ORIGINAL RESEARCH

Staphylococcal Cassette Chromosome *mec* Typing and Multilocus Variable Number Tandem Repeat Analysis of Methicillin Resistant *Staphylococcus aureus* Clinical Isolates with Vancomycin Creep Phenomenon

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Background: The association of treatment failure and mortality with vancomycin minimum inhibitory concentration creep (MIC) is a matter of serious concern in patients with severe methicillin resistant *Staphylococcus aureus* (MRSA) infections. The purpose of the study was to identify and characterize staphylococcal cassette chromosome *mec* (SCC*mec*) and clonal types of MRSA strains, exhibiting the vancomycin MIC creep phenomenon.

Methods: A total of 3305 *S. aureus* strains were isolated from various clinical samples of Lahore General Hospital, Lahore, Pakistan. MRSA strains were identified by cefoxitin resistant (≤ 21 mm) followed by *mecA* and *mecC* gene genotyping. Vancomycin MIC creep was determined by E-test. Isolates having MIC values >1.5 µg/mL were further subjected for SCC*mec* typing (I–V and XI) and multiple-locus variable number tandem repeat analysis (MLVA) by amplification of *spa, sspA, clfA, clfB*, and *sdrCDE* genes. A dendrogram was created based on the similarity index using bioneumerics software.

Results: About 13.3% (440/3305) isolates were MRSA with 99.3% (437/440) and 0.7% (3/440) carried *mecA* and *mecC* genes, respectively. In 120 MRSA isolates, the MIC of vancomycin was >1.5µg/mL. In MRSA isolates with high vancomycin MIC (>1.5µg/mL), the most common SCC*mec* type was SCC*mec* III (38.3%), followed by SCC*mec* IVa (15.8%), SCC*mec* IIIa (13.3%,), SCC*mec* IVc (7.5%), SCC*mec* IVe (5.8%), SCC*mec* IVd (5.8%), SCC*mec* IVb (4.2%), SCC*mec* II (2.5%), SCC*mec* V (1.7%), SCC*mec* I (1.7%) and SCC*mec* XI (1.7%). MLVA revealed 60 genotypic groups of MRSA isolates having a 92% similarity index.

Conclusion: SCCmec III was the most common type in genetically related MRSA isolates showing vancomycin MIC creep. The presence of SCCmec XI may further add burden to infection control measures.

Keywords: methicillin resistant *Staphylococcus aureus*, staphylococcal chromosomal cassette, multilocus variable number tandem repeat analysis

Introduction

MRSA is the leading cause of morbidity and mortality in both healthcare and community-acquired settings.¹ The emergence and spread of MRSA strains with multidrug resistance genes have severely limited the treatment options for staphylococcal infections around the world.²

For many years, vancomycin has been a staple of treatment for serious MRSA infections.³ An increase in vancomycin minimum inhibitory concentration (MIC) values in MRSA strains even within the susceptible range (MIC creep) is the

cause of serious concern.⁴ This phenomenon of vancomycin MIC creep may be a significant risk factor for public health, posing additional challenges in the management of MRSA infections.⁵ Microbiological and clinical data indicate treatment failure in infections with MRSA strains exhibiting vancomycin MIC creep. Several studies found poor clinical outcomes despite a vancomycin MIC in the susceptible range of $\leq 2\mu g/mL$ especially when ranging MIC 1.5–2 $\mu g/mL$.⁶ Rapid and reliable typing methods are thus required for the identification of such precarious MRSA clones, particularly those with high vancomycin MIC values.⁷ SCC*mec* typing is one of the most important methods for studying not just MRSA epidemiology but also the evolution and global spread of MRSA clones.⁸ SCC*mec* acquisition by methicillinsensitive *S. aureus* strains has the potential to transform them into hazardous multidrug resistant pathogens with the ability to adapt and thrive in a hospital setting with poor infection control.⁹ Several different SCC*mec* types have been recognized worldwide.¹⁰ Thirteen allotypes of SCC*mec* have been identified, ranging from I to XIII, based on the combination of complexes of the *mec* and *ccr* gene.¹¹ Human infections have been linked to SCC*mec* types from I–VIII.¹² However, SCC*mec* types IX and X, are linked to livestock infection.¹³ All these SCC*mec* types have now been disseminated worldwide and new variants are constantly emerging.¹⁴

