Characterization of sal(A), a Novel Gene Responsible for Lincosamide and Streptogramin A Resistance in Staphylococcus sciuri

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Natural resistance to lincosamides and streptogramins A (LSA), which is a species characteristic of Bacillus subtilis and Enterococcus faecalis, has never been documented in the Staphylococcus genus. We investigate here the molecular basis of the LSA phenotype exhibited by seven reference strains of Staphylococcus sciuri, including the type strains of the three described subspecies. By whole-genome sequencing of strain ATCC 29059, we identified a candidate gene that encodes an ATP-binding cassette protein similar to the Lsa and VmlR resistance determinants. Isolation and reverse transcription-quantitative PCR (qRT-PCR) expression studies confirmed that Sal(A) can confer a moderate resistance to lincosamides (8 times the MIC of lincomycin) and a high-level resistance to streptogramins A (64 times the MIC of pristinamycin II). The chromosomal location of sal(A) between two housekeeping genes of the staphylococcal core genome supports the gene’s ancient origins and thus innate resistance to these antimicrobials within S. sciuri subspecies.

M acrolides, lincosamides, and streptogramins (MLS) constitute the group of antibiotics employed to treat staphylococcal and streptococcal infections in human and veterinary medicine. Resistance to these antibiotics is increasing among clinical isolates, mainly as a result of acquisition of exogenous genes carried by mobile elements, plasmids, and/or transposons. The validated list of MLS resistance (MLS-R) determinants includes rRNA methylases, efflux pumps, and inactivating enzymes (1).

Efflux-mediated resistance to MLS antibiotics in staphylococci relies on the ATPase activity of a very special kind of ATP-binding cassette (ABC) protein (2–4). These MLS-R ABC proteins form a robust cluster that corresponds to the antibiotic resistance (ARE) protein subfamily, closely related to, but distinct from, the regulatory (REG) subfamily in the phylogenetic and functional classification of ABC systems (5). Their protein designations depend on both the resistance phenotype and the bacterial origin of the genes. Two resistance spectra have been reported in the literature for ABC-based MLS efflux among Gram-positive cocci; the first one is encoded by msr genes and leads to macrolide and sometimes streptogramin B (M/MSB) resistance, while the other is encoded by lsa and vga genes and leads to resistance to lincosamides and streptogramin A (LSA). Thus far, three classes of msr, five classes of lsa, and five classes of vga genes have been shown to be clinically relevant because of their detection in pathogens such as Staphylococcus, Streptococcus, and Enterococcus strains (1, 6, 7).

One of these genes is intrinsic to Enterococcus faecalis; this gene is lsa(A), whose integrity and constitutive expression have been linked to the LSA phenotype, useful as a species identification marker in enterococci (8).

Staphylococci can acquire the LSA phenotype via mobile genetic elements. Though lsa(B)- and lsa(E)-carrying plasmids have been described, vga genes, especially vga(A), carried by plasmids and/or transposons, have been found predominantly among clinical strains isolated in human medicine (9). Our most recent study failed to identify the resistance gene responsible for the LSA phenotype within a Staphylococcus sciuri isolate (10). We report here the nucleotide sequence, genome localization, and antibiotic-inducible expression of sal(A), a new category of ARE-encoding genes that we found in all our reference strains belonging to the three subspecies described for the S. sciuri species.

