35). The corresponding mutant was less virulent in the *Galleria* model but survived better inside macrophages, which may be correlated with its greater fit-

TABLE 2 Growth phenotypes and promoter activities under different stress conditions

Growth conditions	$\Delta ef0408-0409$ strain		$\Delta ef0605-0606$ strain		$\Delta ef1368-1369$ strain		$\Delta ef3314-3315$ strain	
	Growth effect ^a	Promoter activity ^b	Growth effect	Promoter activity	Growth effect	Promoter activity	Growth effect	Promoter activity
M17glu	ND ^c	ND	ND	ND	ND	ND	ND	ND
1.5 mM H ₂ O ₂	+	-13.0	+	-3.0	ND	ND	ND	ND
cc17 MOPS glycerol	+	-2.0	+	-8.0	ND	ND	ND	ND
8% NaCl	+	-1.3	ND	ND	-	ND	ND	ND
0.08% bile salts	+	-3.0	+	-3.0	ND	ND	+	-2.0
100% horse serum	ND	ND	ND	ND	+	ND	ND	ND

^a +, higher, and -, lower growth rates of the sRNA mutant than of the wild-type strain. All the values are available in Fig. S2 to S5 in the supplemental material.

^b Ratio of the promoter activities (in Miller units) of the pVEPhoZPsRNAs strains (where *phoZ* transcription is under the control of sRNA promoters) at T90 (90 min in the presence of the stress) to the measure at T0 (just before the stress). All the values are available in Fig. S7 in the supplemental material.

^c ND, no difference.

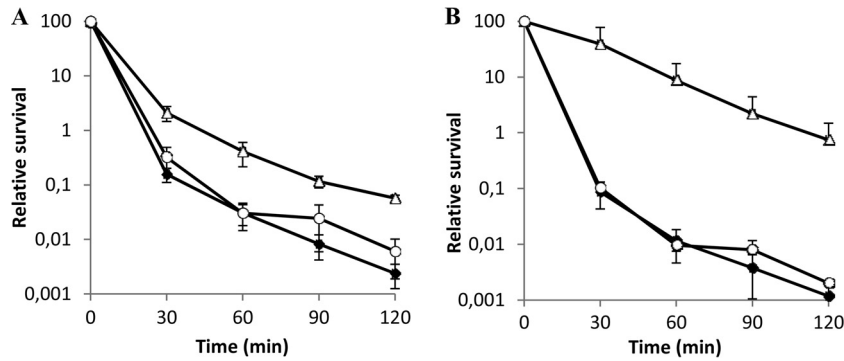


FIG 4 Survival under acid stress conditions (M17/7 medium adjusted to pH 2.8) of the sRNA mutant strains. The *E. faecalis* wild type, the $\Delta ef0408-0409$ mutant (A) or the $\Delta ef1368-1369$ mutant (B), and the complemented strains are represented by diamonds, triangles, and circles, respectively. At the indicated times, the viable-cell count was determined by plating dilutions onto M17glu; 100% survival corresponds to the viable-cell count determined just prior to exposure to the indicated stress. For all assays, each data point represents the mean \pm standard error (SE) of 3 independent experiments.

ness in the presence of H_2O_2 . The decreased virulence model of the mutant in the worm could be due to greater sensitivity to antimicrobial molecules present in the hemolymph of the larvae.

The $\Delta ef1368-1369$ mutant also showed interesting phenotypes under stress conditions (Table 2). It had a somewhat reduced growth rate in the presence of 8% NaCl ($\mu_{\Delta} = 0.16$ and $\mu_{wt} = 0.28$) (see Fig. S5 in the supplemental material) but had remarkably better survival (more than 2 orders of magnitude) in an acid environment (pH 2.8) than the wild-type and the complemented strains (Fig. 4B). Its better ability to cope with this stress may be due to its better survival inside macrophages and ability to colonize mouse organs. Furthermore, the strain lacking *ef1368-1369* sRNA grew to a higher final OD_{600} than the wild type ($OD_{600} = 0.5$ and 0.24, respectively) when cultured in horse serum (see Fig. S5 in the supplemental material). Finally, the sRNA mutant lacking *ef3314-3315* showed a reduced lag phase when grown in the presence of 0.08% bile salts compared to the wild type and the complemented strain (Table 2; see Fig. S6 in the supplemental material). However, no difference in growth between the wild type and the mutant was observed with the other stresses used in this study (data not shown).