Analysis of variable number tandem repeats (VNTR) loci identified in the genomes of several bacterial species has resulted in advances in molecular typing. Multiple-locus VNTR analysis (MLVA) refers to a molecular typing system established by analysing the number of repeats on different VNTR loci. The MLVA system is used to determine the relationship of isolates and can also provide insights into population structure.¹⁵ MLVA can well differentiate between phylogenetically unrelated strains.¹⁶ For the major clonal complexes of *S. aureus*, the MLVA scheme has high-quality type ability and excellent discriminatory power.¹⁷ MLVA has also been reported as a reliable tool for understanding the evolution of MRSA.¹⁸

The current study aimed to identify and characterize SCCmec and clonal types of MRSA strains that exhibited vancomycin MIC creep in clinical isolates.

Materials and Methods

Sample Collection

The study was approved by the ethical review board of the University Health Sciences and Lahore General Hospital (LGH), Lahore, Pakistan (UHS/REG-18/ERC/3590). A total number of 3305 *S. aureus* isolates were collected from the laboratory of LGH during five years (2016–2020). Isolates were recovered from different clinical samples such as pus, wound swabs, blood, sputum, urine, and aspirates from patients admitted to the wards such as medicine, surgery, gynecology, eye, ear, nose and throat, pediatric, and intensive care units.

Culture and Identification of S. aureus Isolates

S. aureus isolates were identified after observing colony morphology on blood agar plates and biochemical testing (catalase, coagulase, and DNase positive). ATCC 12600 *S. aureus* was used as control strains.

Identification of MRSA by Kirby-Bauer Disc Diffusion Method

Antimicrobial susceptibility was accomplished according to clinical laboratory and standard institute (CLSI) guidelines (2019) using cefoxitin disc ($30\mu g$). All the strains with cefoxitin zone diameter of ≤ 21 mm were considered methicillin resistant. MRSA ATCC 33591 and MSSA ATCC 25923 were used as control strains.

Extraction of DNA from Isolates

DNA was extracted by the boiling method as described previously.¹⁹ The quality and quantity of extracted DNA were determined by nanodrop (spectrophotometer, Bio Rad, USA). The quality of the extracted DNA was predicted from the ratio of absorbance at 260 and 280 nm.

Amplification of mecA and mecC Gene

mecA and mecC genes were amplified using 50ng extracted genomic DNA.^{20,21} MRSA ATCC 33591 and BAA 2312 were used as a control strain for the mecA and mecC genes, respectively.

Determination of Vancomycin MIC for MRSA Isolates

Vancomycin MICs for MRSA isolates were determined using the E-test method. Based on MIC results, isolates were classified as susceptible or resistant according to criteria set by CLSI. Isolates with vancomycin MICs $\leq 2\mu g/mL$ were considered as sensitive, $4-8\mu g/mL$ were regarded as vancomycin intermediate *S. aureus* (VISA), and $\geq 16\mu g/mL$ were considered as vancomycin resistant *S. aureus* (VRSA). Isolates with vancomycin MIC >1.5 µg/mL were subjected for molecular characterization by SCC*mec* typing and MLVA.

Determination of SCCmec Types I-V and XI

SCC*mec* gene identification I–V and SCC*mec* XI were carried out as described previously by Zhang et al and Stegger et al, respectively.^{22,23} The primers for SCC *mec* I–V and SCC*mec* XI are given in the <u>Supplementary Table</u>. BAA 2312 was used as a control strain for SCC*mec* XI.

MLVA

MLVA was performed by multiplex PCR to simultaneously amplify the hypervariable VNTR regions of the *spa, sspA, clfA, clfB*, and *sdrCDE* genes.¹⁶ The primers for MLVA are given in the <u>Supplementary Table</u>.

Data Analysis

SPSS version 27.0 was used to analyze the data. The MIC population distribution for each year was graphically evaluated to assess vancomycin MIC creep. The SCC*mec* typing findings for each year were given in the form of percentages for each SCC*mec* category including SCC*mec* type I–V and SCC*mec* XI. A dendrogram was built based on variation in the base pairs (bp) of repeats for MLVA using the bionumerics 8.0 program to show the relatedness and genetic diversity among MRSA. Number of repeats that were obtained at specific loci for isolated strains were fed into Bionumeric software version 8.0 to draw the minimum spanning tree (MST), also known as a phylogenetic tree. In this program, the order of each allelic repeat that fed into software was (*clfA, clfB, sdrD, sdrC, sdrE, spa,* and *sspa*).

