A mecC allotype, mecC3, in the CoNS Staphylococcus caeli, encoded within a variant SCCmecC

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Background: Methicillin resistance in staphylococci is conferred by an alternative PBP (PBP2a/2') with low affinity for most β -lactam antibiotics. PBP2a is encoded by *mecA*, which is carried on a mobile genetic element known as SCC*mec*. A variant of *mecA*, *mecC*, was described in 2011 and has been found in *Staphylococcus aureus* from humans and a wide range of animal species as well as a small number of other staphylococcal species from animals.

Objectives: We characterized a novel *mecC* allotype, *mecC3*, encoded by an environmental isolate of *Staphylococcus caeli* cultured from air sampling of a commercial rabbit holding.

Methods: The *S. caeli* isolate 82B^T was collected in Italy in 2013 and genome sequenced using MiSeq technology. This allowed the assembly and comparative genomic study of the novel SCC*mec* region encoding *mecC3*.

Results: The study isolate encodes a novel *mecA* allotype, *mecC3*, with 92% nucleotide identity to *mecC. mecC3* is encoded within a novel SCC*mec* element distinct from those previously associated with *mecC*, including a *ccrAB* pairing (*ccrA5B3*) not previously linked to *mecC*.

Conclusions: This is the first description of the novel *mecC* allotype *mecC3*, the first isolation of a *mecC*-positive *Staphylococcus* in Italy and the first report of *mecC* in *S. caeli*. Furthermore, the SCC*mec* element described here is highly dissimilar to the archetypal SCC*mec* XI encoding *mecC* in *S. aureus* and to elements encoding *mecC* in other staphylococci. Our report highlights the diversity of *mecC* allotypes and the diverse staphylococcal species, ecological settings and genomic context in which *mecC* may be found.

Introduction

Methicillin resistance in *Staphylococcus aureus* is typically conferred by the gene *mecA* along with two variants, *mecB* and *mecC*.¹⁻⁴ *mec* gene resistance is mediated by an alternative PBP with reduced affinity for almost all β-lactam antimicrobials.^{5,6} Since its first discovery in bulk tank milk on an English dairy farm,¹ *mecC* has been found in *S. aureus* isolates from a wide range of host species, including human carriage and infection and various wildlife, companion and livestock species^{7,8} with genomic analysis indicating zoonotic transmission from livestock to humans.^{9,10} *mecC*-MRSA have been reported from a range of countries and while this geographical distribution has centred on Europe,⁸ *mecC*-MRSA have also been reported in Australia.¹¹ In addition to *S. aureus, mecC* has been described in other species of staphylococci, albeit in only a limited number of species and a small number of isolates, which have all come from animals: *Staphylococcus* *xylosus*,¹² Staphylococcus sciuri,¹³ Staphylococcus stepanovicci¹⁴ and Staphylococcus saprophyticus^{15,16} or in the case of Staphylococcus edaphicus from environmental sampling.¹⁷ mecC in MRSA is found within a staphylococcal cassette chromosome mec (SCCmec) type XI while in other staphylococci it is found in a range of genomic contexts, although always within the orfX region and with features common to SCCmec elements. Divergent mecC allotypes mecC1 and mecC2 have been described in *S. xylosus*¹² and *S. saprophyticus*,¹⁵ respectively, but have not been reported in *S. aureus*, suggesting a greater diversity and ancestral association of mecC with non-aureus staphylococci.

Here we describe a novel *mecC* allotype, *mecC3*, encoded within a distinct and novel SCC*mec* element in a newly described species, *Staphylococcus caeli*,¹⁸ which furthers our understanding of the diversity of *mecC* genes and the diverse species and genetic elements that carry it.

