



**Microbiology**

AIMS Microbiology, 3(4): 908-914.

DOI: 10.3934/microbiol.2017.4.908

Received: 13 August 2017

Accepted: 06 November 2017

Published: 14 November 2017

<http://www.aimspress.com/journal/microbiology>

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*Research article*

## **Intrinsic role of coagulase negative staphylococci *norA*-like efflux system in fluoroquinolones resistance**

**Ligozzi Marco, Galia Liliana, Bertocelli Anna, and Mazzariol Annarita \***

Department of Diagnostics and Public Health, University of Verona, Verona, Italy

\* **Correspondence:** Email: [annarita.mazzariol@univr.it](mailto:annarita.mazzariol@univr.it); Tel: +39-045-8027690; Fax: +39-045-8027101.

**Abstract:** NorA is a *Staphylococcus aureus* multidrug transporter that exports structurally distinct compounds including fluoroquinolones. In this study *norA*-like genes of *Staphylococcus epidermidis* (*norA*<sub>SEP</sub>) and *Staphylococcus haemolyticus* (*norA*<sub>SHAE</sub>) were identified and sequenced. The nucleotide identity of *norA*<sub>SEP</sub> and *norA*<sub>SHAE</sub> with *norA* was 75.3 and 74.1%, respectively, and the amino acid identity 87.7 and 86%, respectively. Inactivation of *norA*<sub>SEP</sub> increased the ciprofloxacin susceptibility of *E. coli* DH5 $\alpha$  carrying the pB SK 198 *norA*<sub>SEP</sub> EZ cat *norA*<sub>SEP</sub> plasmid.

**Keywords:** *S. epidermidis*; *NorA*-like gene; efflux pumps; *S. haemolyticus*; coagulase negative staphylococci

### **Abbreviations:**

FQ	fluoroquinolones;
FQR	fluoroquinolones resistance;
CoNS	coagulase-negative staphylococci;
MIC	minimal inhibitory concentration;
EUCAST	European Committee of Antimicrobial Susceptibility Testing

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## 1. Introduction

Efflux-mediated fluoroquinolone resistance has been described in Gram-positive species [1]. In *Staphylococcus aureus*, *Streptococcus pneumoniae*, viridans streptococci, enterococci, and *Bacillus subtilis* FQ exporting systems belong to the MSF family, the best characterized being NorA of *S. aureus* and Bmr/Blt of *B. subtilis*, responsible for resistance to FQ, basic dyes, puromycin, chloramphenicol, and tetraphenylphosphonium [2,3].

The most information about the efflux-mediated mechanisms of FQR in staphylococci is available for *S. aureus*. The *norA* gene is expressed weakly in wild-type *S. aureus* cells, and *norA*-mediated resistance probably depends upon mutational upregulation of the gene expression, concomitant increase in production of the *norA* efflux pump [4,5] and target site mutations.

Less is known about *Staphylococcus epidermidis* and other coagulase-negative staphylococci (CoNS). Target site mutations have been described [6]. Active efflux, as suggested by blocking by reserpine, contributes substantially to the resistance phenotype in some strains of CoNS [7,8], and role of efflux overexpression of a mutation in an untranslated sequence before *norA*-like gene [9].

The present study was undertaken to investigate the role in intrinsic fluoroquinolones resistance of homologues of *norA* MFS-type efflux transporter in CoNS, namely *S. epidermidis* and *S. haemolyticus*. For this purpose, the *norA*-like gene was first sequenced, insert in a plasmid and then cloned in *E. coli* DH5 $\alpha$  and finally the cloned gene was destroyed by transposon mutagenesis.