Results

Genotypic evaluation of cefoxitin resistant *S. aureus* isolates confirmed 440 MRSA harboring the *mecA* and *mecC* genes. About 13.3% (440/3305) isolates were MRSA with 99.3% (437/440) and 0.7% (3/440) carried *mecA* and *mecC* genes, respectively. Percent of MRSA isolates with vancomycin MIC 0.5 μ g/mL, 0.75 μ g/mL, 1 μ g/mL, 1.5 μ g/mL and 2 μ g/mL is depicted in Figure 1. The vancomycin MIC "1.0" μ g/mL had declined from 28.6% of isolates in 2016 to 14.8% in 2020. The percentage of isolates with MIC "2.0" μ g/mL increased from 19.0% to 39.8% during 5 years. A total of 120 MRSA isolates were found to have MIC values of >1.5 μ g/mL during 5 years. However, none of the isolates was found to be VISA and VRSA.

When isolates with vancomycin MIC >1.5 μ g/mL were tested for SCC*mec* typing, it was observed that SCC*mec* III was found to be the most common type in 38.3% of these isolates, followed by SCC*mec* IVa (15.8%), SCC*mec* IIIa (13.3%), SCC*mec* IVc (7.5%), SCC*mec* IVe (5.8%), SCC*mec* IVd (5.8%), SCC*mec* IVb (4.2%), SCC*mec* II (2.5%), SCC*mec* V (1.7%), SCC*mec* I (1.7%) and SCC*mec* type XI (1.7%). In addition, 1.7% of MRSA isolates were untypeable [Table 1]. Amplification of SCC*mec* types I–V and SCC*mec* XI elements are depicted in Figures 2 and 3, respectively.

By MLVA analysis using Bionumeric 8.0 software, all the tandem repeats of 120 isolated strains get shuffled and presented the relatedness by amplified bands at their specific loci (*clfA, clfB, sdrD, sdrC, sdrE, spa, and sspa*) in the dendrogram [Figure 4]. Sixty MLVA haplotypes were reported out of 120 MRSA stains with vancomycin MIC value >1.5µg/mL [Figure 4]. Concerning to 100% similarity between the strains, the stains having the same genetic clusters were placed under one genotype. The strains with the same genotype were exactly like each other for their band position at specific loci although they have different strain sources. Therefore, using MLVA analysis even a slight difference of base pairs and their position at specific loci was detected and placed in the separate genotypic category. MLVA shuffled all the tandem repeats of strains and then arranged them according to the similarity pattern. This makes closely related or similar genotypic strains lie close together while unidentical repeats at their loci go apart.

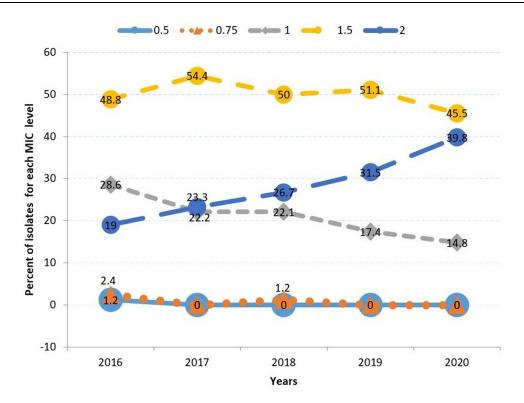


Figure 1 Distribution trend of vancomycin MIC values (2016–2020) (n = 440). MRSA isolates with vancomycin MIC values of 0.5µg/mL and 0.75µg/mL remained at very low levels throughout the years (lower two lines). Isolates with MIC 1µg/mL show an evident decline as indicated with the grey line while isolates with MIC value of 2µg/mL show a clear rising trend (dark blue line). At the top of the figure, the yellow line depicts the highest percentage of MRSA isolates with vancomycin MICs values of 1.5 µg/mL.