Materials and Methods

Bacterial strains and growth conditions. Seven reference strains of S. sciuri were included in this study. They were previously assigned by ribotype delineation to the three validly described subspecies: S. sciuri subsp. sciuri, S. sciuri subsp. carnaticus, and S. sciuri subsp. rodentium (12, 13). Four strains were listed by the World Federation of Culture Collections: ATCC 29059 (CIP 105824) and type strain ATCC 29062 (CIP 81.62T), which belong to S. sciuri subsp. sciuri, type strain ATCC 700058 (CIP 105826T) of S. sciuri subsp. carnaticus, and type strain ATCC 700061 (CIP 105829T) of S. sciuri subsp. rodentium. The three other strains, SS226, BL2, and SVv1, originally described by Wesley Kloos and which belong to S. sciuri subsp. sciuri, S. sciuri subsp. carnaticus, and S. sciuri subsp. rodentium, respectively, have since obtained the designations BM12487, BM12489, and BM12494, respectively. Escherichia coli strain Top10 was used as a cloning host for all plasmid constructs, and Staphylococcus aureus strain RN4220 was used as a recipient for plasmid transformation. Staphylococcus saprophyticus strain ATCC 15305 (CIP 76.125T) was used as a control. When required, ampicillin at 100 μg/ml or chloramphenicol at 10 μg/ml was added to the culture media (Becton, Dickinson, Le Pont de Claire, France) for plasmid maintenance in E. coli or S. aureus, respectively. These antibiotic compounds, as well as lincomycin (Lc), were purchased from Sigma (Saint-Quentin Fallavier, France), while the two others, rifampin and pristinamycin IIA (PIIA), were kindly pro-
vided by Sanofi-Aventis (Croix-de-Berry, Antony, France). Bacterial strains were grown at 37°C in Luria-Bertani medium for *E. coli* or in brain heart infusion for *staphylococci*. Broth cultures were vigorously shaken, and cell density was monitored by measuring the absorbance at 600 nm.

**Antimicrobial susceptibility assays.** MLS resistance patterns of bacteria were determined on Mueller-Hinton agar using antibiotic disks purchased from Bio-Rad (Marnes-la-Coquette, France) or loaded manually with 20 µg PI. MICs of *Lc* and *PIA* were obtained with serial 1:2 dilutions of the antibiotics in Mueller-Hinton agar using 10⁴ CFU per spot (2). Each MIC determination was carried out in triplicate from two independent series of experiments, with plates incubated for 18 h at 37°C. Reference breakpoints were those published in the Communiqué CASFM, which are annually revised and updated (see [http://www.sfm-microbiologie.org](http://www.sfm-microbiologie.org) for details). The control strain was *S. aureus* RN4220.

**DNA manipulations.** Genomic DNA of *S. sciuri ATCC 29059* was extracted and purified by using the QIAamp DNA minikit (Qiagen GmbH, Hilden, Germany) with lyostaphin (Sigma) added to the lysis buffer. Plasmid DNA was isolated from *E. coli* using the QIAprep spin system (Qiagen). Gene cloning in pRB474 (14) was achieved in successive steps, starting from a blunt-ended DNA fragment obtained by PCR and then purified by the QIAquick PCR purification kit (Qiagen) before being inserted into the pCR4-Blunt Topo vector (Invitrogen, Carlsbad, CA). Bacterial DNA was amplified using PfuUltra II fusion high-fidelity DNA polymerase (Agilent Technologies, Santa Clara, CA) from boiled colonies resuspended in sterile water, of which 5 µl was incorporated into 50-µl reaction mixtures. PCR was performed in an Eppendorf vapo.protect Mastercycler pro system using primer pairs described in Table 1 and classic thermal cycling parameters, such as 1 min at 52°C for the annealing segment and 2 min at 72°C for the elongation segment. The sizes of the PCR products were estimated by ethidium bromide staining after 1% agarose gel electrophoresis using a 1-kb DNA ladder (Fermentas, Vilnius, Lithuania) as a double-stranded DNA standard. The clean PCR products were either directly sequenced or cloned into pCR4 for sequencing, and the PstI-EcoRI DNA fragment was later subcloned into a pRB474 shuttle vector (14) using the instructions, reagents, and enzymes contained in a rapid DNA dephosphorylation and ligation kit (Roche Applied Science, Indianapolis, IN). The resulting recombinant plasmid and the empty shuttle vector were introduced into *S. aureus* by electroporation as previously described (2). Each of the two categories of transformants was analyzed by antimicrobial susceptibility assays, as described above.