Of note, the sensitivities of the wild-type strain and all the sRNA mutants to cold ($8^{\circ}C$) and heat ($62^{\circ}C$) shock, autolysin treatment, and exposure to antibiotics were tested, but no difference was observed (data not shown). In addition, the profiles of carbohydrate fermentation (using bioMérieux API test kits [Craponne, France] for bacterial identification of 50 vials of dehydrated substrates) were the same for all strains.

sRNA promoter activities under different stress conditions.

Transcription of sRNA is usually affected by specific environmental conditions (83). Therefore, we were interested in analyzing the promoter activities of the candidate sRNAs. The corresponding promoter regions were fused to the AP gene of *E. faecalis*, and AP activities, expressed in MU, were then determined under those conditions in which the sRNA mutants showed the most pronounced phenotypes.

Without a promoter fused to AP, only low activity of the enzyme (from 4 to 10 MU) was detected regardless of the growth condition analyzed (data not shown). The promoter activities of *Pef0408-0409* and *Pef0605-0606* were comparable under nonstress growth conditions (between 193 and 225 MU, respectively). However, both promoters showed 3-fold-decreased activity in the

presence of bile salts, 3-fold (*Pef0605-0606*) and 13-fold (*Pef0408-0409*) decreases in the presence of 1.5 mM H_2O_2 , and 2-fold (*Pef0408-0409*) and 8-fold (*Pef0605-0606*) reductions in expression during growth on glycerol in comparison to nonstress growth conditions (Table 2; see Fig. S7 in the supplemental material). The activity of the *ef0408-0409* promoter also showed a slight decrease after 90 min under NaCl conditions (1.3-fold). Furthermore, 2-fold reduction in comparison to the expression under nonstress growth conditions (166 versus 122 MU) was measured for *Pef3314-3315* in the presence of bile salts (Table 2; see Fig. S7 in the supplemental material). As previously mentioned, analysis of the different mutant strains revealed that the lack of the transcripts led to better growth under stress conditions. Thus, in the wild-type strain, it seems appropriate to observe that the stressing agents provoked reduced activities of the corresponding sRNA promoters, allowing cells to cope better with these challenges.

Finally, although the $\Delta ef1368-1369$ mutant strain showed a slightly reduced growth rate under osmotic stress, no change in the AP activity of the *Pef1368-1369* construct was found under these conditions (Table 2; see Fig. S7 in the supplemental material). In serum, the growth rates for the mutant and the wild type were similar, and no modification in the activity of *Pef1368-1369* has been detected (Table 2; see Fig. S7 in the supplemental material). However, the difference observed in the final OD_{600} under serum growth conditions suggests that *ef1368-1369* sRNA could mainly act during stationary phase.

Identification of the sRNA regulon. Using quantitative real-time PCR, we checked the effects on the transcription of the flanking genes of the deleted candidate sRNAs. For the two mutant strains $\Delta ef0408-0409$ and $\Delta ef3314-3315$, no relevant impact on the transcription of neighboring genes was observed (Table 3). On the other hand, the *ef0605* (encoding a hypothetical protein) and *ef0606* (encoding a Dps protein) genes were, respectively, 4.4- and 2.7-fold overexpressed in the $\Delta ef0605-0606$ mutant strain (Table 3). The increased transcription of the *ef0606* gene, encoding an enzyme involved in the protection of DNA against oxidative stress, in the sRNA mutant appears consistent with its phenotype toward H_2O_2 (84). Dps is a DNA-binding protein that protects the genome against oxidative damage and also acts as an iron chelator, reducing its free cellular concentration and consequently the formation of reactive oxygen species (ROS). Indeed, Fe^{2+} ions are involved in the Fenton reaction that converts H_2O_2 into OH^{\cdot} ,