In the analysis of MST, order of each allelic repeats that fed into software was (*clfA*, *clfB*, *sdrC*, *sdrE*, *spa*, *and sspa*). Out of 120 isolated strains, a total of 60 clusters were connected following the distribution of VNTRs. The distribution of strains occurred based on the variation in the base pair size of each strain at single or multiple loci and lead to the formation of a new MLVA strain [Figure 5]. In this MST it was seen that circles represent the number of isolates; circles of similar colors represent that they belong to one or the same genotype. Furthermore, more closely related genotypic clusters of isolated MLVA strain were connected with a solid bold line. On the other hand, distant or less closely related genotypes were joined with thin or light lines. In this phylogenic tree, closely related genotypes were encoded with same color such as strain ID 416, 418, 422 and 420 clustered and connected with solid lines and represented with orange color and it belong to genotypic group 44.

Discussion

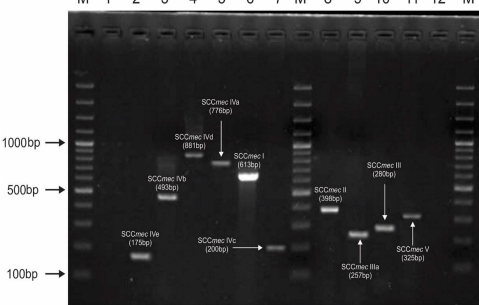
A significant shift toward higher vancomycin MIC values (MIC creep) has been observed, which raises serious concerns.²⁴ MIC creep has been associated with therapeutic failure in patients with serious MRSA infections. High mortality was seen in patients of MRSA bacteremia having vancomycin MIC $\geq 1.5 \,\mu$ g/mL. The physician must be warned of the potential risk of treatment failure at high MIC value and alternative treatments should be opted if necessary.^{25,26} It is critical to assess the occurrence and trend of vancomycin MIC creep phenomenon in settings to generate evidence-based knowledge for clinicians and better management of patients. Even though new drugs such as dalbavancin and ceftaroline have been developed to be an effective alternative therapy for multi-drug resistance MRSA strains, vancomycin remains the first choice in the majority of countries around the world.²⁷ Moreover, by determining the molecular characteristics of such infectious and treatment challenging strains exhibiting MIC creep with high MIC values, we will be able to focus on our infection control practices to avoid their dissemination in the hospital and community.

SCCmec characterization of these isolates revealed that SCCmec III and IIIa were the most prevalent mec types among MRSA isolates as indicated in studies of Iran and Malaysia.^{28,29} A high prevalence of SCCmec type III/IIIa has

SCCmec Types	2016	2017	2018	2019	2020	Total
	n (%)	n (%)	n (%)	n (%)	n (%)	n (%)
1	0 (0.0)	I (4.8)	0 (0.0)	I (3.6)	0 (0.0)	2 (1.7)
П	0 (0.0)	I (4.8)	I (4.5)	0 (0.0)	I (2.9)	3 (2.5)
ш	7 (46.7)	6 (28.6)	8 (36.4)	11 (39.3)	1 (41.2)	46 (38.3)
Illa	I (6.7)	3 (14.3)	3 (13.6)	4 (14.3)	5 (14.7)	16 (13.3)
lva	2 (13.3)	5 (23.8)	3 (13.6)	4 (14.3)	5 (14.7)	19 (15.8)
IVь	I (6.7)	0 (0.0)	2 (9.1)	I (3.6)	I (2.9)	5 (4.2)
IVc	2 (13.3)	2 (9.5)	I (4.5)	2 (7.1)	2 (5.9)	9 (7.5)
IVd	0 (0.0)	2 (9.5)	I (4.5)	3 (10.7)	I (2.9)	7 (5.8)
IVe	I (6.7)	I (4.8)	I (4.5)	2 (7.1)	2 (5.9)	7 (5.8)
v	0 (0.0)	0 (0.0)	I (4.5)	0 (0.0)	I (2.9)	2 (1.7)
ХІ	0 (0.0)	0 (0.0)	l (4.5)	0 (0.0)	I (2.9)	2 (1.7)
NT	I (6.7)	0 (0.0)	0 (0.0)	0 (0.0)	I (2.9)	2 (1.7)
Total	15 (100.0)	21(100.0)	22 (100.0)	28 (100.0)	34 (100.0)	120 100.0)

