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Genomic analysis of multi‑drug resistant coagulase‑negative staphylococci from healthy humans and animals revealed unusual mechanisms of resistance and CRISPR‑Cas system

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Abstract

Background Coagulase-negative staphylococci (CoNS) are evolving as major reservoirs and vectors of unusual and critical antimicrobial resistance (AMR) mechanisms.

Materials and methods In this study, the genomic characterization of 26 multidrug-resistant (MDR)-CoNS (*S. borealis*, *S. saprophyticus*, *S. sciuri*, *S. hominis*, *S. epidermidis*, *S. pasteuri*, *S. hyicus*, *S. simulans*, *S. haemolyticus*, and *S. arlettae*) previously obtained from the nasal cavity of healthy nestling storks, humans who had no contact with animals, pigs, and pig farmers, as well as dogs and dog owners from Spain was performed. High-quality draft genomes obtained by Illumina sequencing technology were used to determine their resistome, virulome, mobile genetic elements, and CRISPR-Cas types. The relatedness of three CoNS species with publicly available genomes was assessed by core-genome single nucleotide polymorphisms (SNPs).

Results AMR genes to all classes of antibiotics in staphylococci were detected including unusual ones (*mecC*, *ermT*, and *cfr*), of which their corresponding genetic organizations were analyzed. About 96.1% of the MDR-CoNS strains harbored diverse adherence or immune evasion genes. Remarkably, one enterotoxin-C and -L-carrying *S. epidermidis*-ST595 strain from a nestling stork was detected. Moreover, various plasmid bound-biocide resistance genes (*qacACGJ*) were identifed in 34.6% of the MDR-CoNS. Two genes that encode for cadmium and zinc resistance (*cadD*, *czrC*) were found, of which *czrC* predominated (42.3%). Complete CRISPR-Cas system was detected in 19.2% of the CoNS strains, of which *cas*-1, -2, and -9 predominated, especially in 75% of the *S. borealis* strains. The phylogenetic analysis identifed clusters of related *S. epidermidis* lineages with those of other countries (SNP < 100). Also, highly related *S. borealis* isolates (SNP < 10) from pigs was confrmed for the frst time in Spain.

Conclusion These fndings showed that various ecological niches harbor CoNS that presented MDR phenotypes mediated by multiple AMR genes carried by mobile genetic elements with relatively low frequency of intact CRISPR-Cas systems. Furthermore, the transmission of some CoNS species in humans and animals is strongly suggested.

Keywords Staphylococci · *cfr* · Linezolid resistance · CRISPR-Cas systems · Plasmids · Enterotoxins

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Introduction

The members of the *Staphylococcus* genus are part of the normal microbiota of the nose and skin of humans and animals (including many avian species) (Szczuka et al. [2023](#page-21-0)). In addition, coagulase-positive staphylococci could occasionally cause clinical diseases mediated by highly potent virulence genes (Aqel et al. [2023\)](#page-19-0). However, not every staphylococcal virulence gene is expressed. Instead, the expression of the genes is usually restricted to times and places and regulated by bacterial factors (Grazul et al. [2023\)](#page-20-0). Over the last years, some coagulase-negative staphylococci (CoNS) species (as *S. epidermidis*, *S. haemolyticus*, or *S. hominis*) emerged as a cause of opportunistic infections such as those in septicaemic children, or in patients with immunosuppression or medical implants, among others (França et al. [2021](#page-20-1); Heilmann et al. [2019](#page-20-2)). Most of other CoNS species are unfrequently implicated in human or animal infections, being often highly susceptible to antimicrobial agents (Merrild et al. [2023](#page-21-1); Santoiemma et al. [2020;](#page-21-2) Argemi et al. [2019](#page-19-1)). However, there have been sporadic reports of some *S. pasteuri* causing endocarditis, whereas *S. hyicus*, *S. chromogenes*, *S. lentus*, and *S. sciuri* are considered etiological agents of exudative epidermitis with zoonotic potentials (Kirk et al. [2022;](#page-20-3) Kalai et al. [2021;](#page-20-4) Li et al. [2021](#page-20-5)). Moreover, *S. saprophyticus* contracted from contaminated food have long been implicated in urinary tract infections in young teenagers (Lawal et al. [2021a,](#page-20-6) [b\)](#page-20-7). Much more recently, whole-genome sequence data of CoNS species have led to the identifcation and characterization of numerous putative virulence factors (Argemi et al. [2019\)](#page-19-1). Furthermore, CoNS could acquire clinically relevant and critical antimicrobial resistance (AMR) genes and transmit them across other species and hosts through various mobile genetic elements (mobilome) (Rossi et al. [2020](#page-21-3)). Specifcally, *S. haemolyticus* has been ranked as the most antibiotic-resistant species among the CoNS (Kranjec et al. [2021\)](#page-20-8). The transferability of AMR genes between diferent *Staphylococcus* species has been strongly suggested by the sequence similarity of their associated mobilome, especially plasmids (Souza-Silva et al. [2022\)](#page-21-4).

The *mecA* gene, its staphylococcal cassette chromosome (SCC*mec*) carrying element, and the arginine catabolic mobile element (ACME) originated from CoNS were acquired by *S. aureus* (Shokrollahi et al. [2022](#page-21-5)). *mecC*-carrying CoNS have also been reported from many countries but in very low frequencies (Loncaric et al. [2019\)](#page-21-6). Previously thought to be a wildlife MRSA trait, the continuous spread of the SCC*mec*-bound *mecC* gene in CoNS highlights their potential role in the evolutionary origin and genetic transfer to MRSA (Abdullahi et al. [2023a\)](#page-19-2).