**RMAs.** The purified genomic DNA of *S. sciuri ATCC 29059* was amplified, fragmented, and labeled using the GeneChip resequencing assay kit (Affymetrix Inc., Santa Clara, CA). The labeled sample was loaded onto a preheated PathogenID v.1 microarray cartridge (Affymetrix) containing 250 antibiotic resistance genes (15). Table S1 in the supplemental material lists the nature of the genes tiled on the microarray. MLS-R genes represent 26% of the total gene number (66 of the 250 nonredundant targeted sequences). After an overnight hybridization at 45°C, the GeneChip was washed, stained, and scanned according to Affymetrix’s instructions. The raw image file (.DAT) obtained after the scan was transformed into a .CEL file. Nucleotide sequences were delivered in FASTA format by GeneChip sequence analysis software (GSEQ v4.1), which uses a derivative of the ABACUS base-calling algorithm (16). This algorithm consists of an automated statistical method that analyzes raw microarray assay (RNA) hybridization data and optimizes the base-calling process. For each base call, a quality score is defined. If this score is below a chosen threshold, an undetermined base, N, rather than A, T, C, or G is displayed. Sequence information is then analyzed manually by using the continuous or discontinuous stretch of resolved bases as a query for database similarity searches.

**WGS.** High-throughput whole-genome sequencing (WGS) of the genomic DNA of *S. sciuri ATCC 29059* was performed using Roche 454 GS Titanium chemistry (GATC Biotech, Constance, Germany). The 83,000 reads were assembled by CLC Workbench 5.2 software, which generated 371 contigs. The nucleotide and deduced protein sequences were analyzed with the tBLASTn and BLASTp programs available at the Institut Pasteur Biology IT Center through the Mobyle platform (17). Search for a candidate *S. sciuri* ARE protein(s) was conducted using Vga(A) and VmlR as queries (GenBank accession numbers AF171259 and BSU05610, respectively). The start codon of the presumptive gene and its putative Shine-Dalgarno sequence were determined by visual inspection.

**Multiple-sequence alignments.** Nucleotide and amino acid polymorphisms were analyzed using the Clustal X v2.1 package (18). Comparison of a set of 25 representative ARE proteins with a confirmed or likely role in MLS resistance was performed using the Muscle algorithm (19). Phylogenetic analysis was done using PhyML 3.0 software, and the resulting tree was displayed with FigTree.

**Induction tests and qRT-PCR.** The impact of 3-h sub-MIC treatments on resistance properties of *S. sciuri ATCC 29059* was tested twice in broth before plating. Antibiotic was added at an optical density at 600 nm (OD₆₀₀) of 0.1. Total RNA of bacterial cells was extracted at an OD₂₆₀ of 0.6 with TRIzol reagent (Invitrogen) as previously described (20) and then treated with the Turbo DNA-free kit (Invitrogen) to remove any trace of DNA contamination. Quantitation of target gene expression was performed in relation to 16S rRNA expression using reagents of the LightCycler RNA amplification kit SYBR green I (Roche) according to the manufacturer’s instructions. The amount of nucleic acid material was calculated by measuring the absorbance at 260 nm of the RNA samples. Each of the two independent series of experiments was analyzed in quadruplicate using 50 ng or 1 ng of input RNA and a 0.5 µM concentration of gene-specific primers (Table 1). Reverse transcription was carried out at 55°C for 10 min. PCR mixtures were subjected to 1 cycle at 95°C for 30 s and 45 cycles at 95°C for 5 s, 54°C for 10 s, and 72°C for 20 s. Identification of the PCR products was carried out by DNA melting profiling; profiles were automatically converted into melting peaks by Light Cycler instrument software (Roche).

**Nucleotide sequence accession number.** The nucleotide sequence of *sal(A)* found within contig 252 of the CLC workbench assembly of the

<table>
<thead>
<tr>
<th>Table 1 List of primers used for this study</th>
<th>Target</th>
<th>Primer pair</th>
<th>Primer sequence from 5’ to 3’</th>
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* Nucleotides in boldface correspond to the PstI site used for *sal(A)* gene cloning into pRB474.
WGS of *S. sciuri* strain ATCC 29059 has been deposited into the GenBank database under accession no. KC693025.