TABLE 3 Transcriptional modifications of genes neighboring the sRNA deletions

Neighboring sRNA gene	Function	Fold change in expression ^a (<i>P</i> < 0.01)
sRNA <i>ef0408-0409</i>		
<i>ef0407</i>	Transcriptional regulator	ND
<i>ef0408</i>	Phosphotransferase (PTS) system, IIA component	ND
<i>ef0409</i>	Hypothetical protein	ND
<i>ef0411</i>	PTS system, mannitol-specific IIBC component	ND
sRNA <i>ef0605-0606</i>		
<i>ef0604</i>	Gls24 protein	ND
<i>ef0605</i>	Conserved hypothetical protein	+4.4
<i>ef0606</i>	Dps protein family	+2.7
sRNA <i>ef1368-1369</i>		
<i>ef1368</i>	Conserved hypothetical protein	ND
<i>ef1369</i>	Transcriptional regulator, Cro/CI family	-113.8
<i>ef1370</i>	Drug resistance transporter, EmrB/QacA family protein	-3.3
sRNA <i>ef3314-3315</i>		
<i>ef3311</i>	Glucose-inhibited division protein A	ND
<i>ef3312</i>	tRNA modification GTPase TrmE	ND
<i>ef3313</i>	Hypothetical protein	ND
<i>ef3314</i>	Cell wall surface anchor family protein	ND
<i>ef3315</i>	CitG family protein	ND

^a The expression values are the averages of at least 3 different RNA extractions from logarithmic-growth-phase cells. All the gene expression values were normalized to the level of *gyrA* and 23S RNA, and those that differed by more than 2-fold with *P* values of <0.01 were considered significant. ND, no difference.

which is a highly deleterious ROS. These results showed that this sRNA is implicated in the regulation of the expression of the adjacent genes that correspond to the first member of the *ef0605-0606* sRNA regulon and that *ef0605-0606* may act as a repressor.

Since Δ *ef1368-1369* also corresponds to the deletion of *ef1368*, as expected, no expression of the latter gene was observed. However, *ef1370* (encoding an EmrB drug resistance transporter) appeared to be 3.3-fold repressed in the mutant strain (Table 3). Also, the Δ *ef1368-1369* mutant is not more sensitive to all the antibiotics tested than the wild-type strain (data not shown).

In general, sRNAs act at the posttranscriptional level of regulation by interacting with mRNA (3, 85). Therefore, in order to identify members of sRNA regulons in *E. faecalis*, i.e., targets whose expression is directly or indirectly controlled by a given sRNA, a global proteomics approach was used to compare the profiles of the sRNA mutant and the parental strains. Using two antibodies (anti-GroEL and anti-Ers), we performed Western blot experiments and confirmed that some proteins were deregulated according to the mutant strain. Thus, GroEL was induced in the Δ *ef0408-409* mutant and repressed in the mutant Δ *ef3314-3315*, and Ers was induced in the Δ *ef0408-409* and Δ *ef0605-0606* mutant strains (see Fig. S8 in the supplemental material).

The proteomics experiments revealed that, in general, induced proteins were twice as abundant as repressed proteins in the sRNA mutant strains (112 induced targets versus 61 repressed [at least 2-fold-change]). In the Δ *ef0408-0409*, Δ *ef0605-0606*, Δ *ef1368-1369*, and Δ *ef3314-3315* strains, we observed, respectively, 1.7-, 2-, 3-, and 1.5-fold more induced proteins than repressed polypeptides (see Table S2 in the supplemental material). The putative direct or indirect targets of the candidate sRNAs studied seemed to belong to a large number of cellular-role categories, including cellular process, energy metabolism, regulatory functions, and transcription (Table 4). Of note, the regulation of metabolic path-

ways (i.e., entry into stationary phase, energy production, and biofilm formation) is necessary for bacterial adaptation to adverse conditions and for pathogenicity. Thus, the fact that several genes encoding enzymes linked to different metabolic pathways were members of the candidate sRNA regulons illustrates their roles in the fitness of bacterial cells, which can be correlated with a role in colonization and/or virulence.