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Materials and methods

Isolate collection and WGS

The *S. caeli* isolate 82B^T (=NCTC 14063^T=CCUG 71912^T) was collected from air sampling in a commercial rabbit holding in Italy in 2013 as part of a previous study¹⁹ and has been described elsewhere, including its genome sequencing.¹⁸ MRSA was also present on the sampled farm.¹⁹ Six months prior to the isolation of 82B^T, ST398 MRSA belonging to t034 and t5210 were isolated from farm workers, their relatives and rabbits on the farms.¹⁹ At the time of 82^T isolation MRSA belonging to t034, t5210, t1190 and t2970 were isolated from rabbits and humans with non-typed MRSA isolates found in air samples and surface swabs.¹⁹

Sequence assembly and identification of SCCmec

Sequencing reads were *de novo* assembled using Velvet²⁰ and annotated using the NCBI Prokaryotic Genome Annotation Pipeline.²¹ Contiguous sequences (contigs) containing the *orfX* and *mecC* genes were identified using BLAST, using the respective genes from *S. aureus* LGA251 (accession number FR821779) as query sequences. Contigs NZ_FMPG01000005.1 and NZ_FMPG01000008.1 were identified as containing the *orfX* and *mecC* genes, respectively. Primers were designed for the end of contig NZ_FMPG01000005.1 and start of contig NZ_FMPG01000008.1, followed by PCR, with the resulting amplicon being ABI sequenced (Source BioScience, Cambridge, UK), to close the gap between the contigs. Primers were designed using Primer3 (http://primer3.ut.ee/).

Phylogenetic analysis of mecA homologues

Phylogenetic analyses were carried out in MEGA7.²² All nucleotide sequences were obtained from NCBI databases, using the following accession numbers: *S. aureus* LGA251, FR821779, *mecC; S. xylosus* S04009, HE993884, *mecC1; S. saprophyticus* 210, KF955540, *mecC2; S. caeli* 82B, *mecC3*, FMPG01000008; *S. sciuri* K11, Y13094, *mecA1; S. aureus* N315, NC_002745, *mecA; Staphylococcus vitulinus* CSB08, AM048810, *mecA2; Macrococcus caseolyticus* IMD0819, KY013611, *mecD;* and *M. caseolyticus* JCSC5402, NC_011996, *mecB.* The sequences were aligned using MUSCLE and a maximum likelihood tree was generated, using a general time reversible model. Site substitution rates were calculated using a discrete gamma distribution model.

Nucleotide accession numbers

The whole genome nucleotide sequences for *S. caeli* 82B^T have been deposited previously¹⁸ in the NCBI database under accession numbers NZ_FMPG00000000 (assembly) and ERR473447 (sequence reads). The assembled SCC*mec* region of 82B^T generated in this work has been deposited under accession number MH155596.

Antimicrobial susceptibility testing

Antimicrobial susceptibility testing was performed by Vitek2 using AST P260 cards following the manufacturer's instructions. Disc diffusion was performed following the EUCAST disc diffusion method Version 6.0 January 2017. Growth on MRSA Brilliance (Oxoid, Basingstoke, UK) was tested by spreading a 1 μ L loop taken from a 0.5 McFarland suspension ($\sim 1.5 \times 10^5$ cfu) on to the Brilliance plate and incubating at 30°C, 35°C and 37°C for 24 h. Interpretation was done based on the criteria for CoNS with *S. aureus* NCTC 12497 and NCTC 12493 used as control strains for susceptibility testing. PBP2a detection was performed using the PBP2' Latex Agglutination Test Kit (Oxoid) following the manufacturer's instructions.