**RESULTS**

**LS$_A$ resistance is intrinsic to *S. sciuri* species.** Antimicrobial susceptibility testing was performed on the seven *S. sciuri* strains using the disk diffusion technique. *S. saprophyticus* strain ATCC 15305 was MLS susceptible, like the control strain RN4220. All *S. sciuri* strains exhibited an unambiguous LS$_A$ phenotype, with growth up to the PIIA-containing disk and a zone inhibition diameter around the Lc disk ranging from 12 to 16 mm. Thus, they were all categorized as resistant to both antibiotics. They all showed intermediate susceptibility to clindamycin, with diameters of the inhibition zones ranging from 19 and 21 mm, and full susceptibility to pristinamycin, which is comprised of a mixture of 70% S$_A$ and 30% S$_B$ compounds. The MICs of Lc and PIIA for strain ATCC 29059 were 16 μg/ml and 32 μg/ml, respectively. Precultivation of the strain for 3 h at one-eighth of the MIC of Lc led to a 2-fold increase in both values. This strain was chosen for an in-depth genetic study in order to characterize the molecular basis of LS$_A$ resistance in *S. sciuri*.

**No LS$_A$ resistance gene was identified by RMA.** A total of 250 resistance genes were selected from the GenBank database to be represented in the PathogenID v.1 microarray (see Table S1 in the supplemental material). A set of 28 sequences coding for ARE proteins which can be either intrinsic or acquired among Gram-positive bacteria were tiled on the array. Only the sequence that codes for VmlR yielded a short fingerprint of 30 nucleotides with the labeled and fragmented DNA of strain ATCC 29059. When translated, this region of homology provided the peptide signature of the conserved H-loop motif shared by some categories of ABC proteins, including those of the ARE subfamily. Thus, neither the length of the deduced nucleotide sequence (Fig. S1) nor the discriminative power of the deduced peptide stretch (IVVSHDR) was strongly indicative of the presence of an ARE-encoding gene in the genome of *S. sciuri* ATCC 29059. Indeed, there are at least three ABC proteins of the REG subfamily coded by the staphylococcal core genome: SACOL2036 (homologous to the YdiF protein of *Bacillus subtilis*), SACOL1427 (homologous to YkpA of *B. subtilis*), and SACOL0779 (homologous to YfmR of *B. subtilis*). None of these three ubiquitous proteins have a demonstrated role in MLS resistance. Rather, due to their similarity to the *Saccharomyces cerevisiae* regulatory protein GCN20, they might be involved in global gene regulation networks (21). Only fine-tuned analysis of full-length polypeptides would distinguish ARE from REG members. The data sequences recovered here by RMA for ARE scanning were not large enough for such a precise characterization. This was in sharp contrast to those obtained for other gene targets, such as *mecA*, *rpoB*, *gyrA*, or *rrs*, with which the accuracy of the identification process was optimal (data not shown). It was therefore anticipated that either a shotgun cloning or a complete genome sequencing might decipher the nature of the LS$_A$ resistance mechanism in strain ATCC 29059 and, more broadly, in *S. sciuri* species.

**One candidate ARE gene was identified by WGS.** From *de novo* assembly of the entire genome of strain ATCC 29059, a tBLASTn search identified a putative ARE protein encoded by contig 252. The encoding gene had 1,626 bp and appeared to be inserted between two housekeeping genes, *iscS* and *mmmA*, of the staphylococcal core genome (Fig. 1). All features of a class 2 ABC protein were found in the deduced protein sequence (Fig. 2): two Walker A motifs (positions 35 to 43 and 347 to 355), two Walker B motifs (positions 131 to 136 and 431 to 436), and two H-loop switches (positions 185 and 485). The closest NCBI matches were ≈70% amino acid identity with two other hypothetical proteins of unassigned staphylococcal genomes recently released in databases: WP_016910816 of *Staphylococcus vitulinus* and WP_016998509 of *Staphylococcus lentus*. The highest scores of amino acid identity with referenced members of the ABCISSE database (http://www.pasteur.fr/abcisse) were 28% with VmlR, formerly ExpZ, 27% with Vga(A), and 24% with Lsa(A). Such high-scoring results with ARE proteins conferring an LS$_A$ phenotype strongly suggested that the candidate gene found within contig 252 was responsible for the LS$_A$ resistance properties observed in strain ATCC 29059 and possibly in the other tested *S. sciuri* strains. In contrast, none of the other assembled staphylococcal genomes at the same chromosomal location contained an ARE-encoding gene. Only the genome of *S. saprophyticus*, strain ATCC 15305, harbors a similar genetic structure at this locus (Fig. 1). However, the presence of a stop codon at the end of the first quarter of the presumptive resid-
tance gene would disrupt the full-length reading frame. Without such a mutation, the deduced polypeptide from this pseudogene would have 27% and 25% amino acid identity with Vga(A) and VmlR, respectively (Fig. 2). As mentioned above, strain ATCC 15305 is unambiguously susceptible to all MLS antibiotics.