Table I Distribution of SCCmec Gene in MRSA Isolates (2016-2020)

been reported in three different geographical areas of India and Pakistan.³⁰ A previous study of 11 Asian countries revealed that SCC*mec* type II was the most prevalent in Japan and Korea while SCC*mec* type III was most prevalent in Saudi Arabia, Singapore, Indonesia, Vietnam, Sri Lanka, Thailand, India, and the Philippines. A low prevalence of



M 1 2 3 4 5 6 7 M 8 9 10 11 12 M

Figure 2 Gel electrophoresis pattern for identification of SCCmec types and subtypes by multiplex PCR for clinical isolates of MRSA. Lane 1 is negative control. Lanes 2 represents SCCmec type IVe (175bp); lane 3 represents SCCmec type IVb (493bp); lane 4 represents SCCmec type IVd (881bp); lane 5 represents SCCmec type Iva (776bp); lane 6 represents SCCmec type I (613bp); lane 7 represents SCCmec type IVc (200bp); lane 8 represents SCCmec type II (398bp). SCCmec type III a (257bp); lane 10 represents SCCmec type V (325bp). Lane 12 is empty. M is the 100 bp DNA ladder (Thermo Scientific).

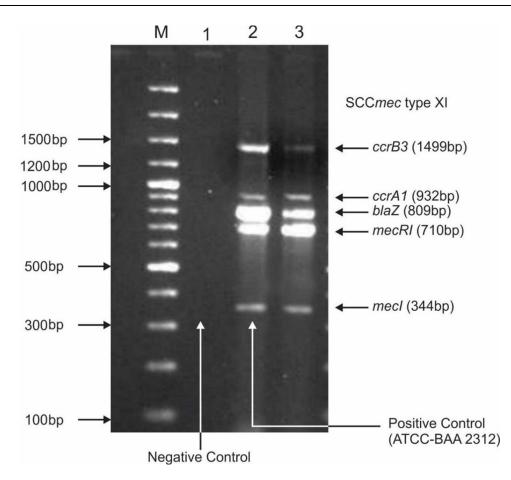


Figure 3 Gel electrophoresis pattern for identification of SCCmec type XI by multiplex PCR for clinical isolates of MRSA. M is a 100 bp DNA ladder (Thermo Scientific). Lane 1: negative control; lane 2 represents positive control of SCCmec XI ATCC- BAA 23122, Lane 3: MRSA strain represents SCC mec type XI carrying mec1 (344bp); mecR1 (710bp); blaZ (809bp); ccrA1 (932bp) and ccrB3 (1499bp).

SCC*mec* type I in these Asian countries.³¹ SCC*mec* III is commonly carried by healthcare associated MRSA (HA-MRSA). HA-MRSA is multidrug resistant and causes infections commonly in elderly patients with underlying bronchopulmonary and cardiovascular diseases.³²

The predominance of SCC*mec* III and IIIa in our setting might be because of antibiotic selective pressure, which led to clonal selection and dissemination. SCC*mec* is important not only in antimicrobial resistance but also in the molecular epidemiology and evolution of MRSA. As a result, a thorough understanding of the prevalence and characteristics of SCC*mec* types may play a potential role in the investigation, surveillance, and implementation of MRSA, ultimately assisting in the development of precautionary and therapeutic methodologies.³³ Similarly, another study in Pakistan reported a high percentage of SCC*mec* type III and its variants.³⁴ SCC*mec* type IV, sub types (Iva-IVe) were the next leading types in this study. Similarly, another study reported that SCC*mec*IV types and subtypes were most common after SCC*mec* type III.³⁵ However, a study in Pakistan revealed that the most prevalent SCC*mec* type was SCC*mec* type IV followed by SCC*mec* type II and then III.³⁶ Another study reported majority of isolates carrying SCC*mec* type IV followed by SCC*mec* type V.¹⁴ SCC*mec* IV and V are main types in community-acquired MRSA (CA-MRSA). CA-MRSA is well recognized for its ability to cause diseases, such as skin and soft tissue infections and necrotizing pneumonia.³⁷

Concerning the significance of SCC*mec* typing, a study described that SCC*mec* associated virulence factors may have an impact on disease outcome, with SCC*mec* type IVa being associated with more metastatic diseases and SCC*mec* II being associated with increased patient mortality. The study recommended that more research is needed to determine which virulence factors of SCC*mec* types influence disease outcomes.³⁸