Most methicillin-resistant CoNS (MRCoNS) strains are often found to be resistant to other non-betalactam antibiotics except for glycopeptides, which have long been utilized in the treatment of staphylococcal infections (Chajęcka-Wierzchowska et al. [2023](#page-20-9)). As the AMR epidemic keeps expanding, the few methicillin-resistant staphylococcal infections that are treated using oxazolidinones (Gostev et al. [2021\)](#page-20-10) could have promoted the emergence, spread, and persistence of linezolid resistance, as some mechanisms mediated by ARGs (*cfr*, *poxtA*, and *optrA*) are carried by plasmids (Bai et al. [2019](#page-19-3); Dortet et al. [2018](#page-20-11)). However, high-level linezolid resistance could be caused by nontransferable mechanisms mediated by mutations in the 23S rDNA, and in the ribosomal proteins L3, L4, and L22 (Ruiz-Ripa et al. [2021\)](#page-21-7).

The clustered regularly interspaced short palindromic repeats (CRISPR) and CRISPR-associated proteins (Cas) are RNA-based adaptive immunity to protect and are utilized by many bacteria against invading mobile genetic elements (MGEs) (Tao et al. [2022a\)](#page-22-0). Hence, the CRISPR–Cas system might be a potential means to prevent the acquisition of plasmid and phage invasion and even horizontal transfer of AMR genes in staphylococci (Murugesan and Varughese [2022](#page-21-8)). There are two categories of CRISPR-Cas, which are based on their proteins' structures, constituents, and modes of action (Nishimasu and Nureki [2017](#page-21-9)). The Class 1 CRISPR-Cas uses multiple protein efector complexes to break down nucleotides and can be subdivided into types -I, -III, and -IV, whereas the Class 2 CRISPR-Cas utilizes single-protein efector complexes to break down nucleotides, of which it is subdivided into types -II, -V, and -VI (Shmakov et al. [2015](#page-21-10); Makarova et al. [2015\)](#page-21-11). The types II-CRISPR-Cas systems have largely been studied and have successfully been used to delete antimicrobial resistance genes (ARGs) due to their relatively simple structures (Tao et al. [2022a](#page-22-0)). Moreover, the Type I CRISPR-Cas systems have been developed and manipulated to eliminate ARGs (Tao et al. [2022a](#page-22-0)). In this regard, certain CRISPR-Cas system prevents foreign nucleotides (such as plasmids and phages) from evading the bacteria thereby limiting the acquisition of ARGs (Tao et al. [2022a](#page-22-0)).

The genetic characterization of CoNS is necessary to understand their evolution and source distribution, reservoir hosts, and vectors of AMR transmission. In this regard, certain animal hosts such as the pigs and human workers in pig farm environments are believed to be under high antibiotic pressure and carry staphylococci presenting a high-level multidrug resistance (MDR) phenotype. However, animals in the wildlife may be at low antibiotic pressure as they are rarely exposed to antimicrobial agents (Abdullahi et al. [2021\)](#page-19-4). It is worth mentioning that the ecology and epidemiology of AMR in CoNS could be diferent from that of *S. aureus* because the CoNS species could present diferent and diverse AMR profles*.* In this study, the genomic characterization of 26 multidrug resistant-CoNS (resistant to≥four classes of antimicrobial agents) previously obtained from the nasal cavity of healthy humans without animal contact, nestling storks, pigs and pig owners, as well as dogs and their owners from Spain were performed by Illumina technology.

Materials and methods

Coagulase‑negative staphylococci strains in this study

A total of 516 non-repetitive CoNS strains were obtained in previous studies (Abdullahi et al. [2023a](#page-19-2); [b](#page-19-5); [c](#page-19-6); [2024a\)](#page-19-7) from nasal samples of healthy animals and healthy humans with diferent types of animal contact: (a) healthy nestling storks (NS) (268 isolates); (b) healthy pigs (H-P) and pig farmers (H-PF) (75 isolates); (c) healthy dogs (H–D) and dog owners (H-DO) (130 isolates); and (d) healthy humans who had no contact with animals (HH−) (113 isolates). The antimicrobial susceptibility of these isolates was previously determined by disk difusion tests, and the presence of ARGs by PCR (Abdullahi et al. [2023a](#page-19-2); [b;](#page-19-5) [c;](#page-19-6) [2024a](#page-19-7)). From this collection, 26 CoNS isolates of 10 species (*S. borealis*, *S. saprophyticus*, *S. sciuri*, *S. hominis*, *S. epidermidis*, *S. pasteuri*, *S. hyicus*, *S. simulans*, *S. haemolyticus*, and *S. arlette*) were selected to be further characterized in the present study by whole genome sequencing (WGS), and they were of the following origins: NS (*n*=4); H-P and H-PF $(n = 14)$; H–D and H-DO $(n = 4)$; and HH⁻ $(n = 4)$. The selection criteria of the strains included were as follows: (i) CoNS that presented an MDR phenotype for four or more classes of antibiotics, selecting one species each per host carrying this resistance phenotype; and (ii) MDR-CoNS isolates with similar AMR genes detected from humans and animals in the same ecological niche to detect potential transmission events.

The study protocols in which these isolates were recovered were reviewed and approved by the ethical research committees of the University of Zaragoza, the University of La Rioja and the University of Castilla La Mancha (Spain).