## 2. Materials and Method

The presence of *norA*-like sequences was investigated in CoNS strains from our collection (*S. epidermidis* 198, *Staphylococcus capitis* 92, *Staphylococcus chonii* 147, *Staphylococcus haemolyticus* 256) using PCR and degenerate oligonucleotide primers based on the highly-conserved motif of MSF-type efflux pumps (*norA* deg1: 5'-AATGTTTCAAAGWCAGAT-3'; *norA* deg2: 5'-KTTGCWGGWRCATTAGGT-3', W = A, T; K = G, T; R = A, G). PCR was performed in 0.2 ml tubes in an MJ Research (BioRad, Hercules, CA). Standard PCR reactions were carried out in 50  $\mu$ l with the following final concentrations: 50 mM KCl, 10 mM Tris-HCl (pH 9.0 at 25  $^{\circ}$ C), 0.1% Triton X-100, 1.5 mM MgCl<sub>2</sub>, 100  $\mu$ M each of dNTP, 0.5  $\mu$ M of each primer, 0.5 U AmpliTaqGold DNA polymerase (Applied Biosystems, Foster City, CA). Standard amounts of DNA were added: 30 ng genomic DNA or plasmid DNA. PCR cycling conditions were as follows: an initial denaturation at 94  $^{\circ}$ C for 5 min followed by 5 cycles of 94  $^{\circ}$ C for 30 sec, annealing at 37  $^{\circ}$ C and extension at 72  $^{\circ}$ C for 60 sec. This was followed by 30 cycles consisting of 94  $^{\circ}$ C for 30 sec, 50  $^{\circ}$ C for 30 sec and 72  $^{\circ}$ C for 60 sec and a final 5-min extension step at 72  $^{\circ}$ C.

Transposon mutagenesis, performed with EZ:TN transposon system (Epicentre, Biotechnologies, Madison, WI) was used to obtain a *norA*<sub>SEP</sub> mutant in accordance with the manufacturer's instructions. The entire *norA*-like gene of *S. epidermidis* (*norA*<sub>SEP</sub>) was amplified using primers derived from the gene sequence (*norA*<sub>SEP</sub> fw 5'-CATAACCACGCACTACTTTCT-3'; *norA*<sub>SEP</sub> rev 5'-GACACAGAATTTCGTCTTGAAC-3') and cloned in the pBluescript SK plasmid (Stratagene, La Jolla, CA), resulting in plasmid pB SK 198 *norA*<sub>SEP</sub> Transposon insertion into *norA*<sub>SEP</sub> was done by incubating the plasmid containing *norA*<sub>SEP</sub> with an equal molar amount of

the EZ:TN <CAT> transposon, encoding chloramphenicol resistance, and EZ:TN transposase for 2 h at 37 °C according to the manufacturer's instructions. Following transformation of chemically competent *E. coli* DH5 $\alpha$  cells (Stratagene, La Jolla, CA) with *in-vitro* insertion reaction, clones were selected by growth on 10  $\mu$ g/ml chloramphenicol agar plates. Chloramphenicol-resistant clones were submitted to PCR analysis with *norA*<sub>SEP</sub> primers.

The MIC of ciprofloxacin for both *E. coli* DH5 $\alpha$  and *S. epidermidis* 198 were determined in triplicate with E-test strips according to the EUCAST guidelines [10].

SDS-polyacrylamide gel electrophoresis of NorA of *E. coli* DH5 $\alpha$  wild-type and harboring the recombinant plasmid. pB SK 198 *norA*<sub>SEP</sub> was done. *E. coli* strains were grown in LB broth (5 ml) with ampicillin (100  $\mu$ g/ml) at 37 °C with shaking (300 rpm). At the absorbance of 600 nm, cells were harvested and resuspended in a 0.1 volume of loading buffer and incubated for 2 min at 100 °C.

### 3. Results

An amplification product of 190 bp from total DNA both of *S. aureus* SA 1199 (kindly provided by G. W. Kaatz), and of different species of the CoNS were obtained.

*S. epidermidis* 198 was selected for subsequent cloning experiments. The 190 bp PCR product was sequenced by the *Taq* dye-deoxy terminator method with a 377 DNA Sequencing System (Applied Biosystems, Foster City, CA). Sequence analysis and alignments were done using the Genebase version 1 computer software (Applied Maths, Kortrijk, Belgium) and revealed a high degree of homology with the corresponding sequence of *S. aureus norA*.

