An observational cross-sectional study of nasal staphylococcal species of medical students of diverse geographical origin, prior to healthcare exposure. Prevalence of SCCmec, fusC, fusB and the arginine catabolite mobile element (ACME) in the absence of selective antibiotic pressure.

Paulo Eduardo Budri, Anna C. Shore, David C. Coleman, Peter M. Kinnevey, Hilary Humphreys, Deirdre Fitzgerald-Hughes.

1Department of Clinical Microbiology, Royal College of Surgeons in Ireland, Education and Research Centre, Beaumont Hospital, Dublin 9, Ireland.
2Microbiology Research Unit, Division of Oral Biosciences, Dublin Dental University Hospital, University of Dublin, Trinity College, Dublin 2, Ireland
3Microbiology Department, Beaumont Hospital, Dublin 9, Ireland

Keywords; Staphylococcus aureus, Coagulase-negative staphylococci, Nasal colonization, Healthy human nares, Antimicrobial resistance, MRSA.

Running title: Antimicrobial resistance and virulence genes among staphylococcal carriage isolates.

*Corresponding author: Paulo Eduardo Budri, Department of Clinical Microbiology, RCSI Education and Research Centre, Smurfit Building, Beaumont Hospital, Dublin 9, Ireland. Tel. +353 1 8093711, Fax +353 1 8092871, Email paulobudri@rcsi.com
Abstract

Objective: The aim of this study was to investigate co-located nasal Staphylococcus aureus and coagulase-negative staphylococci (CoNS) (mainly Staphylococcus epidermidis), recovered from healthy medical students in their pre-clinical year, prior to exposure to the healthcare environment, for the carriage of genes and genetic elements common to both species and that may contribute to S. aureus and methicillin-resistant S. aureus (MRSA) evolution.

Design: Prospective observational cross-sectional study. Carriage of antimicrobial resistance and virulence-associated genes in the absence of significant antibiotic selective pressure was investigated among healthy medical students from geographically diverse origins who were nasally co-colonised with S. aureus and CoNS. Clonal lineages of S. aureus isolates were determined.

Setting/Participants: Dublin-based international undergraduate medical students

Results: Nasal S. aureus carriage was identified in 137/444 (30.8%) students of whom nine (6.6%) carried MRSA (ST59-MRSA-IV (6/9), CC1-MRSA-V-SCCFus (3/9)). The genes mecA, fusB, ileS2, qacA/qacC and the arginine catabolite mobile element (ACME)-arc were detected among colonizing nasal staphylococci and had a significantly greater association with CoNS than S. aureus. The rate of co-carriage of any of these genes in S. aureus/CoNS pairs recovered from the same individual was <1 %.

Conclusions The relatively high prevalence of these genes among CoNS of the healthy human flora in the absence of significant antibiotic selective pressure is of interest. Further research is required to determine what factors are involved and whether these are modifiable to help prevent the emergence and spread of antibiotic resistance amongst staphylococci.
Strengths and limitations of this study:

- Global evaluation of antibiotic resistance gene carriage among Staphylococci among healthy medical students in preclinical years through DNA microarray analyses.
- Pairs of staphylococcal species were isolated from the same colonisation site (nares) of multiple participants to allow investigation of shared antibiotic resistance and virulence in the same human niche in a community setting.
- A single centre study design.
- CoNS was investigated only in students co-colonised with S. aureus.
- The study design did not facilitate follow-up of this cohort during clinical training.

Introduction

*Staphylococcus aureus* and *Staphylococcus epidermidis* are significant colonisers of healthy human skin and nares and are among the leading causes of healthcare-associated infection (HAI). Morbidity, mortality and the financial burden associated with methicillin-resistant *S. aureus* (MRSA) infections are well documented. Furthermore, coagulase-negative staphylococci (CoNS) including *S. epidermidis* are reported reservoirs of antimicrobial-resistance genes and their associated mobile genetic elements, most notably the staphylococcal cassette chromosome (SCC) harbouring the *mec* gene (SCC*mec*) \(^1\).