Genome sequencing, assembly, and phylogenetic analyses

Whole genome sequencing of the selected 26 CoNS isolates was carried out on the Illumina NextSeq platform. The MagNA Pure 96 DNA Multi-Sample Kit (Life Technologies, Carlsbad, CA, USA, 4413021) was used to extract genomic DNA according to instructions provided by the manufacturers. The Qubit 1X dsDNA HS Assay Kit (Thermo Fisher Scientifc, Scoresby, VIC, Australia) was used for DNA quantifcation, while Sequencing libraries were prepared using the Illumina Nextera XT DNA Library Preparation Kit (Illumina, San Diego, CA, USA, FC-131–1096) and sequenced on the

NextSeq 500 platform (Illumina, San Diego, CA, USA) using a 300-cycle kit to obtained paired-end 150 bp reads, as previously described (Abdullahi et al. [2023d](#page-19-8)).

All the genomes analyzed in this study were de novo assembled using SPAdes (v.3.15.5), performing the in silico typing with the settings of a minimum of 90% coverage and 80% identity. First, core-genome single nucleotide polymorphisms (SNPs) between the eight *S. epidermidis* strains in this study were detected with the NASP pipeline v.1.0.0 (Sahl et al. [2016\)](#page-21-12) after they were mapped together with a reference strain ATCC 14990 (GenBank accession number: GCA_006094375) and 31 previously published *S. epidermidis* genomes from diferent countries with similar genetic lineages from the PubMLST database [\(https://pubml](https://pubmlst.org/bigsdb?db=pubmlst_sepidermidis_strains&page=query&genomes=1) [st.org/bigsdb?db=pubmlst_sepidermidis_strains&page=](https://pubmlst.org/bigsdb?db=pubmlst_sepidermidis_strains&page=query&genomes=1) [query&genomes=1](https://pubmlst.org/bigsdb?db=pubmlst_sepidermidis_strains&page=query&genomes=1)) (identifcation [id] numbers: 32110, 32113, 32116, 41749, 42109, 43340, 43421, 43426, 43427, 43436, 43455, 43466, 43518, 43568, 43636, 43643, 43656, 43697, 43720, 43770, 43771, 43774, 43786, 43800, 43816, 43823, 43921, 44294, 44298, 44496, 44521) to obtain an *S. epidermidis* phylogenetic trees. GATK (v.4.2.2) was used to call SNPs and excluded positions featuring<90% unambiguous variant calls and $<$ 10 depth. IQ-TREE (v.2.1.2) was used to construct the phylogenetic trees using ModelFinder with 100 bootstraps. The graphical data was added to the phylogenies with iTOL v.6.6 (Letunic and Bork [2021\)](#page-20-12). To determine the relatedness of the *S. saprophyticus* from a pig and pig farmer, we used a web-based CSI phylogeny database to obtain the SNPs by mapping the genomes to a reference *S. saprophyticus* ATCC 15305 (GenBank accession no. AP008934.1) with the default parameter, except for the minimum distance between SNPs which was disabled. Also, the SNPs of the *S. borealis* from four pigs were determined by comparing them with 16 additional publicly available genomes of *S. borealis* strains available from NCBI (GenBank accession numbers: GCA_030362885, GCA_030362875, GCA_003580835, GCA_003580835, GCA_034103225, GCA_024580895, GCA_030501495, GCA_035788295, GCA_035791815, GCA_035791575, GCA_013345165, GCA_009735325, GCA_013345185, GCA_013345175, GCA_013345205, GCA_013345195) mapped with a reference strain 7067_4#69 (GenBank accession number: GCA_001224225.1) by using the web-based CSI phylogeny database following settings similar to the ones used for *S. saprophyticus.*

Annotation, typing, and in silico analysis of the CoNS genomes

The sequence types (STs) were determined with MLST v.2.16 (Jolley et al. [2018](#page-20-13)). Virulence factors, plasmid replicons, and antimicrobial resistance genes were identified

using ABRicate v.0.9.0 and the respective databases VFDB, Plasmidfinder, and Resfinder databases from the Center for Genomic Epidemiology. Mutations associated with AMR were identified using ResFinder v4.1 (Bortolaia et al. [2020](#page-20-14)) and PointFinder (Zankari et al. [2017\)](#page-22-1). Biocide and heavy metal resistance genes were identified using BACMET (Pal et al. [2014\)](#page-21-13). Phaster was used to identify all prophage elements (Arndt et al. [2016](#page-19-9)). The SCC*mec* types were assigned using SCC*mec*Finder 1.2 (<https://cge.food.dtu.dk/services/SCCmecFinder/>). The genetic environment of the *ermT*, *cfr*, and *mecC* genes was illustrated in comparison with the reference strains using the EasyFig software.

Determination of the CRISPR‑Cas system of coagulase‑negative staphylococci

The CrisprCasFinder [\(https://crisprcas.i2bc.paris-saclay.](https://crisprcas.i2bc.paris-saclay.fr/) [fr/\)](https://crisprcas.i2bc.paris-saclay.fr/) was used to identify the numbers of CRISPR, Cas proteins, and spacers of all the MDR-CoNS (Couvin et al. [2018](#page-20-15)). Specifcally, the size of the fanking region and other parameters were set to default values. Moreover, three CoNS strains that contained larger sequences than CrisprCasFinder could handle were analyzed by the CRIS-PRCasMeta ([https://crisprcas.i2bc.paris-saclay.fr/Crisp](https://crisprcas.i2bc.paris-saclay.fr/CrisprCasMeta/Index) [rCasMeta/Index\)](https://crisprcas.i2bc.paris-saclay.fr/CrisprCasMeta/Index) applying all the default settings.