To perform the complete sequence of the *norA*-like gene an inverse PCR approach [11] starting from the 190 bp sequence found in the *S. epidermidis* 198 chromosome was followed. The *norA*-like gene was found to be located in a 1.7-kb fragment whose nucleotide sequence showed one open reading frame (nucleotides 568 to 1728) long enough to encode a polypeptide of 387 amino acids (accession number AJ621598). Putative promoter sequences were found at nucleotides 450 to 455 (TACAAT) and nucleotides 426 to 431 (TTGTCA), which well match the consensus sequences (TATAAT and TTGACA) for the -10 and -35 regions of *E. coli* promoters. An inverted repeat, which might act as a transcription terminator, was found at nucleotides 1797 to 1834. The gene was 1161 bp nucleotides in length and consisted of 387 amino acids. The sequence revealed a nucleotide identity of 75.3% with *norA* of *S. aureus* 1199. The complete *norA* gene of *S. epidermidis* 198 was designated as *norA*<sub>SEP</sub>. Using a similar strategy, the sequence of the *norA*-like gene of *S. haemolyticus* 256 (*norA*<sub>SHAE</sub>) was also performed (accession number AJ621601).

Figure 1 shows the alignments between the deduced amino-acid sequence of *S. aureus* NorA protein and that of NorA<sub>SEP</sub> and NorA<sub>SHAE</sub>. The nucleotide identity of *norA*<sub>SEP</sub> and *norA*<sub>SHAE</sub> with *norA* was 75.3 and 74.1% and the amino-acid identity 87.7 and 86%, respectively. These results indicate a high degree of homology between the *norA* genes of CoNS and the *norA* gene of *S. aureus*.

S. aureus SA1199	MNKQILVLYFNIFLI FLGIGLVI PVLFPVYLKDLGLTGSDDLGLLVAAAFALSQMIISPFGGT
S. epid 198	MKKQLFILYFNIFLI FLGIGLVI PVLFPVYLKDLGLKGSDDLGLMVAAFALSQMIISPFGGT
S. haem 256	MKKQLFILYFNIFLI FLGIGLVI PVLFPVYLKDLGLKGSDDLGLMVAAFALSQMIISPFGGT
	* ** .*****
S. aureus SA1199	LADKLGKKLIICIGLILFSVSEFMFAIQNFLILMLSRVIGGMSAGVMMPGVTGLIADIS
S. epid 198	LADKLGKKLIICIGLVFFAVSEFMFAAGQSFTILISRVLGGFSAGVMMPGVTGMIADIS
S. haem 256	LADKLGKKLIICIGLIFFAVSEMLAAGRSFTILISRVLGGFSAGVMMPGVTSVIANIS
	*****. *.** * * *. * ** .***. * ***** .** **
S. aureus SA1199	PSHQKAKNFGYMSAI INSGFILGPGIGGFMAEVSHRMPFFYFAGALGILAFIMSIVLIHDP
S. epid 198	PGADKAKNFGYMSAI INSGFILGPGFGFLAEISHRLPFYVAGTLGVVAFIMSVLLIHNP
S. haem 256	PGADKAKNFGYMSAI INACFILGPGGLGFLSEISHRLPFYVAGTLVGVAFIMSVLLIHNP
	* *****. ***** **..*.***.*** **.* . *****.*** *
S. aureus SA1199	KKVSTNGFQKLEPQLLTKINWKVFITPVILTLVLSFGLSAFETLYSLYTADKVNYS PKDI
S. epid 198	QKATTDGFHQYQPELFTKINWKVFITPVILTLVLAFLSAFETLFSLYTADKVNYPKDI
S. haem 256	HKATTDGFHQYQPELFTKINWKVFITPVILTLVLAFLSAFETLFSLYTADKVNYPKDI
	.* .* **.. .*.* *****.*****.*****.*****.****
S. aureus SA1199	SIAITGGGIFGALFQIYFFDFKFMKYFSELTFAWSLIYSVIVLVLLVIADGYWTIMVISF
S. epid 198	SIAIIGGGVFGALFQVFFDFKFMKYMSELNFIAWSLLYSIVLVMLVLANGYWTIMII SF
S. haem 256	SIAIIGGGVFGALFQVFFDFKFMKYMSELNFIAWSLLYSIVLVMLVLANGYWTIMII SF
	**** **..*****.***** **..*****.*** **..**.* *****.***
S. aureus SA1199	VVFIGFDMIRPAITNYFSNIAGDRQGFAGGLNSTFTSMGNFIGPLIAGALFDVHIEAPIY
S. epid 198	VVFIGFDMIRPALTNYFSNIAGKRQGFAGGLNSTFTSMGNFIGPLVAGALFDVNLEFPLY
S. haem 256	VVFIGFDMIRPALTNYFSNIAGKRQGFAGGLNSTFTSMGNFIGPLVAGALFDVNLEFPLY
	*****.***** *****.*****.*****.*** **.*
S. aureus SA1199	MAIGVSLAGVIVLIEKQHRAKLKQDDL
S. epid 198	MAIAVSLSGIIIFIEKGLKSRKEAN-
S. haem 256	MAIAVSLSGIIIFIEKGLKSRKEAN-
	*** **..**.*. *** ... *