Results and discussion

S. caeli 82B^T contains a novel mecC allotype, mecC3

Isolate 82B^T has been described as the type strain of a novel staphylococcal species, S. caeli, most closely related to S. xylosus, S. saprophyticus and S. edaphicus.¹⁸ As part of the characterization of 82B^T it underwent WGS. Further investigation of the 82B^T genome by BLAST analysis using the mecC gene from S. aureus LGA251 as the guery sequence revealed that 82B^T encodes a mecC homologue on contig NZ FMPG0100008.1. Alianments of the 82B^T mecC gene with known mecC allotypes showed that it shares 92% nucleotide identity with mecC from LGA251, 93% with mecC1 from S. xylosus S04009 and 94% with mecC2 from S. saprophyticus 210. Based on these results and in line with auidelines for the classification for mecA homologues,²³ the mecC gene for *S. caeli* 82B^T is a new allotype of *mecC* and therefore designated mecC3. The phylogenetic relationship of mecC3 to the other mecA homologues is illustrated in Figure 1; this suggests that the more common *mecC* has evolved from the more ancestral forms (mecC1, mecC2, mecC3) and that, of the three, mecC3 is closest to the ancestral form. The *mecC* complex of *S*. *caeli* 82B^T shares 93% and 92% nucleotide identity with the complete mecC complex of LGA251 and S04009, respectively, with the partial mecC complex of *S. saprophyticus* 210 sharing 94%. 82B^T gave a negative reaction using the commercial PBP2a latex agglutination test kit from Oxoid. This reaction was not altered after cefoxitin induction. mecC3 in 82B^T was, however, detected by the published mecC primer pairs 1A and 1B²⁴ and mecC-Uni-F and mecC-Uni-R.¹³

S. caeli 82B^T mecC3 is encoded within a distinct SCCmec-like element

mecC genes are typically found within SCC*mec* elements, ^{13,14} which insert into the genome at the 3'-end of a 23S rRNA methyltransferase gene, often referred to as *orfX*.^{25,26} BLASTn analysis using *orfX* from LGA251 as the query sequence identified the *orfX* region of 82B^T within contig NZ_FMPG01000005.1. PCR and amplicon sequencing confirmed that the two contigs (NZ_FMPG01000005.1 containing *orfX* and NZ_FMPG01000008.1 containing *mecC3*) are contiguous in the 82B^T genome with a gap of 140 bp between the ends of the two contigs. The assembled sequence generated in this study from these two contigs, encoding the *orfX* region of 82B^T and *mecC3* including a SCC*mec*-like element, has been deposited in NCBI with accession number MH155596.

The integration site on the genome for SCC*mec* elements within *orfX*, known as the attachment (*attB*) site, is identified by the recombinase CcrAB/CcrC via a 14 bp sequence.²⁷ Insertion into the *attB* generally results in the SCC*mec* being flanked by direct repeats, *attR* and *attL*.^{13,27,28} We therefore searched for these repeat sequences within the *orfX* region of 82B^T to define the limits of its putative SCC*mec*-like element. The search sequences; gc[ag]-tatca[tc]aaatgatgcggttt and aacc[tg]catca[tc][tc][at][ac]c[tc]gataag[ct], produced from previously identified *attR* and *attL* sequences, respectively, identified an *attR* site 51 kbp downstream of *orfX* and an *attL* site 3.8 kbp downstream of *mecC3* (Figure 2a). This indicates an SCC*mec* in 82B^T, which is 127 kbp in size, with the *orfX* and *mecC3* regions at opposite ends of the element (Figure 2a). However, it is not clear if this represents a single entity or a composite element generated by consecutive insertions.





Figure 1. Phylogenetic relationship of representative *mec* allotypes. Nucleotide sequences were aligned using MUSCLE, with the maximum likelihood method, based on the general time reversible model, being used to build the tree. The highest log likelihood (-9785.4010) tree is shown. The percentage of trees in which the associated genes clustered together, based on bootstrapping with 500 replicates, is shown at the branches. A discrete gamma distribution was used to model site substitution rates, with branch lengths measured in the number of substitutions per site.

Within this SCCmec region, 82B^T contains two ccr complexes and three joining (J) regions (Figure 2a). The ccrA and ccrB genes of 82B^T share 90% and 91% nucleotide identity with ccrA5 and ccrB3, respectively, from Staphylococcus pseudintermedius KM241 (accession number AM904731). This represents a type 6 ccr allotype,^{25,28} which is novel for an SCCmec encoding a mecC. Indeed this pairing is rare among SCCmec in staphylococci, with a search of NCBI finding only two other staphylococcal isolates that contained this ccrAB pairing [Staphylococcus cohnii WC28 (accession number GU370073)²⁹ and S. cohnii SNUDS-2 (CP019597; locus tags BZ166_04625 and BZ166_04620)]. In addition to the ccrAB genes described above, the SCC*mec* of 82B^T also contains a *ccrC* gene, sharing 90% nucleotide identity with S. aureus PM1 (SCCmec VII, accession number AB462393, gene ccrC8). The attR site was identified 51 kbp away from the orfX gene (Figure 2a), which deviates from typical SCCmec elements and therefore the orfX region was examined to find any additional attR-like site.