Genome availability

All the raw genome reads generated from this study have been deposited at the European Nucleotide Archive under Study Accession no. PRJNA1023081.

Statistical analysis

Data generated from this study reported frequencies and were presented in tables. Univariate logistic regression was to compute the odd ratio (OR) at a 95% confdence interval (95%CI) between the presence of MDR-CoNS genomes, and various mobilome with the ecological niches. Significant association at $p < 0.05$ was considered.

Results and discussion

CoNS have long been considered reservoirs of ARGs; however, very few genomic studies have elucidated the infuence of diferent ecological niches on the levels of ARGs and their MGEs. Moreover, there is a paucity of phylogenomic data on the transmission pathways of CoNS species and their ARGs between humans and animals.

Resistome, mobilome, and relatedness of the 26 CoNS analyzed in this study

The phenotypes of resistance of the 26 CoNS isolates characterized in this study are shown in Supplementary Table S1, and their resistome, virulome, genetic lineages, and mobile genetic elements are represented in Table [1.](#page-4-0) As identifed, all the isolates presented an MDR phenotype to 4 to 9 classes of antimicrobial agents. In this regard, the CoNS isolates with the least were those from nestling storks and with the highest those from pigs and pig farmers (Table [1\)](#page-4-0). The mechanisms of resistance to most of the antibiotics were mediated by combinations of multiple antibiotic resistance genes (ARGs).

Concerning the genetic lineages of *S. epidermidis*, STs belonging to the clonal complexes CC2 and CC5 were identifed. For the *S. haemolyticus,* the three isolates were of the lineage ST30 and ST68. Moreover, *S. sciuri*-ST212 and *S. hominis*-ST33 were detected. The genetic lineages of other species were not identifed as no MLST scheme has been developed and validated for them yet.

Multiple virulence genes that mediate host immune evasion, adhesion, and haemolysis among others were identifed (Table [1\)](#page-4-0). It is important to remark on the detection of an *S. epidermidis* strain that carries the enterotoxin genes, *sec* and *sel.*

Relatedness of the coagulase‑negative staphylococci strains

The phylogenetic analysis identifed clusters of related strains of various CoNS species with other countries. Specifically, the *cfr*-carrying *S. epidermidis-*ST16 strain (X5485) was related to an *S. epidermidis-*ST16 strain from a human blood sample $(SNP = 70)$ from Canada (id-41749). The *S. epidermidis*-ST35 from a dog owner is related to an human strain from Portugal (id-43340) (SNP = 90). Moreover, the *S. epidermidis*-ST297 from a healthy human in our study is related to three human strains from Germany, the UK, and Switzerland (SNP<80**) (**Fig. [1\)](#page-10-0). Also, the *S. epidermidis*-ST173 strain (X9066) was related to an animal strain in Thai-land (id-44496) (SNP = 76) (Fig. [1](#page-10-0)). Furthermore, the *sec/sel* carrying-*S. epidermidis*-ST595 strain (X4430) is not related (> 3000 SNPs) to previously described *sec/ sel*-carrying strains from Portugal and Italy (ID-43921, Id-43401). It is important to remark that despite the few SNP differences $(< 85$ SNPs) between some strains from Portugal (id-43340) and Canada (id-41749) with our two linezolid-resistant strains (X5485 and X6049b), none of them from the two countries was linezolid-resistant. This

The pigs (10 per farm) are named P1-P10 in each farm (A–D). In the case of humans working on the farm, they are designated as F1, F2, F3 and the farm (A–D)

The pigs (10 per farm) are named P1-P10 in each farm (A-D). In the case of humans working on the farm, they are designated as F1, F2, F3 and the farm (A-D) * All strains were of nasal origin, except S. haemolyticus X378

*All strains were of nasal origin, except *S. haemolyticus* X3784 of nestling stork which was from tracheal sample

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Fig. 1 Phylogenetic tree based on core genome SNP analysis of eight *S. epidermidis* isolates of this study with 31 publicly available genomes with similar lineages

suggests that our strains might have acquired the gene and mutation following antibiotic pressure in the livestock niche. These fndings highlight the international circulation of related *S. epidermidis* strains between humans and animals as confrmed by the phylogenetic analysis (SNP<100) (Fig. [1](#page-10-0)**,** Supplementary Table S2).

Aside from the *S. epidermidis* strains, we found related *S. borealis* (SNP<10) between pigs (Fig. [2](#page-11-0)**,** Supplementary Table S3**)**. However, the relatively high SNP (*n*=346) between the *S. saprophyticus* strains from a pig and pig farmer on the same farm suggests that they are unlikely related **(**Supplementary Table S4).