Twelve SCC*mec* types and numerous subtypes have been described among MRSA isolates to date. The more prevalent and diverse range of SCCs and SCC*mec* among CoNS further supports CoNS as a reservoir for antimicrobial resistance genes \(^1\). The identification of SCC, SCC*mec* and SCC-associated elements with other antimicrobial and virulence genes and their epidemiological relationships among clinical staphylococcci has advanced our understanding of the role of CoNS in the evolution of MRSA \(^2\). For example, the fusidic acid resistance gene *fusC* is associated with SCC*mec*
IV-SCC and other SCC-like elements have been identified in \textit{S. aureus}, MRSA and CoNS and may contribute to MRSA emergence in countries with significant fusidic acid usage.\textsuperscript{3-5} Furthermore, the SCC-like arginine catabolic mobile element (ACME) which enhances acid tolerance, is abundant among clinical CoNS isolates, in particular \textit{S. epidermidis} and \textit{S. haemolyticus}.\textsuperscript{6} Among \textit{S. aureus}, ACME has mainly been detected among isolates of the community-associated (CA) USA300 clone.\textsuperscript{7} CoNS are also a putative reservoir of the high level mupirocin resistance encoding gene \textit{ileS2}, which is also increasing among \textit{S. aureus}/MRSA in healthcare and community environments related to horizontal gene transfer or expansion of specific clones.\textsuperscript{8,9} Increasingly, MRSA clones previously associated with the community, such as clonal complex (CC) 1 are spreading to healthcare settings making the differentiation between healthcare-associated (HA) MRSA and CA-MRSA unclear.\textsuperscript{10} Therefore, detailed investigation of the genetic and phenotypic traits of colonizing staphylococcal species in community settings are important to identify those with features that may contribute to their evolution into potentially successful and formidable healthcare-associated clones.

The aim of this study was to investigate co-located nasal \textit{S. aureus} and CoNS (mainly \textit{S. epidermidis}) recovered from healthy medical students in their pre-clinical year, prior to exposure to the healthcare environment, for the carriage of genes and genetic elements common to both species and that may contribute to \textit{S. aureus} and MRSA evolution.

\textbf{Methods}

\textbf{Study setting, participants and sample collection}

This observational cross-sectional study was conducted at the Royal College of Surgeons in Ireland (RCSI) from December 2014 – January 2016. Nasal swabs (eSwab Copan\textsuperscript{®}, Italy) were collected anonymously from undergraduate medical students. Eligible students were those attending the RCSI medical centre to submit a swab for...
mandatory MRSA screening in the week before they began their clinical attachments. In total 444/450 eligible medical students (250 (56.3%) male, 194 (43.7%) female) participated in this study. All participants reported no previous hospital contact in the six weeks prior to recruitment. The student volunteers were from the second year of the undergraduate medical programme and as such, all participants were domiciled in Ireland for a minimum of two years prior to recruitment. Data was collected anonymously from each participant, including age range, region of origin and previous healthcare contact. Ethical approval (approval number REC949) was obtained from the Institute’s Ethics Committee and informed consent was obtained from each participant.

Sample preparation

Swabs were processed to recover *S. aureus* (including MRSA) and pathogenic CoNS species using a modification of a published method\(^\text{11}\). Swabs were enriched in brain heart infusion (BHI) supplemented with 6% (w/v) NaCl for 24 h at 37°C followed by further enrichment in mannitol salt broth for 24 h at 37°C. The enriched culture was diluted 1/1000 and 100 μl was spread onto SaSelect agar (Bio-Rad®, Hercules, CA, USA). Plates that yielded pink/orange colonies (presumptive *S. aureus*) were inspected for growth of colonies of relevant CoNS based on colony colour (e.g. light pink colonies of various sizes, presumptive *S. epidermidis*; white/yellow colonies, *S. haemolyticus*, *S. hominis*, *S. capitis*, *S. warneri*, *S. caprae*, *S. lugdunensis*). Presumptive CoNS species and *S. aureus* were sub-cultured from these plates onto Columbia blood agar (CBA) and identified by matrix-assisted laser desorption/ionization-time-of-flight mass spectrometry (MALDI-TOF) using a MALDI Biotyper (Microflex LT, Bruker). Matched isolates (where *S. aureus* and a CoNS species were recovered from the same swab) were cryopreserved and stored at -20°C (Protect™ bacterial preserver beads (Technical Service Consultants, UK)).
Characterisation of *S. aureus* and CoNS isolates