Some of the identified polypeptides, such as enzymes known to be involved in the oxidative-stress response and/or virulence that may, at least in part, explain the phenotypes of the mutant strains, are of particular interest. This may be true for proteins encoded by the *arc* operon, overproduced around 2-fold in the Δ *ef0408-0409* mutant and more than 4-fold in the Δ *ef0605-0606* mutant strain. Proteins encoded by the *arc* operon are enzymes of the arginine deiminase pathway but may also be involved in the oxidative-stress response. In *Lactococcus lactis*, the disruption of the *arcA* or *arcB* gene abolished H₂O₂ resistance (86). Other proteins linked to oxidative stress were observed. Indeed, the transcriptional regulator Ers, known to play roles in virulence, H₂O₂ resistance, and glycerol metabolism, has been found to be induced more than 3-fold and 4-fold in the Δ *ef0408-0409* and Δ *ef0605-0606* mutant strains, respectively (24, 53). Interestingly, in *E. faecalis*, Ers has been shown to participate in the regulation of the *arc* operon (53). It should be noted that *ers* (*ef0074*) and *ef0605-0606* sRNA are both located near an operon structure that includes a *gls24* paralogue encoding a general stress protein and virulence factor in *E. faecalis* (87–89).

In addition, the NADH peroxidase enzyme (Npr) was overexpressed 2-fold in the Δ *ef0408-0409* mutant strain. Npr is implicated in the oxidative-stress response in *E. faecalis* and has been shown to be the most important peroxidase for protecting cells against external or internal H₂O₂ stress (39, 90, 91).

Expression of some polypeptides belonging to the category

TABLE 4 Cellular role categories of putative targets up- and downregulated on the sRNA mutant strains

Cellular role category	No. of targets ^a											
	<i>ef0408-0409</i>			<i>ef0605-0606</i>			<i>ef1368-1369</i>			<i>ef3314-3315</i>		
	T	R	I	T	R	I	T	R	I	T	R	I
Amino acid biosynthesis	3	0	3	2		2						
Biosynthesis of cofactors, prosthetic groups, and carriers	1	0	1	2		2						
Cell envelope	6	3	3	2	1	1	1	0	1			
Cellular processes	5	2	3	4	2	2	1	1	0	3	2	1
Central intermediary metabolism	1	0	1				1	1	0	1	1	0
DNA metabolism	3	3	0							2	0	2
Energy metabolism	10	3	7	4	0	4	5	1	4	5	2	3
Fatty acid and phospholipid metabolism	1	0	1	2	0	2	2	0	2	1	0	1
Hypothetical proteins	4	0	4	3	0	3						
Mobile- and extrachromosomal-element functions	1	0	1	1	0	1	2	0	2	1	0	1
Protein fate	5	2	3	3	1	2	1	0	1	2	1	1
Protein synthesis	7	4	3	6	4	2	2	1	1	4	2	2
Purines, pyrimidines, nucleosides, and nucleotides	8	4	4	10	6	4	2	1	1	1	1	0
Regulatory functions	5	2	3	3	0	3				1	1	0
Transcription	3	1	2	3	2	1	1	0	1	2	0	2
Transport and binding proteins	1	1	0									
Unknown function	11	3	8	8	2	6	2	0	2	2	0	2
Total	75	28	47	53	18	35	20	5	15	25	10	15

^a T, total; R, repressed; I, induced.