Mobilome‑bound antimicrobial resistance in coagulase‑negative staphylococci

Generally, MDR was the criteria of selection, and so, all isolates need to be resistant to at least 4 classes of antibiotics **(**Supplementary Table S1**)**. In this sense, the resistome profle of the strains was mainly to beta-lactam, macrolidelincosamide-streptogramin-B (MLS_B) , tetracyclines, sulfamethoxazole-trimethoprim, aminoglycosides, and phenicols as previously detected by PCR and presented in our previous study (Abdullahi et al. [2023a\)](#page-19-2). MDR to a maximum of 5 antibiotic classes was found from the previous study on CoNS strains from nestling storks (NS) (Abdullahi et al. [2023a](#page-19-2)). For the healthy dogs (H–D), healthy dog owners (H-DO), pigs (H-P), and pig farmers (H-PF), resistance to a maximum of nine antibiotic classes was obtained. In the case of isolates of healthy humans without animal contact (HH−) resistance to a maximum of seven antibiotic classes were obtained. Resistance genes already detected by PCR were found (Abdullahi et al. [2023a](#page-19-2), [b](#page-19-5), [c,](#page-19-6) [2024a,](#page-19-7) [b](#page-19-10), [c](#page-19-11)), but others not previously identifed such as *lsaB*, *lsaE*, *vgaA*(*LC*), *vga*(*E*), *bleO*, *str*, and *dfrC* were identifed (Table [1](#page-4-0)). Concerning plasmid bound-AMR genes, all the MRCoNS from pigs and pig farmers had *mecA* genes carried by SCC*mec* type-Vc except the two *S. saprophyticus* strains that had *mecA* in SCC*mec* type-IVb. The predominance of the SCC*mec* type-Vc in these isolates strongly suggests the interspecies transmission of *mecA* gene by the same SCC*mec* element. Thus, it has been speculated that the SCC*mec*

Fig. 2 SNIP-based phylogenetic tree of the four *S. borealis* isolates of this study mapped with all the 16 publicly available genomes

type-Vc in LA-MRSA originated from MR-CoNS carried in the same ecological niche (such as nostrils in this case) (Matuszewska et al. [2022](#page-21-14)). This is corroborated by the SCC*mec* types carried by LA-MRSA-CC398 isolates from the same pigs and pig farmers (Abdullahi et al. [2024b\)](#page-19-10). Whereas the SCC*mec* type-IV (a common community SCC*mec* type in MRSA) in *S. saprophyticus* from the pig and pig farmer suggests community-associated strains brought to the pig farm. Moreover, *S. saprophyticus* is known to cause uncomplicated urinary tract infections in the community (Lawal et al. [2021a,](#page-20-6) [b\)](#page-20-7). In nestling storks, the MDR-*S. arlettae* and *S. epidermidis* isolates were methicillin-susceptible, whereas the MR-*S. haemolyticus* carried *mecA* gene located in SCC*mec* type-V. Moreover, the MR-*S. lentus* carried the *mecA/ mecC* genes located in SCC*mec-mecC* hybrid. It is important to remark that the *mecA* gene might be intrinsic in *S. lentus* (Saber et al. [2017](#page-21-15)). Of the MR-CoNS strains from dogs/ owners and healthy humans, both the classical hospital and community-associated SCC*mec* elements were detected. This shows the SCC*mec* type in these hosts aside from pigs/ farmers has no categorical predilection.

Concerning genes that encode MLS_B resistance, the *ermB*, *ermC*, *erm45*, *vgaA*(*LC*), and *vga*(*E*) genes were identifed in single or in combination among over 50% of the CoNS isolates (Table [1\)](#page-4-0). Specifcally, the *ermC* gene in most of the *ermC*-positive strains was located in small plasmids that were 99.8% identical to those previously described in an *S. aureus* isolate, plasmid pMSA16 (GenBank accession number: JQ246438.1) and in an *S. saprophyticus* isolate, pSES22 (GenBank accession number: AM159501.1). Moreover, it is important to remark on the detection of the unusual *ermT* gene in two staphylococcal species: *S. borealis* (carried by plasmid *repUS18*) and *S. hyicus* (with no associated plasmid). The $ermT$ gene is not a common mechanism for MLS_B resistance in CoNS. It appears *ermT* gene is silently evolving in CoNS causing a constitutive MLS_B resistance phenotype.

Tetracycline resistance was found in all the pigs' and pig farmers' isolates and mediated by diferent combinations of genes. In this regard, *tet*(K), *tet*(L), *tet*(M), and *tet*(45) were found in most of the pigs/pig farmers isolates **(**Table [1](#page-4-0)**)**. Moreover, the *tet*(L) gene was found in one *S. epidermidis* (X6293) isolate from a dog owner. It is important to mention that the *tet*(L) gene was located in plasmid *rep22* in all the pigs and farmers isolates. However, no MGE was detected to be associated with all the *tet*(M) and *tet*(45) carrying MDR-CoNS strains. The absence of MGE associated with *tet*(M) gene in the MDR-CoNS is diferent from the transposon-linked *tet*(M) gene found in the *S. aureus* strains (Abdullahi et al. [2024b\)](#page-19-10), and this is subject to further investigations to unravel the reasons for the diferences. Perhaps, this plasmid *rep22* located *tet*(L) gene is coincidentally predominant in the pig farm niche. The *tet*(K) gene in most of the CoNS isolates was located in *rep7a* while in only one strain (X6049b) was located in plasmid *rep20,* and three others from pigs were not associated with this plasmid replicon (Table [1\)](#page-4-0). It is important to highlight that all the plasmid bound-*tet*(L) genes were linked with the *dfrK* gene in similar plasmid *rep*US12. A similar observation was reported in an MRSA-CC398 strain from a pig (GenBank accession number: FM207105). However, *tet*(L) was not found to be located in any plasmid in one of the *S. hyicus* strains from a pig (X5069) carrying a Tn*559*-bound *dfrK.* This denotes the diference in the pattern of acquisition of *tet*(L) gene and potential inter-species transmission in CoNS and *S. aureus* in a pig farm setting.