Genomic DNA from *S. aureus* and CoNS isolates was extracted using enzymatic lysis using the buffers and solutions provided with the *S. aureus* Genotyping Kit 2.0 (Alere Technologies GmbH, Jena, Germany) and a DNeasy® Blood and Tissue kit (Qiagen, Crawley, UK). Genetic characterisation of isolates was undertaken by DNA microarray profiling using the *S. aureus* Genotyping Kit 2.0 as described previously. The kit detects 333 gene targets including staphylococcal antimicrobial-resistance, virulence, SCC*mec* and ACME-arc genes and assigns *S. aureus* isolates to multilocus sequence type (ST) or clonal complexes (CC)s. MRSA phenotype was confirmed in *S. aureus* and CoNS isolates positive for *mecA* by growth of pink (*S. aureus*) or colorless/white (CoNS) colonies on MRSASelect agar (Bio-Rad®, Hercules, CA, USA). When required, confirmation of carriage of fusidic acid resistance genes *fusC*, *fusB*, and toxic shock syndrome toxin gene (*tst1*) were confirmed by PCR using the primers and conditions described by O’Neill *et al* and Chen *et al*.

Fusidic acid susceptibility testing

Fusidic acid MICs were determined by ETEST® (bioMérieux, Marcy-l’Etoile, France) according to manufacturer’s instructions. Thirty *S. aureus* and CoNS isolates harbouring *fusC* or *fusB* were sub-cultured twice before testing. Results were interpreted according to EUCAST (http://www.eucast.org, assessed May 2015) susceptibility criteria.

Statistical analyses

Fisher’s exact test was used to analyze categorical variables (prevalence of genes) using GraphPad QuickCalcs on-line software. The significance of differences between groups was expressed as two-tailed *p*-values, *p* values of ≤ 0.05 were considered statistically significant.
Patient and Public Involvement statement:

The participants in this study were medical undergraduate students. They were invited to participate in this study when they underwent mandatory MRSA screening prior to commencement of clinical placements. As future clinicians, participants were broadly considered in the development of the research question. As nasal carriage of *S. aureus* and MRSA among healthcare staff contributes to transmission to patients in healthcare environments, medical students have an interest in contributing to this knowledge.

Participants did not contribute to the study design. Participants were informed at recruitment that dissemination of the results would be through the scientific literature.

Results

Nasal carriage of staphylococcal species and regional distribution

Thirty-one percent (137/444) of students were positive for nasal carriage of *S. aureus* of whom 6.6 % (9/137) were MRSA. Eighty-seven percent (386/444) of students were positive for nasal carriage of CoNS (*S. epidermidis* 82% 364/444), *S. haemolyticus* (3% 14/444) or *S. saprophyticus* (2% 8/444) (Table 1). All students positive for *S. aureus* also carried *S. epidermidis*. Methicillin-resistant (MR)-CoNS were investigated in the *S. aureus*-positive cohort only of which 13.1 % (18/137) were MR-CoNS. One student exhibited co-carriage of MRSA and MR-CoNS. The geographical region of origin of students harbouring *S. aureus* and CoNS is shown in Figure 1. The Middle East, Europe and North America accounted for 68.6% of *S. aureus* carriers. For regions represented by ≥ 12 participants, the rate of nasal carriage of *S. aureus* varied geographically between 17 % (South East Asia) and 44 % (Africa).

Clonal lineages among *S. aureus* isolates
The ST or CC distribution among 137 *S. aureus* isolates is shown in Figure 2. Isolates belonged to a variety of CCs with 46/137 (33.5%) assigned to internationally disseminated CC5, CC8, CC22, CC30, CC45. A further 24/137 (17.5%) isolates belonged to CC1, CC59, CC88 or CC398.