**Figure 1.** The deduced amino acid sequence alignment of *S. aureus* NorA, amino acid sequence, *S. epidermidis* 198 NorA and *S. haemolyticus* 256 NorA alignments. Asterisks and dots indicate residues that are identical and similar to the three amino-acids sequences respectively.

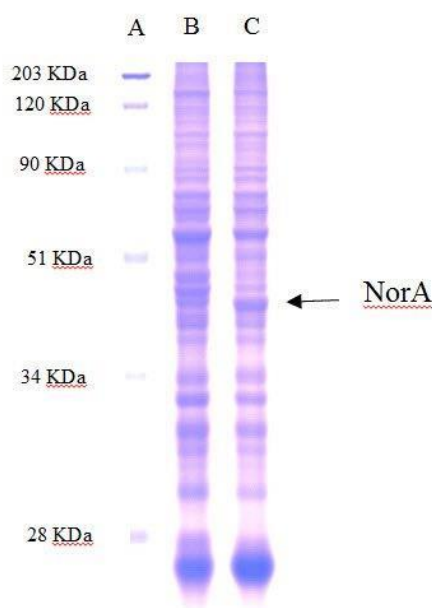
The insertional mutagenesis approach to inactivate the *norA* gene was used to determine the physiological function of the protein encoded by *norA*<sub>SEP</sub>.

Several mutants of *S. epidermidis* were obtained by *in-vitro* transposition techniques [12]. The entire *norA*<sub>SEP</sub> gene was amplified using primers derived from the gene sequence and cloned in the pBluescript SK plasmid (Stratagene, La Jolla, CA), resulting in plasmid pB SK 198 *norA*<sub>SEP</sub>. Transposon insertion into *norA*<sub>SEP</sub> was done with the EZ:TN <CAT> transposon and clones in chemically competent *E. coli* DH5 $\alpha$  cells (Stratagene, La Jolla, CA) with *in-vitro* insertion reaction were selected by growth on 10  $\mu$ g/ml chloramphenicol agar plates. Chloramphenicol-resistant clones were submitted to PCR analysis with *norA*<sub>SEP</sub> primers. Insertion of the transposon in *norA*<sub>SEP</sub>

increased the amplicon length from 1.6 kb (*norA<sub>SEP</sub>* gene without the transposon insertion) to 2.4 kb. Several clones were obtained which gave amplicons of the expected length. From one of these clones (*E. coli* DH5 $\alpha$  198 *norA<sub>SEP</sub>*) the recombinant plasmid containing the *norA<sub>SEP</sub>::cat* fragment (pB SK 198 *norA<sub>SEP</sub>* EZ cat) was purified and sequenced.