Aside from these plasmid-bound AMR genes, other genes that mediate resistance to aminoglycosides (such as *ant4′* and *bleO*, located in plasmid *repUS12*), clindamycin (e.g., *lnuA*, in *rep22*), and sulfamethoxazole-trimethoprim (e.g., *dfrK*, in *repUS12* and *rep22*) were occasionally identifed. In some instances, these AMR genes were not associated with any plasmid. We cannot categorically infer the reason some AMR genes are located in plasmids in some CoNS strains while in the bacterial chromosome of others. It could be that the bacteria lost the plasmids during horizontal transfer but the recipient bacteria retained the AMR genes (Dimitriu [2022](#page-20-16)). The similarity in plasmids that carry many AMR genes in all the CoNS strains demonstrates their impact on bacterial ftness for survival and capability to transfer these resistant genes intra-species (the same species), interspecies, and between diferent hosts. The transferability of AMR genes between diferent *Staphylococcus* species has been strongly suggested by the sequence similarity of their associated mobilome, especially plasmids (Souza-Silva et al. [2022](#page-21-4)). Moreover, some plasmids appeared to carry multiple AMR genes from diferent classes of antibiotics (such as *repUS12* and *rep22*).

Aside from these mobilome-bound AMR genes, the aminoglycoside and MLS_B resistance genes *ant9'* and *ermA* were also carried by Tn*554* in an *S. epidermidis* strain from a dog owner (X3617). Similar fndings (i.e., Tn*544*-linked *ant9′* and *ermA* genes) was reported but in a diferent CoNS species, *S. lugdunensis* (Chang et al. [2019,](#page-20-17) [2021\)](#page-20-18). This suggests potential inter-staphylococcal species transmission of the ARGs. Chloramphenicol resistance is an important phenotypic marker for linezolid resistance, especially in pig farm settings. Chloramphenicol has long been prohibited for the treatment of animal and human infections in Spain. However, forfenicol is still used for livestock. The *fexA* and *fexB* genes confer resistance to both forfenicol and chloramphenicol and could be responsible for the frequent co-resistance to chloramphenicol found in CoNS strains from pigs and pig farmers. In this study, only *fexA* which was carried by *Tn554* and *Tn558* was identifed in four pigs' strains and this illustrates the infuence of pig farm setting on the persistence of phenicol resistance genes especially the *fexA* that could be carried by two diferent transposons. Of clinical and public health concern is that other critical AMR genes such as those that mediate transferable linezolid resistance could be co-selected. In this regard, two *cfr*-carrying *S. epidermidis* and *S. saprophyticus* isolates from a pig previously identifed were identifed (Abdullahi et al. [2023b\)](#page-19-5). Upon genomic characterization, the *cfr* gene in *S. saprophyticus* strain was located in a plasmid *rep15*, while in *S. epidermidis* was not associated with any plasmid but was flanked by ISSau9 (Table [1](#page-4-0)).

Antimicrobial resistance mediated by chromosomal point mutations

Twelve of the 26 CoNS isolates analyzed (46.2%) carried one or more mechanisms of ciprofoxacin resistance mediated by DNA topoisomerase IV point mutations at GyrA (S84L) and DNA gyrase at GrlA (S80F) **(**Table [1](#page-4-0)**)**. Interestingly was the detection of 21 non-synchronous mutations on the GyrA on one *S. simulans* strain from a healthy pig (X5777) **(**Table [1](#page-4-0)**)**. A major diference in the ciprofoxacin resistance rate was observed between the isolates from the pigs and pig farmers and those of the other hosts: 7 (50%) of the CoNS isolates from pigs and pig farmers showed one or more of the mutations on quinolone-resistance-determining region, whereas three CoNS isolates from healthy humans (*S. epidermidis* and *S. haemolyticus*) and one *S. arlettae* isolate from a nestling stork exhibited this mutation **(**Table [1](#page-4-0)**)**. These highlight the infuence of pig farm antibiotic pressure on ciprofoxacin resistance on the CoNS isolates. Moreover, mutation-mediated AMR related to linezolid resistance was found in ribosomal proteins L3, L4, and L22 in a *S. epidermidis*-ST15 strain from a dog owner, as previously identifed by PCR-sequencing (Abdullahi et al. [2023c\)](#page-19-6).

Plasmid‑bound biocide and metal resistance among the CoNS isolates

Concerning biocide resistance, various plasmid bound-biocide resistance genes (such as *qacA* [*rep20*, *rep22*], *qacC* [*rep13*], *qacG* [*rep21*], and *qacJ* [*rep21*]) were detected in 34.6% of the 26 MDR-CoNS isolates characterized in this study. The acquisition of *qacG* gene carried on plasmid *rep*21 was previously found in the majority of *S. aureus* strains from our previous study (Abdullahi et al. [2024b](#page-19-10)). This plasmid-bound resistance to quaternary ammonium compounds could facilitate the persistence and co-selection of MDR in CoNS, as these genes make it very difficult for their eradication (Seier-Petersen et al. [2015](#page-21-16)). In addition, *smr* gene that encodes resistance against cationic antiseptic compounds (Damavandi et al. [2017\)](#page-20-19) was identifed in four strains (Table [1\)](#page-4-0). Two genes that encode for cadmium and zinc resistance (*cadD* and *czrC,* respectively*)* were identifed, of which *czrC* predominated (42.3%).