**SCCmec types and fusidic acid resistance among *S. aureus* and CoNS**

Of the 333 staphylococcal genes detected by the microarray, the two most prevalent antibiotic resistance genes among nasal staphylococci were those encoding resistance to β-lactams and fusidic acid. The most common SCCmec type among nasal MRSA (n=9) and MR-CoNS (n=18) was SCCmec type IV (class B mec *mecA, DmecR1, ugpQ* and *ccrA-2, ccrB-2*). The nine MRSA isolates belonged to ST59-MRSA-IV (6/9) and CC1-MRSA-V-SCCfus (3/9) (class C *mec mecA, ugpQ* and *ccrC and fusC Q6GD50*) and cassette chromosome recombinase (*ccr A-1, ccrB-1*). Among 18 MR-CoNS identified (17 *S. epidermidis* and 1 *S. saprophyticus*), half harboured SCCmec type IV (8 *S. epidermidis* and the single *S. saprophyticus*). SCCmec types II, V and VII were identified in three, five and one of the remaining *S. epidermidis* isolates, respectively. Isolates from the one individual who exhibited nasal co-carriage of MRSA and MR-CoNS (*S. epidermidis*) both harboured SCCmec type IV (Table 2).

In addition to the three CC1-MRSA-V isolates that carried SCCfus, the fusidic acid resistance genes *fusC* and *fusB* were identified in 28/128 (21.8%) and 2/128 (1.5%) of methicillin-susceptible *S. aureus* (MSSA) isolates, respectively. Ten of the 28 *fusC*-positive MSSA isolates belonged to CC1-MSSA-SCCfus, 11 were CC88-MSSA and 7 CC8-MSSA. All 10 CC1-MSSA-SCCfus isolates harbour a combination of SCCfus with the *ccr* genes, *ccrA-1* and *ccrB-1*. The two *fusB* positive isolates belonged to CC5-MSSA and CC8-MSSA (Table 2). Among MR-CoNS, 27.7 % (5/18) *S. epidermidis* isolates carried *fusC* (two of them also carried *ccr* genes *ccrA-1 ccrB-1*) and 50 % *fusB* (9/18,
eight *S. epidermidis* and the one *S. saprophyticus*). Among methicillin susceptible CoNS isolates, the *fusC* and *fusB* genes were identified in 20/119 (16.8%, 18 *S. epidermidis* and two *S. saprophyticus*) and 18/119 (15.1%, all *S. epidermidis*), respectively. One participant had nasal co-carriage of *fusC*-positive *S. aureus* (CC88-MSSA) and CoNS (*S. epidermidis*). Among SCCmec positive staphylococci were confirmed to have an MRSA/MR-CoNS phenotype. However, there was poor correlation between *fusC/fusB* carriage and phenotypic fusidic acid resistance. Fusidic acid MICs for all *fusC* or *fusB*-positive *S. aureus* and CoNS isolates are shown in Table 3. Phenotypic fusidic acid resistance was confirmed (based on EUCAST breakpoints, MICs ≥1 µg/ml) in 23/32 (71.8%) *S. aureus* and 20/38 (52.6%) CoNS nasal isolates harbouring either *fusC* or *fusB* (DNA microarray result confirmed by PCR). Eight nasal isolates (three *S. aureus*, five *S. epidermidis*) positive for *fusB* exhibited high level fusidic acid resistance (MIC ≥ 32 µg/ml). Fusidic acid resistance was inducible in a further three *S. aureus* and seven *S. epidermidis* isolates following incubation with 0.01 µg/ml fusidic acid BHI agar.

Other notable antimicrobial resistance genes among nasal *S. aureus* and CoNS

Apart from SCCmec element and *fus* genes, other antimicrobial genes detected among staphylococcal nasal flora were identified by DNA microarray. Tetracycline resistance genes, *tet(K)* or *tet(M)*, were detected in 13/137 (9.5 %) of *S. aureus* isolates and 6/137 (4.3%) of the CoNS isolates. The quaternary ammonium compound resistance genes (*qacA/qacC*), encoding antiseptic resistance, were significantly more prevalent among CoNS isolates compared to *S. aureus* isolates (29/137 (21.2%) Vs 2/137 (1.4%), p<0.0001). Significantly more CoNS than *S. aureus* isolates carried *ileS2* encoding high-level mupirocin resistance (11/137 (8%) vs 1/137 (0.72%), p<0.01). However, none of these genes were common to *S. aureus*/CoNS pairs recovered from the same individual. The β-lactamase genes were abundant among *S. aureus* and CoNS; *blaZ* was present in 101/137 (73.72%) *S. aureus* isolates and 92/137 (67.1%) CoNS isolate and in 74/137
(54%) of individuals, these genes were common to *S. aureus*/CoNS pairs from the same nares. A summary of the antibiotic resistance genes found among *S. aureus* and CoNS is shown in Table 2. The staphylococcal isolates were negative for all other antibiotic resistance genes spotted on the microarray.