In Table 1 are reported the results of MICs of some fluoroquinolones that are NorA efflux substrate as ciprofloxacin, levofloxacin and ofloxacin. MICs are measured also for substrate as the moxifloxacin that effect NorB but NorA efflux pumps. All antibiotics were tested alone and in presence of carbonyl *m*-chlorophenylhydrazone (CCCP) an efflux pumps inhibitor. *E. coli* DH5 $\alpha$  carrying the plasmid pB SK 198 *norA<sub>SEP</sub>* had a ciprofloxacin MIC of 0.25  $\mu$ g/ml, eight times higher than the MIC of *E. coli* DH5 $\alpha$  carrying the plasmid pB SK 198 *norA<sub>SEP</sub>* EZ cat *norA<sub>SEP</sub>* (MIC = 0.032  $\mu$ g/ml) and *E. coli* DH5 $\alpha$  without plasmid. Similar effect is register for levofloxacin and ofloxacin. There are no effects indeed in the MICs of moxifloxacin as well the tetracycline, since they are not substrates of NorA pump.

In order to confirm expression of the efflux pump protein we performed a SDS-polyacrylamide gel electrophoresis (PAGE) and NorA expression analysis in *E. coli* DH5 $\alpha$  wild-type and harboring the recombinant plasmid pB SK 198 *norA<sub>SEP</sub>*. In Figure 2 we showed the presence of around 42 KDa protein in the strains harboring the recombinant plasmid only.



**Figure 2.** SDS-PAGE and NorA expression analysis in *E. coli* DH5 $\alpha$ . Line A: Pre-stained Molecular Weight (Biorad, Milan Italy); line B: *E. coli* DH5 $\alpha$  recipient strain; line C: *E. coli* DH5 $\alpha$  pB SK 198 *norA<sub>SEP</sub>*.

**Table 1.** MICs value of *E. coli* DH5 $\alpha$  and its trans-conjugants for fluoroquinolones and tetracycline, in absence and presence of CCCp.

Strains	MIC (mg/L)									
	Ciprofloxacin		Levofloxacin		Ofloxacin		Moxifloxacin		Tetracycline	
	-CCCp	+CCCp <sup>a</sup>	-CCCp	+CCCp <sup>a</sup>	-CCCp	+CCCp <sup>a</sup>	-CCCp	+CCCp <sup>a</sup>	-CCCp	++CCCp <sup>a</sup>
<i>E. coli</i> DH5 $\alpha$	0.032	0.032	0.015	0.015	0.015	0.015	0.0075	0.0075	0.5	0.5
<i>E. coli</i> DH5 $\alpha$ pB SK 198 <i>norA</i> <sub>SEP</sub>	0.25	0.015	0.12	0.015	0.06	0.015	0.015	0.015	1	1
<i>E. coli</i> DH5 $\alpha$ pB SK 198 <i>norA</i> <sub>SEP</sub> EZ cat	0.032	0.032	0.015	0.015	0.015	0.015	0.015	0.015	0.5	0.5

<sup>a</sup> Carbonyl *m*-chlorophenylhydrazine (CCCp) was added with concentration of 1  $\mu$ g/ml.

#### 4. Conclusion

Our results demonstrated that *norA*-like genes play an important role in the intrinsic FQR in CoNS. The resistance level to FQ due to NorA efflux pumps is not elevated. Like in other species such as *S. aureus*, *S. pneumoniae* the overexpression of efflux pumps combined with other mechanisms may contribute to increase the resistance at high level.

#### Acknowledgments

We thank G.W. Kaatz for the kind gift of *S. aureus* SA 1199.

#### Conflict of Interest

All authors declare no conflicts of interest in this paper.

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