Within the 3' region of *orfX* an *att* site with sequence homology to the *attR SCCmec* type VII of PM1 was identified. As there are two *ccr* regions within the 82B^T SCC*mec*, it was proposed that the *attR* within the 3'-end of *orfX* (Figure 2, *attR2*) might be linked to the CcrC recombinase. Therefore, a search sequence: [at][at] [at][ct][ct][ga][cg][atc][ta][ca][at][ta][ct][ac]act[ga][ga][tc]a, based on the *attL* sequence of SCC*mec* elements containing only the *ccrC* gene, was used to identify the corresponding *attL* site linked to *attR2*, *attL2*. This revealed only one potential *attL2* site, located 16.8 kbp upstream of the *mec* region (Figure 2, *attL2*). Sequence alignment of the *attR/L* sites of 82B^T with *attR/L* sites of *ccrAB*-containing SCC*mec*, compared with the alignment of the *attR/L2* sites of 82B^T versus those *attR/L* sites of *ccrC*-containing SCC*mec*, suggest a varied *att* site for CcrAB/CcrC (Figure 3). Though it appears that the

highly conserved central 8 bp sequence is consistent, there is notable variation between the ccrAB- and ccrC-associated att sites within those bases that flank the 8 bp region (Figure 3). Although the location of *attL2* was identified with the predicted region, the only other potential *attL* site identified was *attL'*, recently described for a highly conserved type XI SCCmec found in S. xylosus 47-83.³⁰ As with the *attL'* site identified for *S. xylosus* 47–83, the *attL'* of $82B^{T}$ lies downstream of the lip gene; however, when compared with the attL of the ccrC-associated SCCmec attL sequences, it shares little similarity and lacks the central cysteine required for *attB/attSCC* recombination (Figure 3b).²⁷ Indeed, the *attL* of 82B^T shows more sequence similarity to that of attL2 from S. scuiri GVGS2 and the other *attL* sites related to CcrAB (Figure 3a). This suggests that *attL2* is most likely linked with *attR2*, though its unusual genomic location within the SCCmec suggests various DNA recombination events may have occurred.

SCCmec include J regions, defined by the areas between the orfX, ccr and mec genes, with the SCCmec of 82B^T having three such J regions. The J1 region contains primarily hypothetical genes and the ccrC gene and shares 86% identity with the J1 region of *S. aureus* PM1 SCCmec. The J2 region upstream of *attL2*, contains genes similar to those of SCCmec type V, with the presence of type 1 restriction modification genes, although there is little similarity between the restriction modification genes of 82B^T and those present on SCCmec type V.

The J2 region downstream of *attL'* shows the greatest similarity to the SCCmec XI of LGA251, sharing 89% identity, with ABC transporters, genes associated with arsenic resistance and a lipase gene. Unlike the lipase gene in LGA251 the version in 82B^T appears to be intact. The J3 region is divergent from the rest of the SCCmec of LGA251, with the exception of a putative membrane protein and a putative DNA helicase protein. A notable feature of the J3 region is the presence of a lantibiotic biosynthetic cluster that shares 90% nucleotide identity with one present on the plasmid pETB797 of S. aureus NRL 08/797 (accession number KY436025). The cluster encodes two peptide homologues of Lacticin 3147, produced by Lactococcus lactis, which has been shown to be active against Gram-positive bacteria.³¹ Within the cluster is also a gene encoding a homologue of LtnT, which is required for the transport of the Lacticin 3147 peptides as well as unrelated peptides.³² The J3 region also contains two ISL3-like transposases, with the majority of the other genes found within SCCmec being from different staphylococci.