**Virulence genes among nasal *S. aureus* and CoNS**

A single isolate, CC30-MSSA, was positive for the Panton-Valentine leucocidin genes (*lukF/S-PV*). Among nasal staphylococci, ACME-arc was significantly associated with CoNS compared to *S. aureus* (44/137 (32.1%) Vs 1/137 (0.7%)), p<0.0001. The toxic shock syndrome toxin gene *tst1* was identified in 33/137 (24.1%) nasal *S. aureus* isolates. Unusually, DNA microarray identified *tst1* in two *S. epidermidis* isolates and this was confirmed by PCR. ACME-arc was common to *S. aureus*/CoNS recovered from the nares in one individual only. One hundred and two (74.4%) *S. aureus* isolates encoded one or more enterotoxin genes. The enterotoxin gene cluster (egt), containing *seg, sei, sem, sen, seo, seu* was the most prevalent (48/102, 47%) followed by *seq/k* (13/102, 12.7%) and *sec/l* (7/102, 6.8%). The staphylococcal isolates were negative for all other toxin genes spotted on the microarray.

**Discussion**

Studies of staphylococcal carriage and epidemiology among the healthy population in the absence of significant antibiotic pressure are important in identifying the potential for pathogenic evolution. To our knowledge, this is the first study to co-investigate CoNS and *S. aureus* when recovered together from the nares of healthy pre-clinical medical students. The species distribution of nasal colonizing CoNS was similar to other studies even though the enrichment methods used here favoured *S. aureus* and *S. epidermidis* and may explain the low prevalence of other CoNS species. Our study revealed that, apart from the *bla* genes, which are abundant among staphylococci, the
rates of co-carriage of antibiotic resistance genes in paired *S. aureus/CoNS* from the same individual were low in the community setting at <1%. Rates of simultaneous carriage of antimicrobial resistance among nasal staphylococci are likely to be higher under selective antibiotic pressure but few studies have investigated this among patients. One small study of hospitalized patients with nasal carriage of *S. aureus* and CoNS reported a rate of 12.5% patients carrying MRSA and MR-CoNS [17]. However, the authors reported only two cases where simultaneous carriage of MR-CoNS and MRSA was detected and the strains involved carried different SCCmec types. Despite negligible detection of co-species nasal carriage of these genes in medical students prior to healthcare exposure, based on antimicrobial resistance gene carriage by CoNS from this cohort, there is significant potential for mobilisation of genes to *S. aureus* that may enhance its pathogenic potential in the healthcare setting.

DNA microarray analyses revealed carriage of SCCmec, fusC, fusB, ileS2, qacA/qacC and ACME-arc among colonising nasal staphylococci in individuals with no previous healthcare exposure with greater prevalence among CoNS than *S. aureus*. This pattern among pre-clinical medical students, supports CoNS as a reservoir with potential to subsequently accelerate antimicrobial resistance and pathogenicity among colonizing *S. aureus* in clinical environments under antibiotic selective pressure [16, 18, 19].

Despite considerable geographical distribution of the participants in this study, a *S. aureus* nasal carriage rate in the community of 30.8% was recorded. In this study, CC30, CC88 and CC8 were the most prevalent clones identified among nasal *S. aureus*. CC30 is among the internationally disseminated clones in which SCCmec has been acquired and is a successful colonising lineage, reported among HA and CA-MRSA. Among medical students, these MSSA isolates may therefore represent a significant pool for the uptake of SCCmec in a clinical setting. CC88 is frequently isolated in Australia but in our study the geographical background of isolates was mixed (including Middle East, Europe, South East Asia and Central America). CC8 is associated with MRSA.
infection and is globally disseminated\textsuperscript{20}. Although CC30, CC88 and CC8 were prevalent among community MSSA isolates in this study, among the relatively few MRSA recovered, none belonged to these CCs. Two CC/ST types detected among MRSA recovered from healthy medical students in this study were ST59-MRSA-IV and CC1-MRSA-V-SCC\textit{fusC}. ST59 (Western Australian-MRSA-73) is a sporadic Australian strain and apart from PVL-negativity, is indistinguishable from USA1000\textsuperscript{21}. In this study the geographical background of these isolates was wide (Middle East, North America and South East Asia).