Characterization of the novel \textit{sal}(A) gene. The putative ARE-encoding gene of strain ATCC 29059 was amplified using primers P7 and P8 (Table 1) and cloned into shuttle vector pRB474. The recombinant plasmid was electroporated into \textit{S. aureus} RN4220. In comparison with the recipient carrying the empty plasmid, \textit{S. aureus} RN4220 transformants carrying the \textit{S. sciuri} gene exhibited 8-fold- and 64-fold-increased MIC values of Lc (8 \(\mu\)g/ml) and PIIA (64 \(\mu\)g/ml), respectively. Such a result showed that the cloned gene was capable of conferring an LSA phenotype. In multiple-sequence alignment, the novel characterized ARE protein of \textit{S. sciuri} was confirmed to be similar to the VmlR determinant of \textit{B. subtilis}, and it was found to be much more closely related to the Lsa group than to the Vga group of proteins on the phylogenetic tree (Fig. 3). On the basis of these functional data and this clustering analysis, the novel ARE member received the designation \textit{Sal}(A) from the MLS resistance gene nomenclature center (http://faculty.washington.edu/marilynr/).

Expression and distribution of \textit{sal}(A). To study \textit{sal}(A) gene expression in strain ATCC 29059, a series of qRT-PCR experiments were conducted with and without antibiotic exposure at one-eighth the MICs of Lc and PIIA. Both antibiotics were potent inducers of the targeted RNA transcript(s), since Lc and PIIA led to an 8-fold and a 6-fold increase in \textit{sal}(A) gene expression, respectively. Specificity of induction was tested with chloramphenicol and rifampin under the same conditions, and no change in \textit{sal}(A) expression was observed. Standard PCR experiments checked that \textit{sal}(A) could be amplified from all other studied \textit{S. sciuri} strains (see Fig. S2 in the supplemental material).

**DISCUSSION**

The \textit{sal}(A) gene characterized in this study was found to be highly divergent from all other thus-far-described ARE-encoding genes. It had not been detected either previously by PCR (10) nor here by RMA. Only WGS and BLAST analyses against sequences in ABCISSE, a relational database displaying the properties and total
subunit compositions of experimentally investigated ABC systems (21), pointed out its likely involvement in the LSA resistance phenotype of *S. sciuri* strain ATCC 29059. This assumption was confirmed by its subsequent cloning into *S. aureus* strain RN4220. Based on the gain in MIC values of Lc and PIIA obtained by use of this reference strain, the novel gene has been definitively included in the long list of MLS resistance determinants (22).

Despite its demonstrated inducibility in strain ATCC 29059, the *sal(A)* gene appears to be loosely regulated by the presence of lincosamide or S₅ antibiotics in culture medium. Neither the mechanism of gene induction nor the determination of the transcription start site(s) was investigated here. Indeed, that type of analysis was not considered of high clinical relevance, mainly because the LS₅ resistance trait could be easily evidenced in classical antibiograms without preincubation treatment for all tested *S. sciuri* strains. Since PIIA-loaded disks are not commercially available, the use of Lc disks would be the best way to detect the LS₅ phenotype. While *lnu* (formerly *lin*) resistance genes confer high-level resistance to lincomycin in staphylococci (up to a MIC of 128 μg/ml) and no detectable resistance to clindamycin, the *sal(A)*, *vga*, and *lsa* genes confer low-level resistance to both antibiotics (23, 24). We speculate that pleuromutilin antibiotics, such as tiamulin, preferentially used in veterinary medicine but not tested here because of their unavailability, would be good alternatives to screen for the LSA and pleuromutilin resistance phenotype. In light of the present study, it can be anticipated that the high-level resistance to tiamulin exhibited by the recently collected *lnu(A)*-carrying *S. sciuri* clinical strain (23) resulted from the indigenous *sal(A)* gene. This assumption needs to be further investigated.