Antimicrobial susceptibility of S. caeli 82B^T

Using Vitek 2, isolate $82B^T$ was found to be susceptible to ciprofloxacin, daptomycin, gentamicin, linezolid, mupirocin, rifampicin, teicoplanin, tigecycline and vancomycin. The isolate was resistant to clindamycin, erythromycin, fusidic acid, tetracycline and trimethoprim. Genomic analysis revealed that $82B^T$ has the resistance genes erm(B), erm(C), fusD, tet(L) and dfrK, which is consistent with the resistance profile of the isolate. With regards to β -lactam antibiotics, it was resistant to benzylpenicillin (MIC $\geq 0.5 \text{ mg/L}$) and oxacillin (MIC 1-2 mg/L) but negative in the cefoxitin resistance screen. However, by disc diffusion it was resistant to cefoxitin and displayed an MIC of 3 mg/L when assessed by Etest. The isolate failed to grow on the MRSA screening agar MRSA Brilliance despite a high inoculum and the use of different incubation temperatures.



Figure 2. Overview of *S. caeli* 82B^T SCC*mec.* SCC*mec* Type XI–LGA251 corresponds to *S. aureus* LGA251, accession number FR821779, with SCC*mec*-82B corresponding to *S. caeli* 82B^T. Regions of homology are represented by bands connecting the two sequences, with the percentage identity key shown on the right. Blue denotes normal sequence alignment (N); red denotes inverted sequence alignment (I). Key features associated with SCC*mec* elements are labelled. *att* sites are highlighted by filled circles and labelled above. This figure appears in colour in the online version of *JAC* and in black and white in the print version of *JAC*.

(a)	
attR 82B	***** ** TGTGAGAATTGCTGTTATGTTTTGTGAAGCG TATCATAA ATGATGCGGTTTTTATAAGGTTCTTCTTATT
attR GVGS2	ATCGCGCATTTAAGATCATGCGTGGGGAAGCA TATCATAA ATGATGCGGTTTTTTCAGCCGCTTCATAAAG
attR LGA251	ATAGAGCGTTTAAGATTATGCGTGGAGAA <u>GCGTATCACAAATGATGCGGTTT</u> TTTTAACCTCTTTACGTAT
attR N315	ATAGAGCATTTAAGATTATGCGTGGAGAA <u>GCATATCATAAATGATGCGGTTT</u> TTTCAGCCGCTTCATAAAG
attR_\$04009	AAAACCGCATCATTAACCGATACGCAGAAGCT TATCATAA GTGATGCGTTTT
attR_ZTA09	ATAGAGCGTTTAAGATTATGCGTGGAGAA <u>GCGTATCACAAATGATGCGGTTT</u> TTTTAACCTCTTTACGTAT
attl 82B	*** TAAACCTCATCATCAACTGATAAGCAGAAGCC TAATCCCAA GTGAAACGCTTCTGCCTATTAGAT
attl_GVGS2	TAAACCGCATCATTAACTGATAAGCATAGAGT TATCAATC TTTTTGATAATAAGAAAGTACAGAGCAACT
attl IGA251	AAAACCGCATCATCTACCGATAAGCAGAAGCA TATCATAA GTAGAAGGGGTATTAGCCAATTTAATAAAT
attL N315	AAAACCTCATCATTAACTGATACGCAGAGGCG TATCATAA GTAAAACTAAAAAATTCTGTATGAGGAGAT
attL ZTA09	A <u>AAACCGCATCATCTACCGATAAGC</u> AGAAGCA TATCATAA GTAGAAGGGGTATTAGCCAATTTAATAAAT
attL2 GVGS2	AAAACCGCATCACTACCTGATAAGTAGAGCCA TATAATAA ATAATCCTGAAAATCCACCCAATTTAAAGT
attL'_82B	AAAACCACATCACTACCTGATACGCAGAGCCA TATAATAA ATAAACGAAAAAGTCGTCCAATTTAAATGT
(b)	
(-)	****** **
attR_85_2082	AGAGGATTTAAGATTATGGGTGGAGAAGGG TATGATAA AAGAGTAAAAATTAGGTTGTGTATAATTTA
attp DN41	
attR TSGH17	AGAGCGTTTAAGATTATGCGTGGAGAGGGGT TATCATAA ATAAAAGT AAAA ATTAGATTGGGTGTAAAAATTA
attR_ICSC6082	AGAGCATTTAAGATTATGCGTGGAGAAGCG TATCATAA ATAAAACTAAAAATTAGGTTGTGTATAAATTTA
attR2_82B	CGTGCATTTAAGATTATGCGAGGGGAA <u>GCGTATCACAAATAAAACTAAAA</u> AATAGATTGTGAAAAAATATA
attL 85 2082	*** * ** A <u>AACCGCATCATCAACTGATAAGC</u> AGAAGCG TATCATAA GTAACGGAGGAGTTTTTTACCTTGTGACTT
attL_WIS	T <u>TTTTAGTAAAATCACTGGTAGGG</u> AGAGGCG TATCATAA GTGATGCTTGTTAGAATGATTTTTAACAAT
attL_PM1	T <u>TTTTAGCAAAATCACTGATAGGG</u> AGAAGCG TATCATAA ATGATGGGGTTTTAAGTACGATTTAATAAA
attL_TSGH17	T <u>TTTTAGCAAAATCACTGATAGGG</u> AGAAGCG TATCATAA ATGATGGGGTTTTAAGTACGATTTAATAAA
attL_JCSC6082	T <u>TTTTAGCAATATCACTAACAAGG</u> AGAGGCG TATCATAA GTAAAACTAAAAAATTCTGTATGAGGAGAT
attL2_82B	T <u>TTTTAGTAAAATAACTAATAAGG</u> GGAAGCG TATCACAA GTGATGCGGCTTTTTATCAGTTTTGTAAAG
attL'_82B	A <u>AACCACATCACTACCTGATACGC</u> AGAGCCA TATAATAA ATAAACGAAAAAGTCGTCCAATTTAAATGT

