

Evaluation of Pleuromutilins Resistance Mechanisms Among Surveillance Clinical Isolates: Results from the Worldwide Surveillance Program for Lefamulin in 2019

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INTRODUCTION

- Lefamulin (Xenleta[™]) is a first in human pleuromutilin antibiotic approved in the US, Europe, and Canada for the treatment of community acquired bacterial pneumonia (CABP).
- Both intravenous and oral formulations are available for the treatment of CABP in adults. Lefamulin has a targeted spectrum of activity against key pathogens including typical Grampositive, fastidious Gram-negative and atypical organisms, including strains resistant to standardof-care therapies.
- Lefamulin is a first-in-class, semi-synthetic pleuromutilin that inhibits bacterial protein synthesis by binding to the 50S ribosomal subunit at the A- and P- sites in the peptidyl transferase center (PTC) via an "induced-fit" mechanism, which prohibits the correct positioning of the tRNA.
- The in vitro activity and emergence of resistance to lefamulin are being monitored against a global collection of Gram-positive and fastidious Gram-negative organisms through the SENTRY Antimicrobial Surveillance Program.
- This study characterized possible resistance mechanisms among surveillance isolates with elevated lefamulin MICs collected globally in 2019.

MATERIALS AND METHODS

Bacterial isolates

- A total of 3,975 Staphylococcus aureus, coagulase-negative staphylococci (CoNS), Streptococcus pneumoniae, β-haemolytic streptococci, viridans group streptococci, Moraxella catarrhalis, and Haemophilus spp. were included as part of the surveillance study for 2019 (Table 1).
- A total of 36 (0.8%) isolates met the MIC screening criteria based on the US FDA and EUCAST breakpoints or tentative epidemiological cut-off (ECOFF) values.
- *S. aureus*: ≥0.5 mg/L, with an FDA susceptible breakpoint ≤0.25 mg/L
- CoNS and β-haemolytic Streptococcus spp.: ≥0.5 mg/L
- S. pneumoniae: ≥1 mg/L, with an FDA susceptible breakpoint ≤0.5 mg/L
- H. influenzae: ≥4 mg/L, with an FDA susceptible breakpoint ≤2 mg/L
- Bacterial isolate identification was confirmed by standard algorithms and supported by matrix-assisted laser desorption ionization-time of flight mass spectrometry (Bruker Daltonics, Bremen, Germany) and genome sequencing.

Antimicrobial susceptibility testing

- Isolates were tested for susceptibility by broth microdilution following Clinical and Laboratory Standards Institute (CLSI) M07 (2018) guidelines using frozen-form broth microdilution panels containing CAMHB; supplemented with 2.5-5% lysed horse blood for streptococci; HTM was used for H. influenzae.
- Quality assurance was performed by concurrently testing the CLSI-recommended quality control reference strains.

Characterization of resistance mechanisms by next-generation sequencing

- Selected isolates had total genomic DNA extracted by the fully automated Thermo Scientific™ KingFisher[™] Flex Magnetic Particle Processor (Cleveland, OH, USA).
- DNA libraries were prepared using the Nextera™ library construction protocol (Illumina, San Diego, CA, USA) according to manufacturer instructions.
- FASTQ format sequencing files for each sample set were assembled independently using de novo assembler SPAdes 3.9.0.
- Additional sequences of intrinsic genes associated with the pleuromutilin binding site, including 23S rRNA (PTC), rplC (L3), rplD (L4), and rplV (L22), were evaluated against a susceptible reference strain of the corresponding species.
- All intrinsic 23S rRNA target genes or ribosomal protein amino acid sequences were considered wild type if 100.0% homology with the respective reference sequences was displayed.

Transcriptional levels of AcrA

- Haemophilus spp. were subjected to sequencing analysis of AcrR and the quantification of AcrAB-ToIC expression.
- The total mRNA was extracted and purified using the RNeasy Mini Kit in the QIAcube workstation according to manufacturer instructions and treatment with RNAse-free DNase (Promega, Madison,
- The quantification of mRNA and sample quality were assessed using the RNA 6000 Nano kit from Agilent on the Agilent 2100 Bioanalyzer, according to manufacturer instructions.
- The transcription levels of acrA were determined by relative quantification using real-time PCR (qRT-PCR) assays in the StepOne Plus instrument (Life Technologies, Foster City, California).
- Transcription levels of acrA were measured by the quantification of the target gene mRNA using a normalized expression analysis method with a housekeeping reference gene (gyrA).

Multilocus sequence typing

 Multilocus sequence typing (MLST) was performed by extracting the previously defined set of 7 housekeeping gene fragments (~500 bp) and comparing with allelic variants on the MLST website.