“cellular processes,” likely related to pathogenicity, seemed to be controlled by sRNAs. For example, hemolysin A (EF0982) was overproduced around 3-fold in the *Δef0408-0409* and *Δef0605-0606* mutant strains; the EF2150 protein, a member of the FemAB family, is induced around 5-fold in the two strains; and the superoxide dismutase SodA is induced 2-fold in the *Δef3314-3315* mutant (see Table S2 in the supplemental material).

ef1368-1369 sRNA likely acts as an antisense sRNA. However, the expression of 20 proteins was modified in the corresponding mutant strain, showing that the sRNA may have other targets than *ef1369* mRNA. We did not find EF1369 in our proteomics data, likely because of its small size (around 13 kDa). Indeed, the 2D gel electrophoresis conditions we used allowed a clear separation of proteins from 15 to 140 kDa.

Computational predictions for mRNA targets of candidate sRNAs were performed using two software programs (Predator and Copra) (54, 55). For each candidate sRNA, around 100 putative targets were found. Among them, only a few were members of the regulons identified by proteomics (Table 5). Of note, as expected, *ef0409*, homologous to RAI of the TA system, could be a target of *ef0408-0409* sRNA. Bioinformatics approaches to predict the cellular targets of sRNAs are still plagued by false results, because all the parameters involved in the interaction are not known (2). This may explain the low coverage with genes encoding proteins deregulated in the sRNA mutants, as well as the different data obtained with the two informatics tools tested. Moreover, the members of the regulons identified by proteomics may result from a cascade of regulation events. Nevertheless, further study combining all the approaches and experimental data will provide insight into how these sRNAs function.

Conclusions. These results constitute the first functional analysis of candidate sRNA molecules in enterococci. Some sRNA mutant strains were less virulent, whereas another was more virulent. Commensal bacterial cells, such as *E. faecalis*, need to find equilibrium between behavior favorable to colonization (repressing

virulence) and pathogenicity according to the host environment. Therefore, it is not surprising to observe that the transcription of certain sRNAs was modulated by stresses likely encountered in the different bacterial niches or during infection and that mutants were affected in their growth or survival under stress conditions compared to the wild-type strain. The proteomics approach allowed the identification of some members of the candidate sRNA regulons without distinction between direct and indirect interaction. Nevertheless, these results showed that for *E. faecalis*, an opportunistic pathogen, the implication of sRNAs in virulence appears to be mainly correlated with global metabolism and the stress response. The fact that the expression of some enzymes was altered in different mutant backgrounds underscores the cross talk that may exist between regulatory networks orchestrated by

TABLE 5 Computational identification of putative cellular targets of sRNAs that were member of the candidate sRNA regulons obtained by a proteomics approach

Candidate sRNA	Putative target			
	Predator		Copra	
	Gene	Function	Gene	function
<i>ef0408-0409</i>	<i>ef0409^a</i>	Hypothetical protein	<i>ef0409^a</i>	Hypothetical protein
	<i>ef3303^b</i>	Myosin cross-reactive antigen		
<i>ef0605-0606</i>	<i>ef2353^b</i>	Acetyltransferase		
	<i>ef0982^b</i>	Hemolysin A		
	<i>ef0684^a</i>	<i>cmp</i> binding protein; putative		
<i>ef1368-1369</i>			<i>ef0185^b</i>	Phosphopentomutase
<i>ef3314-3315</i>			<i>ef2353^b</i>	Acetyltransferase
			<i>ef1613^a</i>	Formate acetyltransferase

^a CopraRNA and RNA Predator results with *P* values lower than 0.01.

^b CopraRNA and RNA Predator results with *P* values lower than 0.05.

sRNAs. Because sRNAs are “cost-effective” molecules for cell adaptation to environmental changes, they appear here as key actors for the transition from a commensal relationship with the host to a virulence trait.

The selected candidate sRNAs for this study are among the most highly expressed during the exponential and/or stationary growth phase but correspond to a small fraction of sRNAs present in bacteria (more than 70) (35). We are thus at the first step of the identification of the roles of the sRNAs, which constitutes one of the main future challenges for a better understanding of genetic regulation.

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