Figure 3. Comparison of *attR/L* sites from *ccrAB*- or *ccrC*-containing SCC*mec*. Conserved nucleotide bases within the core 8 bp region, represented in black, bold font, are indicated by an asterisk. The black triangle indicates the position of the central cytosine, thought to be essential for recombination between *attB* and *attSCC.*²⁷ Inverted repeats are marked by the underlined bases. (a) Sequences of known *attR* and *attL* sites associated with *ccrAB* [from: *S. aureus* N315 (N315), NC_002745; *S. aureus* LGA251 (LGA251), FR821779; *S. xylosus* S04009 (S04009), HE993884; *S. sciuri* GVGS2 (GVGS2), HG515014; and *S. aureus* ZTA09 (ZTA09), LK024544] were aligned and compared with those identified in *S. caeli* 82B^T. (b) Sequences of known *attR* and *attL* sites associated with *ccrC* [from: *S. aureus* 85/2082 (85_2082), AB037671; *S. aureus* WIS (WIS), AB121219; *S. aureus* PM1 (PM1), AB462393; *S. aureus* TSGH17 (TSGH17), AB512767; and *S. aureus* JCSC6082 (JCSC6082), AB373032] were aligned and compared with *attR2* and *attL2* from *S. caeli* 82B^T.

This study extends the known distribution and diversity of *mecC* genes in terms of country of isolation, sample type, staphylococcal host species, genomic context and allotypes, all of which are novel to the best of our knowledge. These highlight the complex epidemiology of this resistance determinant, particularly among CoNS. It is interesting to note that the species most closely related to *S. caeli* (*S. xylosus, S. saprophyticus* and *S. edaphicus*) have all been reported to carry *mecC* or *mecC* allotypes, which may indicate a prominent role for this group in the origins, evolution and epidemiology of *mecC* and SCC*mecC*.

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Transparency declarations

None to declare.

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