RESULTS

- Among 873 S. aureus included in the 2019 lefamulin surveillance program, 2 (0.2%) isolates had lefamulin MIC values of 1 or >16 mg/L and were selected for this study (Table 1).
- These 2 non-susceptible isolates harboured vga(A) or lsa(E) 871 (99.8%) S. aureus isolates showed lefamulin MICs of 0.015-0.25 mg/L (MIC_{50/90} of 0.06/0.12 mg/L) and are susceptible based on FDA and EUCAST breakpoints (S ≤0.25 mg/L).
- A total of 24 of 329 (7.3%) CoNS, spanning 4 total species with lefamulin MIC values of 0.5–≥32 mg/L, were selected for genetic characterization.
- Most CoNS isolates carried vga gene variants (19/24; 79.2%) and had lefamulin MIC results of 0.5–≥16 mg/L).

^a Shading represents MICs above the breakpoints or ECOFF values for the respective species or groups of species or groups of species. S. bovis group display higher and broader MIC distributions for lefamulin and were not included in the study.

Table 1. Lefamulin MIC distributions obtained during the 2019 surveillance program

	No. and cumulative % of isolates inhibited at MIC (mg/L) of: ^a											NAIO	NAIC			
Organism/organism group (no. of isolates)	≤0.008	0.015	0.03	0.06	0.12	0.25	0.5	1	2	4	8	16	32	>32	MIC ₅₀	MIC ₉₀
Staphylococcus aureus (873)	0.0	3 0.3	46 5.6	490 61.7	312 97.4	20 99.7	0 99.7	1 99.9	0 99.9	0 99.9	0 99.9	0 99.9		1 100.0	0.06	0.12
Coagulase-negative staphylococci (329)	6 1.8	10 4.9	120 41.3	132 81.5	35 92.1	2 92.7	4 93.9	7 96.0	4 97.3	1 97.6	2 98.2	3 99.1		3 100.0	0.06	0.12
Streptococcus pneumoniae (1,384)	3 0.2	20 1.7	131 11.1	510 48.0	571 89.2	143 99.6	3 99.8	3 100.0							0.12	0.25
β-haemolytic streptococci (367)			253 68.9	94 94.6	17 99.2	2 99.7	0 99.7	0 99.7	0 99.7	0 99.7	0 99.7	0 99.7	1 100.0		≤0.03	0.06
Haemophilus influenzae (490)	2 0.4	5 1.4	1 1.6	5 2.7	18 6.3	73 21.2	214 64.9	123 90.0	43 98.8	3 99.4	3 100.0				0.5	1
Moraxella catarrhalis (300)	6 2.0	5 3.7	5 5.3	177 64.3	103 98.7	4 100.0									0.06	0.12

- Two S. epidermidis isolates from Mexico carried cfr (2/24; lefamulin MIC, 8 mg/L). These 2 isolates were from 2 patients hospitalized in the same medical centre and exhibited the same MLST.
- One S. epidermidis from Germany carried a G2576T mutation (1/24; lefamulin MIC of 0.5 mg/L). One S. haemolyticus and 1 S. saprophyticus with a lefamulin MIC of 0.5–1 mg/L did not show any mechanisms previously associated with lefamulin resistance and MICs represent the upper end of the MIC wildtype distributions.
- Three S. pneumoniae were non-susceptible to lefamulin (MIC, 1 mg/L), but these isolates did not show any clear mechanisms implicated with pleuromutilin resistance and MIC values represent the upper end of the MIC wildtype distributions (Tables 1 and 4)
- A single Streptococcus agalactiae (0.3% of β-haemolytic streptococci) with an elevated lefamulin MIC (32 mg/L) was detected in this study (collected in Italy).
- This strain carried Isa(E) (Tables 1 and 4)
- A total of 6 (1.2%) *H. influenzae* showed lefamulin nonsusceptible MICs of 4–8 mg/L.
- All but 1 isolate (1129277) showed derepression of acrA and acrB expression due to a premature stop codon in *acrR*.
- Additionally, these isolates had either an 23S rRNA (A2058G or A2059G) or a L4/L22 alteration, except for isolate 1126213 that had an acrR with a premature stop codon only (Table 5).

Table 2. Molecular epidemiology and resistance mechanism results for S. aureus isolates

Collection	міст	Country	MICa	Resistance mechanisms								
					Gene		Ribosomal mutations ^b					
no.	IVILOI	Country	(mg/L)	cfr	Isa(E)	vga(A)	23S rRNA	L3	L4	L22		
1099706	9	Mexico	>16		+		WT	WT	V118D	WT		
1127385	9	Brazil	1			+	WT	WT	V118D	WT		

MLS1, multilocus sequence typing; W1, wild type ^a MIC, minimal inhibitory concentration.

The 23S rRNA mutational analysis was performed on nucleotide sequences. Mutations outside of PTC were observed but were considered polymorphisms and suppressed. The 23S rRNA sequence was designated as WT. Protein sequences that were analysed for annotating were L3, L4, and L22.