The identification of a significant reservoir of antibiotic resistance among medical students prior to healthcare exposure in subsequent clinical years, highlights the need for effective infection prevention and control policies in relation to hand hygiene and surveillance. In the absence of antibiotic selective pressure, the colonising MRSA rate appears relatively stable and in this study was 2\% (9/444), similar to rates reported elsewhere\textsuperscript{22}. However, a previous study among medical interns in China reported a nasal MRSA rate of 9.4\% likely reflecting exposure to the healthcare environment\textsuperscript{23}. One study reported an increasing in carriage rates of MR-CoNS from 14\% among medical student pre-internship, to 29.28\% among interns\textsuperscript{24}. Prevalence rates of MR-CoNS in recent community-based surveys are variable but rates of 16.5\%\textsuperscript{19} and 17.2\%\textsuperscript{25} are reported in similar cohorts to this study where, of those colonised with \textit{S. aureus}, 13.1\% carried MR-CoNS. SCC\textit{mec} type IV, the smallest of the SCC\textit{mec} elements, was the most prevalent type among MRSA and MR-CoNS here (66.6\% and 50\%). SCC\textit{mec} IV has been detected in approximately 40\% of methicillin-resistant \textit{S. epidermidis} identified in humans\textsuperscript{26}. However, in this study SCC\textit{mec} type V was also represented among MRSA and MR-CoNS. While only one individual was colonised with MRSA and MR-CoNS in this study (both SCC\textit{mec} type IV), the preponderance of SCC\textit{mec} IV element in nasal MRSA and MR-CoNS suggests the potential for \textit{mecA} gene transfer.
among these species even in the absence of selective pressure. The small size of this
element, which has a low fitness cost, may enhance its dissemination potential. 27

Fusidic acid resistance among *S. aureus* from healthy carriers in nine European
countries in 2014 was reported to be <10 % 28. However, we found a prevalence of 22.6%
of fusC/fusB genes among healthy carriers. Fusidic acid resistance appears to correlate
with increased use of this agent. For example, in New Zealand, where it is used as a
first-line empiric agent for topical treatment of impetigo, prevalence rates of resistance in
community *S. aureus* isolates increased from 17% in 1999 to 29% in 2013 29. In Europe,
fusidic acid is combined with β-lactams for the treatment of staphylococcal bacteraemia,
endocarditis, and osteomyelitis 30 and is used widely in the community for skin and soft
tissue infections (SSTIs). A 2010 study of fusidic acid resistance among *S. aureus* clinical
isolates showed Greece and Ireland to have the highest rates (52.5 and 19.9%) 31.