As judged by its genetic environment in strain ATCC 29059, *sal(A)* cannot be easily disseminated by horizontal gene transfer, and its localization between two housekeeping genes of the staphylococcal core genome is a rather limited way of spreading, especially within the genus *Staphylococcus*, where generalized trans-

**FIG 3** Phylogenetic relatedness among the experimentally tested ARE proteins. The maximum likelihood tree, using YkpA as an outgroup REG protein, was built by PhyML using the best-fitting model, which consists of WAG plus I (0.05) plus G (2.54), and 1,000 bootstraps. The basal group corresponds to ARE proteins from the MLS producers. Three main ARE groups were recovered; they correspond to Lsa proteins, Msr proteins, and Vga proteins. Sal(A) is closer to Lsa than to Vga.
duction and transformation events are scarce for most of the studied chromosomal markers (25, 26). Moreover, the presence of sal(A) in the chromosome of strain ATCC 29059, a strain of *S. sciuri* isolated from the healthy skin of a Virginia opossum in 1972 (27), long before the intensive use of antimicrobials in medicine and agriculture practices, lends support to the hypothesis of a natural resistance trait. Since sal(A) was successfully amplified by the use of external gene primers for the three *S. sciuri* subspecies and since this detection is correlated with both the LS$_A$ phenotype of the strains and the evolution time of the species, stable inheritance of the resistance gene from a common ancestor is likely. Such a progenitor might have belonged to the Gram-‐positive bacterial lineage, prior to the separation of the *Firmicutes* and *Actinobacteria* phyla (28). The latter phylum is recognized as the source of more than two-‐thirds of clinically useful antibiotics, and each MLS producer strain was found to have at least one ARE encoded by its genome in the vicinity of antibiotic biosynthesis genes (29, 30, 38). In these MLS-‐producing strains, ARE proteins confer self-‐resistance or natural immunity against the antibiotic that they synthesize. It might be speculated, in light of recently reported findings, that a ubiquitous microorganism like *S. sciuri*, which has been isolated in soil, sand, and marsh grass (31), has maintained an indigenous Sal(A) determinant to survive within lincosamide-‐containing environments. This is probably what also happens with many other species of the *Bacilli* class, as recently exemplified by molecular characterization of VmlR in *E. faecalis* (32) and Lsa(A) in *E. faecalis* (8), which confer the same intrinsic resistance properties, the LS$_A$ resistance phenotype, upon the bacteria in which they reside.

In contrast to Lsa(A), from which were derived the lso(B) (33), lsa(C) (34), and lsa(E) (7) genes, neither sal(A)-‐ nor vmlR-‐related genes have been detected to date outside their linked species, and the present study did not find a reservoir of LS$_A$ resistance genes in staphylococci. The origin of vga variant genes remains obscure at this time. According to us, it is now clear that most staphylococcal species lost their ancestral ARE-‐encoding gene, represented by sal(A) in *S. sciuri*, during their species evolution; bit by bit, they became highly restricted to the skin and mucous membranes of mammal, fish, reptile, and bird hosts. Staphylococcal strains, currently confronted with much MLS use in a wide variety of human processes, have developed various successful strategies for acquiring a whole range of MLS resistance genes from the environment (35). While *S. sciuri* and sal(A) cannot be seen as immediate threats to public health, both should be included in the monitoring menus for MLS antibiotic resistance surveys. In addition, the report of the presence of a sal(A)-‐related pseudogene in *S. saprophyticus* might be of epidemiological concern, especially since none of the thus-‐far-‐described resistance genes can be found in an ML-‐resistant strain (36). Indeed, it was recently reported that only one nucleotide mutation within a resident cryptic ARE-‐encoding gene of the *Enterococcus faecium* genome would be enough to generate an LS$_A$-‐resistant strain from a susceptible one (37). With respect to these documented examples, ABC-‐mediated MLS-‐R clinical cases may have been underestimated in the literature.

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