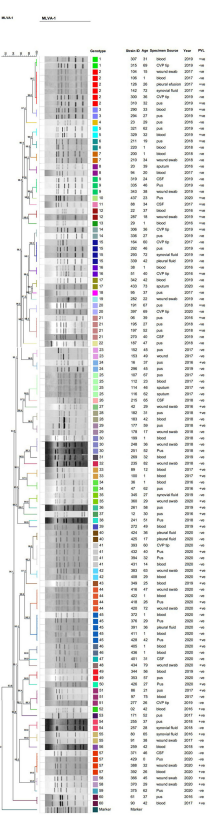


Figure 4 Dendrogram based on the clustering of strains according to similarity index (by Dice coefficient and UPGMA approach) through MLVA analysis of MRSA strains. By MLVA analysis 120 isolates were clustered into 60 haplotypes based on similarity index due to slight variations in VNTRs using Bionumerics Software version 8.0. A dendrogram was produced with the settings of position tolerance (optimization 1% and band position tolerance 0.75%). All strains were shuffled according to their similarity index and clustered into genotypes.

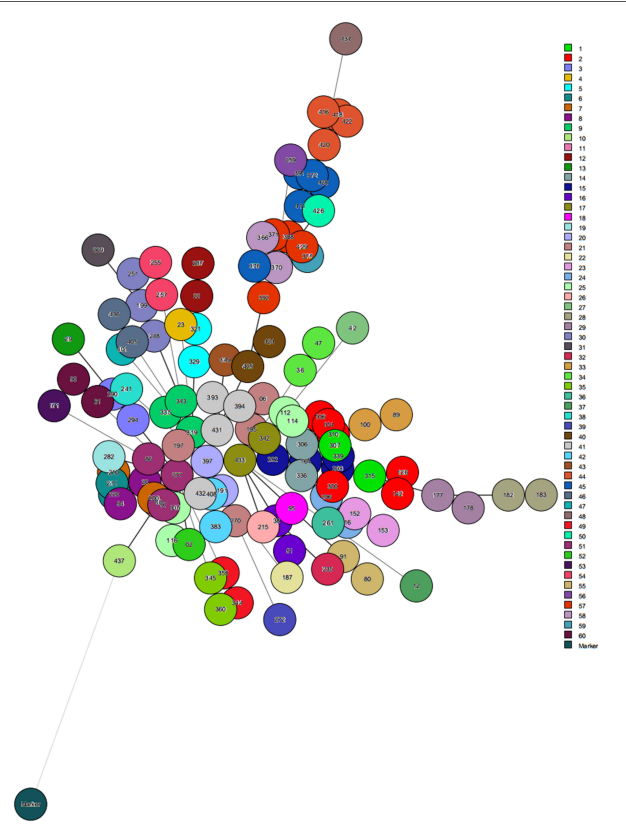


Figure 5 Minimum Spanning Tree analysis of MRSA isolates depicting their phylogenetic relationship according to VNTRs (*dfA*, *dfB*, *sdrD*, *sdrC*, *sdrE*, *spa* and *sspa*) distribution index. It is based on slight base pair variations leading to a new MLVA type strain using Bionumerics software version 8.0. In this MST analysis total 60 clusters were found. Circles are representing MRSA isolates and the same colored circles belong to one genotype. Furthermore closely related genotypes were connected with bold lines while distantly related were connected with thin lines.

SCC*mec* I and II were shown to have a very low prevalence in the current study. In Europe, however, there was a higher frequency of SCC*mec* I and II, as well as a predominance of SCC*mec* IV and V in hospital settings, but a surprising paucity of SCC*mec* III.³⁹ Another study reported a high frequency of SCC*mec* type IA and IIC among MRSA isolates for the first time in Asian countries.⁴⁰

The discrimination of HA-MRSA and CA-MRSA has become more complicated due to changing epidemiology as the community-acquired SCC*mec* types infiltrate in hospital settings and vice versa. This transmission, intermingling, and increasing diversity of SCC*mec* types carrying multiple drug-resistant genes among the community and hospital settings can worsen the clinical outcomes thus limiting the therapeutic options.⁴¹ The detection of *mecA* and *mecC* genes at the molecular level demonstrates an ongoing evolutionary process in MRSA strains. Simultaneous detection of *mecA* and *mecC* genes is critical for correct MRSA strain identification.⁴² If MRSA is not detected in time, it can lead to treatment failure and the spread of resistant strains in the community.⁴³ The situation may impose an additional medical and financial burden.⁴⁴