SCCfus has been identified in the CC1 background and more recently, in other lineages
such as ST239 and ST779 32-34. As highlighted here in the absence of significant
antibiotic pressure in the community, it appears that this element is associated with
MRSA and MSSA in the CC1 background. This genetic platform, particularly when
associated with SCCmec in a composite element (SCCmec V+SCCfus) may enable the
transfer of multi-drug resistance in a single transfer event. The use of fusidic acid is un-
regulated in some countries and hence it may be used inappropriately in a community
setting (for example in short or discontinuous doses). Inappropriate use of fusidic acid
may therefore favour co-selection of methicillin-resistance among *S. aureus*. In addition,
in this study, 14/18 (77%) of MR-CoNS were positive for fusC or fusB. This association
of resistances among the resident flora may provide further opportunity for dissemination
of MRSA driven by fusidic acid selective pressure. Interestingly a positive correlation
between carriage of fusC/fusB and phenotypic resistance was observed in only 71.9 %
and 53.6 % of *S. aureus* and CoNS respectively. However, induction of gene expression
with fusidic acid pre-incubation gave better correlation (82.2% and 76.3 % correlation).
There were limitations to this study, which included; a single centred, relatively small study. Some nasally abundant CoNS species, for example *Staphylococcus lugdunensis* and *Staphylococcus hominis*, were under-represented as the enrichment method favoured pathogenic staphylococci such as *S. aureus* and *S. epidermidis*. CoNS was investigated only in those co-colonised with *S. aureus* and therefore prevalence rates for genes among CoNS do not reflect the entire cohort. CCs and STs were determined only among *S. aureus* as the high rate of genetic recombination among CoNS makes strain typing unreliable. Although the microarray system used is reported effective for staphylococcal species other than *S. aureus*\(^{35}\), some gene targets may be heterologous among staphylococci leading to false negatives. The study design did not facilitate follow-up of this cohort during clinical training which may have revealed further changes in gene carriage among colonising staphylococci. However, the multi-national origin of the student body in our institution facilitated analysis of a relatively broad geographic cohort in a single study and emphasises the role that importation plays in *S. aureus* epidemiology. Unlike other studies of staphylococci in the healthy human nares, pairs of staphylococcal species originating from the same individual were investigated here for their resistance and virulence traits. These data support a low rate of transfer of antibiotic resistance between colonising staphylococcal species in the absence of healthcare contact. However, it is concerning that similar SCC\(_{mec}\) and SCC\(_{fusC}\) types, in addition to *ileS2, qacA/qacB* and ACME are carried among CoNS and *S. aureus* in healthy individuals who will have subsequent roles in healthcare provision. Given the increasing emergence of HA-MRSA with features of community strains, further mobilisation of these elements under selective antibiotic pressure may enhance the transmission and success of *S. aureus* in the healthcare environment.

**Funding**
PEB received funding for the study from Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES), Brazil under the Science without Borders Program. Grant number 9172-13-0.

Competing interests

HH has received funding from Pfizer and Astellas outside the relevance of the submitted work, all other authors report no competing interests.

Contributor Statement

PEB and DFH recruited students to the study, PEB conducted the laboratory work and drafted the manuscript. DFH and HH conceived of the study and contributed to study design. AS, PK and DC provided critical data interpretation and revised the drafted work. All authors contributed to the final approved draft.

Data sharing statement

All data for these analyses are included in the manuscript. No additional data are available.

Acknowledgements

We acknowledge the support of the Mercer’s medical centre, RCSI and Ms Helen Barry, Chief Medical Scientist, St. James’ Hospital Dublin.

Figure Legends

Figure 1. Geographical origin of medical students recruited. The geographical areas of origin of 444 medical students recruited to the study are shown (dark grey bars). Of those recruited, 137 were confirmed nasal S. aureus and CoNS positive. The proportion of recruited students from each geographical origin with nasal S. aureus carriage are also shown (light grey bars).
Figure 2. Genotypic diversity of 137 *S. aureus* nasal isolates using DNA microarray analysis, including 128 MSSA (dark grey bars) and 9 MRSA (light grey bars). Letter (a) indicates internationally disseminated clones into which SCCmec can integrate. CC=clonal complex.
Table 1. Staphylococcus species recovered from 444 nasal swabs.

<table>
<thead>
<tr>
<th>Staphylococcus species recovered</th>
<th>Total n, (%)</th>
<th>Methicillin-resistant phenotype n, (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n=444</td>
<td>n=137(^a)</td>
</tr>
<tr>
<td><strong>S. aureus</strong></td>
<td>137, (30.8)</td>
<td></td>
</tr>
<tr>
<td>MRSA</td>
<td>9, (6.6)</td>
<td></td>
</tr>
<tr>
<td><strong>S. epidermidis</strong></td>
<td>364, (81.9)</td>
<td></td>
</tr>
<tr>
<td>MRSE</td>
<td>17, (12.4)</td>
<td></td>
</tr>
<tr>
<td><strong>S. haemolyticus</strong></td>
<td>14, (3.1)</td>
<td></td>
</tr>
<tr>
<td>MRSH</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td><strong>S. saprophyticus</strong></td>
<td>8, (1.8)</td>
<td></td>
</tr>
<tr>
<td>MRSS</td>
<td>1, (0.72)</td>
<td></td>
</tr>
</tbody>
</table>

**Co-carryage species**

| **S. aureus** + **S. epidermidis**     | 137, (30.8)      |                                       |
| MRSA + MR-CoNS                         | 1, (0.72)        |                                       |
| *fusC* positive **S. aureus** + CoNS  | 1, (0.72)        |                                       |

\(^a\) CoNS were investigated only in those positive for nasal **S. aureus** in the student cohort and not in all those recruited. MRSA = methicillin resistant **S. aureus**, MRSE = methicillin resistant **S. epidermidis**, MRSH = methicillin resistant **S. haemolyticus**, MRSS = methicillin resistant **S. saprophyticus**
Table 2. Resistance/virulence genes detected among 137 co-located nasal S. aureus/CoNS pairs.