Metal resistance has previously been hypothesized to coselect for AMR and they are often linked to SCC*mec* elements (Lawal et al. [2021b\)](#page-20-7) and plasmids in LA-MRSA, *S. epidermidis*, *S. saprophyticus*, *S. haemolyticus*, etc. (Lawal et al. [2021a;](#page-20-6) Argudín and Butaye [2016;](#page-19-12) Schijfelen et al. [2010](#page-21-17)). Specifcally, determinants of copper (*copA*) and zinc (*czrC*) resistance were widespread among our MR-CoNS isolates of the pigs and pig farmers, but absent or minimal in other hosts. This could denote the potential selection of resistance to these

metals due to their persistence in pig farm settings (e.g., in pig feed) especially when plasmid-linked (Huang et al. [2021](#page-20-20); Sliferz et al. [2015\)](#page-21-18). Moreover, the cadmium resistance gene (*cadD*) suggests the involvement of environmental pollution where these staphylococci originated (Rebelo et al. [2021\)](#page-21-19).

Genetic environment of the unusual antimicrobial resistance gene in CoNS strains

The in silico analysis of the *ermT* sequences of three CoNS strains of two diferent species (*S. borealis* and *S. hyicus*) from healthy pigs revealed major diferences in their genetic environment (Fig. [3\)](#page-13-0). The *ermT* gene is in the opposite direction respect to a*nt9′* and both are located in plasmid *repUS18* in *S. borealis* strain. However, the *ermT* gene in the other two *S. hyicus* strains (X5447 and X5069) is not associated with any plasmid, perhaps it is chromosomally located. The *ermT* gene in the three strains produces an erythromycin-clindamycin resistance phenotype of constitutive character and highlights their evolution in MLS_B resistance among CoNS.

The in silico analysis of *S. lentus* strain (X4638) showed that it carried a hybrid SCC*mec-mecC*, which is 100% similar to an *S. sciuri* strain from bovine infection in the UK (Harrison et al. [2014](#page-20-21)). Specifcally, the SCC*mec*-*mecC* hybrid consisted of a class C1 *mec* complex located immediately downstream of a SCC*mec* type-VII element. Moreover, the *cadA*, *cadC*, and *cadD* genes are included in the system **(**Fig. [4](#page-14-0)**).** It has been previously described that most CoNS that carry the *mecC* gene are within a hybrid SCC*mec* element comprised of *mecA* included in SCC*mec* type VII and a *mecC* region consisting of the class E *mec* complex (de Moura et al. [2023](#page-20-22); Belhout et al. [2023;](#page-20-23) Paterson [2020\)](#page-21-20). However, *blaZ-*SCC*mec* XI was initially found to be associated with *mecC* in our *S. lentus* X4630 strain by

Fig. 3 Genetic environment of *ermT* gene of three CoNS isolates of this study (X5447, X5069, and X5409) in comparison with those of four reference strains

Fig. 4 The environment of the *mecC* gene of *S. lentus* (X4638) in comparison with previously described *mecC*-carrying coagulase-negative staphylococci and the *S. aureus*_{LGA251} strain

PCR and amplicon sequencing by Sanger (Abdullahi et al. [2023a](#page-19-2)). Following WGS, the *mecC* gene of the *S. lentus* X4638 strain was noted to be quite diferent from the classical SCC*mec* type XI that was frst demonstrated in *S. aureus*LGA251 (accession number FR821779). The reason for this variation is subject of further analysis. But, it could be hypothesized that a recombination event took place between the SCC*mec* type III (intrinsic for most MR-*S. lentus*) of the *mecA* gene and SCC*mec* type XI of the *mecC* to produce the SCC*mec-mecC* hybrid (i.e., the SCC*mec* type VII). In this regard, there is a need for caution in the use of PCR-based assays for the detection of SCC*mec* types in *mecC*-carrying non-*aureus* staphylococci. To the best of our knowledge, this report represents the frst description of a *mecC* in an *S. lentus* strain from a wild bird. This suggests the expansion of this mechanism of methicillin resistance in CoNS across various ecological niches including wild animals, which were previously proposed to be the major reservoirs of the *mecC* gene in *S. aureus* (Abdullahi et al. [2021](#page-19-4)).

The linezolid resistance mechanism mediated by plasmid pURX4944 (41.6 Kb) (Fig. [5](#page-15-0)) carrying the *cfr* gene located upstream of *lsaB* was identifed in *S. saprophyticus* X4944 strain and it was 96% identical to the plasmid of a clinical *S. epidermidis* strain from Italy (GenBank accession number: KR230047.1). Nevertheless, the *cfr* gene of our *S. epidermidis*-ST16 strain was not associated with a plasmid but was fanked by IS256 upstream of *lsaB* (Fig. [6](#page-15-1))*.* It has been suggested that the emergence and dissemination of the *cfr* gene in animals that have never used any of the oxazolidinones might be due to the selective pressure by the high use of forfenicols, lincosamides, tetracyclines, and pleuromutilins in the livestock sector (Gostev et al. [2021\)](#page-20-10).