Table 3. Molecular epidemiology and resistance mechanisms results for Staphylococcus spp. other than S. aureus

					Resistance mechanisms						
Collection	Organism	МІСТ	Country	MICa		Ribosomal mutations ^b					
no.	Organisin	IVILSI	Country	(mg/L)	Gene	23S rRNA	L3	L4	L22		
1117096	S. capitis	NA	Australia	0.5	vga(A)	WT	T83A	WT	WT		
1118328	S. capitis	NA	Australia	4	vga(A)	WT	T83A	178N	WT		
1074687	S. capitis	23	Mexico	8	vga(A)	WT	S158Y,D159Y	WT	WT		
1090848	S. epidermidis	23	Mexico	8	cfr	WT	S158Y,D159Y	WT	WT		
1090853	S. epidermidis	23	Mexico	8	cfr	WT	S158Y,D159Y	WT	WT		
1091111	S. epidermidis	16	USA	1	vga(A)	WT	WT	WT	WT		
1092976	S. epidermidis	2	Germany	0.5	G2576T	WT	H146R	G71_R72insG	WT		
1101658	S. epidermidis	87	England	16	vga(A)	WT	WT	WT	WT		
1101683	S. epidermidis	35	England	16	vga(A)	WT	WT	WT	WT		
1104351	S. epidermidis	25	Spain	>16	vga(A)	WT	WT	WT	WT		
1107456	S. epidermidis	5	France	1	vga(A)	WT	WT	WT	WT		
1112499	S. epidermidis	87	Wales	1	vga(A)	WT	WT	WT	WT		
1112518	S. epidermidis	142	Wales	2	vga(A)	WT	WT	WT	WT		
1116902	S. epidermidis	87	Australia	1	vga(A)	WT	WT	WT	WT		
1123315	S. epidermidis	87	Belgium	1	vga(A)	WT	WT	WT	WT		
1123338	S. epidermidis	87	Belgium	0.5	vga(A)	WT	WT	WT	WT		
1123389	S. epidermidis	5	Belgium	2	vga(A)	WT	WT	WT	WT		
1126561	S. epidermidis	59	Argentina	2	vga(A)	WT	WT	WT	WT		
1126563	S. epidermidis	59	Argentina	2	vga(A)	WT	WT	WT	WT		
1126635	S. epidermidis	87	Argentina	1	vga(A)	WT	WT	WT	WT		
1099446	S. haemolyticus	29	Italy	16	vga(A)LC	WT	T82A,G137A	V130G,I142V	WT		
1103323	S. haemolyticus	3	USA	0.5		WT	T82A,G137A	V130G,I142V	WT		
1107535	S. haemolyticus	29	France	>16	vga(A)LC	WT	T82A,G137A	V130G,I142V	WT		
1117552	S. haemolyticus	29	USA	>16	vga(A)LC	WT	T82A,G137A	V130G,I142V	WT		
1125669	S. saprophyticus	NA	Taiwan	1		WT	I35V	WT	WT		
	S. saprophyticus sequence typing; NA, MLS			1 on-existent	WT, wild type	WT	I35V	WT	WT		

b The 23S rRNA mutational analysis was performed on nucleotide sequences. Mutations outside of PTC were observed but were considered polymorphisms

and suppressed. The 23S rRNA sequence was designated as WT. Protein sequences that were analysed for annotating were L3, L4, and L22.

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WT, wild type; NA, not available.

protection.

CONCLUSIONS

negative clinical isolates.

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related to preparing this poster.

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Table 4. Molecular epidemiology results and resistance mechanisms obtained for Streptococcus spp.

The 23S rRNA mutational analysis was performed on nucleotide sequences. Additional mutations outside of PTC were observed but were considered

Table 5. Molecular epidemiology results and resistance mechanisms obtained for *H. influenzae*

Q57X

The 23S rRNA mutational analysis was performed on nucleotide sequences. Protein sequences that were analysed for annotating were L3, L4, and L22.

Gram-positive and fastidious Gram-negative isolates causing human infections with

• Lefamulin resistance mechanisms identified in S. aureus, CoNS, and streptococcal

isolates were vga(A) and lsa(E), which have been described to mediate ribosomal

Target site alterations, such as mutations in the large ribosomal proteins or 23S rRNA

Longitudinal surveillance studies will continue to monitor the emergence of resistance

and in vitro stability of lefamulin activity among Gram-positive and fastidious Gram-

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elevated (non-susceptible) lefamulin MIC results remained rare (0.9%).

or methylation of A2503 by the Cfr methyltransferase, were less common.

- The cfr gene was detected in only 2 S. epidermidis surveillance isolates.

E144X A2058G

A2060G

Resistance mechanisms

H45R

R60H

R88 G91del

S. pneumoniae

S. pneumoniae

S. pneumoniae

S. agalactiae

olymorphisms and suppressed. Protein sequences that were analysed for annotating were L3, L4, and L22.

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^a MIC, minimal inhibitory concentration