<p>| Detected Gene(s) | Phenotypic resistance/trait       | No. isolates positive | | | | No. S. aureus/CoNS pairs positive (n) |
|------------------|----------------------------------|-----------------------|------------------|------------------|-----------------------------|
|                  |                                  | n (%)                | S. aureus n=137  | CoNS n=137       | P value                     |
| <strong>Antibiotic resistance gene</strong> |                                  |                       |                  |                  |                             |
| <em>bla</em>Z*          | β-lactam                         | 101 (73.7)           | 92 (67.1)       | 0.289            | 74                          |
| <em>fusB</em>           | Fusidic acid                     | 2 (1.5)              | 27 (19.7)       | 0.0002*          | 0                            |
| <em>fusC</em>           | Fusidic acid                     | 31 (22.6)            | 25 (18.2)       | 0.159            | 1                            |
| <em>mecA</em>           | Methicillin                      | 9 (6.5)              | 18 (13.1)       | 0.103            | 1                            |
| <em>ileS2</em>          | Mupirocin                        | 1 (0.7)              | 11 (8.0)        | 0.005*           | 1                            |
| <em>qacA and qacC</em>  | Quartenary ammonium salts        | 3 (2.2)              | 29 (21.2)       | &lt;0.0001*         | 0                            |
| <em>tet(K) and tet(M)</em> | Tetracycline                   | 13 (9.5)             | 6 (4.4)         | 0.152            | 0                            |
| <em>erm(C)</em>         | Macrolide/lincosamide           | 6 (4.3)              | 5 (3.6)         | 1.000            | 0                            |
| <em>msr(A)</em>         | Macrolide                        | 2 (1.45)             | 15 (10.9)       | 0.002*           | 1                            |
| <em>mph(C)</em>         | Macrolide                        | 0                    | 15 (10.9)       | &lt;0.0001*         | 0                            |
| <em>dfrS1</em>          | Trimethoprim                     | 0                    | 19 (13.8)       | &lt;0.0001*         | 0                            |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>vga</strong></td>
<td>Streptogramin A</td>
<td>1 (0.7)</td>
<td>6 (4.3)</td>
<td>0.120</td>
</tr>
<tr>
<td><strong>Virulence</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ACME-arc</td>
<td>pH tolerance</td>
<td>1 (0.7)</td>
<td>44 (32.1)</td>
<td>&lt;0.0001*</td>
</tr>
<tr>
<td><em>tst</em>&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Toxic shock toxin</td>
<td>33 (24.1)</td>
<td>2 (1.5)</td>
<td>&lt;0.0001*</td>
</tr>
</tbody>
</table>

<sup>a</sup>associated with SCC element, (ccrA-1 and ccrB-1) in 13/137 *S. aureus.*<sup>b</sup>*tst* confirmed by PCR. ACME = Arginine Catabolite Mobile Element. * indicates a statistically significant result by Fisher’s exact test.
Table 3. Fusidic acid MICs for *S. aureus* and CoNS

<table>
<thead>
<tr>
<th>MIC Interpretation*a</th>
<th>S n (%)</th>
<th>R n(%)</th>
<th>HR n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>S. aureus (n = 32)</strong></td>
<td>9 (28.1)</td>
<td>20 (62.5)</td>
<td>3 (9.3)</td>
</tr>
<tr>
<td><strong>CoNS (n = 38)</strong></td>
<td>18 (47.4)</td>
<td>15 (39.5)</td>
<td>5 (13.2)</td>
</tr>
</tbody>
</table>

References


21


Genotypic diversity of *S. aureus* isolates

![Diagram showing genotypic diversity of *S. aureus* isolates with bars representing the number of isolates for each clonal complex/ST, distinguishing between MSSA and MRSA.](image)