Fig. 5 Circular representation of the plasmid-carrying the *cfr* gene in *S. saprophyticus*

Fig. 6 Environment of the *cfr* gene of *S. epidermidis* (X5485) and *S. saprophyticus* (X4944) in comparison with previously described *cfr*-carrying coagulase-negative staphylococci and *S. aureus*

*All strains were of nasal origin, except *S. haemolyticus* X3784 of nestling stork which was from a tracheal sample

All strains were of nasal origin, except S. haemolyticus X3784 of nestling stork which was from a tracheal sample

International Microbiology

Virulome profle of the coagulase‑negative staphylococci strains

We investigated the frequency and distribution of virulence genes among the diferent CoNS isolates from the four hosts. About 96.1% of the MDR-CoNS strains harbored one or more of diverse adherence, exoenzymes, haemolysin, or immune evasion genes (Table [2\)](#page-16-0). Enterotoxins constitute important virulence determinants of the genus *Staphylo coccus*, of which they are rarely detected in CoNS (França et al. [2021](#page-20-1)). Enterotoxins are the most implicated in foodborne gastroenteritis (Grispoldi et al. [2021](#page-20-24)). Moreover, other virulence factors could be responsible for a range of staphylococcal-related infections that are rarely detected in non-*aureus* staphylococci (Nanoukon et al. [2018](#page-21-21)). However, it is important to highlight the detection of a *sec*- and *sel* carrying *S. epidermidis* strain of the lineage ST595. Similar studies have previously reported these virulence genes and their associated pathogenicity islands in *S. epidermidis* (Lin et al. [2021;](#page-20-25) Nasaj et al. [2020;](#page-21-22) Banaszkiewicz et al. [2019](#page-19-13)). Moreover, it has been suggested that only *S. epidermidis* from animals or food but not from humans may typically produce *S. aureus-*related enterotoxins (Podkowik et al. [2016](#page-21-23); Veras et al. [2008;](#page-22-2) Stach et al. [2015](#page-21-24); Nanoukon et al. [2018](#page-21-21)). However, some *sec* and *sel* genes have been identi fed in association with plasmids, phages, and pathogenicity islands. Thus, they can be horizontally transmitted between any host, including humans. It appears that the *sec* and *sel* carrying *S. epidermidis* from nestling stork are not trans ferable as they were not associated with a mobile genetic element. Moreover, simultaneous colonization of the nostril by several *Staphylococcus* spp could promote the transfer of enterotoxin genes from *S. aureus* to commensal *S. epidermidis* (Nanoukon et al. [2018](#page-21-21)).

Aside from the toxins, many CoNS harbored genes such as the *capB* and *capC* (encode capsules) and *adsA*, *galE*, *wbtE*, *wbtP* genes that facilitate immune evasion by CoNS (Naushad et al. [2019;](#page-21-25) Li et al. [2015](#page-20-26)). Furthermore, the *icaABC* operon and its *icaR* were present in fve strains (19.2%) (Table [2](#page-16-0)). This denotes that some of the CoNS spe cies could easily adhere to the mucosa and inanimate sur faces and serve as a fundamental step in colonization and persistence on environmental surfaces and fomites (Idrees et al. [2021](#page-20-27)).

CRISPR‑Cas system distribution among the coagulase‑negative staphylococci

Complete CRISPR-Cas system was detected in 19.2% of the CoNS strains, of which *cas-*1, -2, and -9 predominated in *S. borealis* (75%). In other species, Cas3-type I CRISPR was identifed in two *S. epidermidis* strains (X6590 and X6049b) from humans. Furthermore, the *mecC*-carrying *S.*

a *NS* nestling stork, *H-P* healthy pig, *H-PF* healthy pig farmer, *H–D* healthy dog, *H-DO* healthy dog owner, *HH*− healthy human without animal contact

The pigs (10 per farm) are named P1-P10 in each farm (A–D). In the case of humans working on the farm, they are designated as F1, F2, F3 and the farm (A–D)

* All strains were of nasal origin, except *S. haemolyticus* X3784 of nestling stork which was from a tracheal sample

lentus harbored Cas2-type I and Cas9-type II (Table [3](#page-18-0)). The low frequency of CRISPR-Cas positive strains identifed in our study is closely similar to the 12.3% rate by Rossi et al. [\(2017](#page-21-26)) which consisted of mainly class 1 type IIIA and class 2 type IIC systems. Considering that most CRISPR-Cas reduces or eliminates mobile genetic elements such as plasmids, the low frequency of CRISPR-Cas elements among MDR-CoNS isolates in this study could explain the reason why all the strains had ARGs carried by multiple plasmids. However, large-scale genome-based studies including isolates with diferent profles of antibiotic resistance are necessary to better understand the roles of the CRISPR-Cas system on AMR genes and their plasmids among *S. borealis*.

Conclusion

These findings showed that various healthy ecological niches harbor CoNS that presented MDR phenotype mediated by multiple ARGs with several mobile genetic elements with relatively low frequency of intact CRISPR-Cas systems. Furthermore, our fndings highlight the potential geographical dissemination of some lineages of CoNS species across various hosts. Collectively, our fndings underscore the need to strengthen the genomic epidemiological approach and inclusion of MDR-CoNS from all hosts to adequately control the global fght against AMR and potentially pathogenic ones as identifed in the *sec-* and *sel*-carrying *S. epidermidis.*

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Author contribution I.N.A. and C.T performed the conceptualization of the study; I.N.A., M.S. and C.T. revised the methodlogy; I.N.A. carried out the laboratory experiments; I.N.A. and M.S. performed the sofware anaysis; C.T., I.N.A., J-L-F., M.S., M.Z., C.L. validated results; I.N.A., C.T., R.S., J.L-F., and M.S. did formal analysis; C.T., I.N.A., M.S., J-L-F., M.Z., and C.L. performed data curation; I.N.A. did writing—original draft preparation; C.T., I.N.A., J-L-F., M.S., M.Z., C.L. performed writing—review and editing; C.T. and C.L carried out the supervision of the study; C.T. did project administration; C.T., M.Z.,and I.N.A. funding acquisition; All authors have revised and agreed to the published version of the manuscript.

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Data availability No datasets were generated or analysed during the current study.

Declarations

Conflict of interest The authors declare no competing interests.

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