The situation has become more complicated due to the emergence of the SCC*mec* type XI disseminating *mecC* gene in animals and humans.⁴⁵ SCC*mec* type XI has a wide geographical distribution and has adapted to a variety of host species, including livestock and wildlife. This could be a potential source of zoonotic infections and may complicate accurate diagnosis, thus adding medical and financial burdens.^{2,46} A less frequent SCC*mec* XI was also found in 1.7% of MRSA isolates. SCC*mec* XI is characterized by different components including *mecI*, *mecR1*, *blaZ*, *ccr* A, and *ccrB* genes. SCC*mec* XI has also been reported in Ireland, Germany, and Japan.^{47–49}

Another study documented that even though SCC*mec* type XI is uncommon in human isolates, its presence can be misjudged, necessitating molecular typing to confirm its presence.⁵⁰ Another study also stated that SCC*mec* XI containing toxigenic virulence factors raises the alarm for MRSA isolates with high virulence potential in Europe, emphasizing the importance of proper molecular detection.⁴⁶ Furthermore, a study identified SCC*mec* XI by detecting its various components, but few of those isolates lacked *ccrA* and *ccrB* genes, contrary to our findings. They did, however, speculate that isolates with *ccrA* and *ccrB* genes might have minor sequence differences in their SCC*mec* type XI *ccr* specific alleles.²³

MLVA was performed on MRSA isolates in this study to obtain phylogenetic relationships in the form of a dendrogram. By using the MLVA technique with Bionumeric software 8.0 we were able to separate the PCR amplicons even with small differences in a band size that allowed us to obtain an accurate product size. MRSA isolates produce a PCR product with specific VNTRs converted into tandem repeats at their specific loci. The genotypes were formed by MLVA and shrunk down to 60 haplotypes. A minor change in the base pair of the VNTR created a new MLVA strain, which now belongs to a different genotypic group. Similarly, 63 MLVA types were identified in 123 MRSA isolates in an earlier study in Pakistan. MLVA was found an effective technique to understand the phylogenetic relationship and distribution of MRSA isolates.¹⁸ Moreover, in another study, MLVA was used to type *S. aureus* isolates. MLVA performed exceptionally well in distinguishing between unrelated isolates.⁵¹

Low genetic diversity was found among MRSA strains in this study because all strains had a similarity index above 92%. A similar observation of diversity was documented by another study but in different bacterial species in India.⁵² MLVA can be used as a standardized and portable molecular typing technique. This technique can contribute to our understanding of the genetic diversity of various pathogens, particularly in areas where new isolates are emerging.⁵³ A previous study also documented that MLVA can differentiate between strains of similar lineages in clonal complexes.¹⁵

Conclusion

In conclusion, SCCmec III was found to be the most common type in MRSA isolates exhibiting the vancomycin MIC creep phenomenon. Discovery of mecC gene and the presence of SCCmec XI indicates zoonotic origin of these MRSA isolates which highlights the complexity of the antimicrobial resistance scenario. A similarity index above 92% by MLVA analysis indicated that genetic diversity among the MRSA population was very low and strains were related to each other. Data on identification and characterization of clones of MRSA with high MIC values will provide insight to clinicians and health professionals to formulate local antibiotic policy in each hospital and

implementation of infection control measures which consequently will hamper the distribution and spread of MRSA in Pakistan.

Data Sharing Statement

The database generated and analysed during the current study is available from the corresponding author on reasonable request.

Ethics Approval

Clinical isolates of *S. aureus* were obtained after getting an ethical approval of the ethical review committee of Lahore General Hospital and University of Health Sciences, Lahore, Pakistan ref# (UHS/REG-18/ERC/3590).

Consent for Publication

Permission was taken from the hospital to publish the findings of study.

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Author Contributions

All authors contributed significantly to the work reported, whether in the conception, study design, execution, data acquisition, analysis, and interpretation, or in all of these areas; participated in the drafting, revising, or critical review of the article; gave final approval of the version to be published; agreed on the journal to which the article was submitted; and agreed to be accountable for all aspects of the work.

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Disclosure

The authors have no relevant financial or nonfinancial interest to disclose in this work.

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