Genomic diversity of *mec* regulator genes in methicillinresistant *Staphylococcus aureus* and *Staphylococcus epidermidis*

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(Accepted 2 May 1996)

SUMMARY

Low-affinity penicillin-binding protein PBP-2a encoded by mecA is closely related to methicillin resistance in staphylococci, and expression of PBP-2a is controlled by regulator elements encoded by mecR1 and mecI which are located adjacent to mecA on the chromosome. Deletion or mutation which occurred in *mec* regulator gene is considered to be associated with constitutive production of PBP-2a. The distribution of the mec regulator genes in 176 strains of Staphylococcus aureus and 33 strains of S. epidermidis isolated from a single hospital was studied by polymerase chain reaction amplification. Most clinical isolates of methicillin-resistant S. aureus (MRSA) (94.3%) and S. epidermidis (MRSE) (83.9%) possessed both mecI and mecR1 genes (type I), whereas no mec regulator genes were detected in mecA-negative isolates. In contrast, 7 MRSA and 5 MRSE isolates were found to have incomplete regulator genes, and they were classified into three groups; strains which lacked only mecI gene (type II), strains which lacked *mecI* and 3'-end of *mecR1* gene (type III), and strains which lacked both regulator genes (type IV). Analysis of mecI gene from all the strains having mecI by restriction fragment length polymorphism after Mse I digestion indicated that three MRSA strains possessed one of the known point mutations identified previously. These findings indicated the predominance of a single type of MRSA possessing both mecl and mecR1 in the study period and also suggested a high genomic diversity in *mec* regulator region of staphylococci.

INTRODUCTION

Methicillin-resistant S. aureus (MRSA) and S. epidermidis (MRSE) are recognized as pathogens that can cause serious nosocomial infection and contribute significantly to mortality and morbidity in hospitals around the world [1, 2]. Resistance to methicillin in staphylococci is primarily mediated by penicillinbinding protein (PBP-2a) which is a unique cell wallsynthesizing enzyme [3–6]. Due to the low affinity for beta-lactam antibiotics, PBP-2a can compensate for the transpeptidation function of all other PBPs in staphylococci and enable the organism to survive in the presence of lethal concentrations of beta-lactams [7]. PBP-2a is encoded by the chromosomal structural gene mecA which is highly conserved in clinical isolates of methicillin-resistant staphylococci [8–10]. However, the level of methicillin resistance is affected by other chromosomal factors (fem or aux factors) encoded by femA, femB, femC, and femD genes which are not linked to the mec locus [11–13].

Expression of PBP-2a is considered to be controlled by regulator elements which are putative transmembrane-signalling protein and repressor protein encoded by *mecR1* and *mecI* genes, respectively [14– 16]. These regulator genes, *mecR1* and *mecI* are located upstream of the mecA gene on the chromosome. The expressions of these regulators strongly repress mecA transcription in the absence of inducer [15, 17]. Hence MRSA possessing intact mecR1 and mecI as well as mecA are phenotypically methicillin susceptible because of the repression of PBP-2a production by mec regulator elements. However, deletion or mutation in the mec regulators was detected in clinical MRSA isolates and methicillin-resistant coagulase negative staphylococci (MR-CNS) highly resistant to methicillin [18]. Such genomic changes in mec regulator genes are considered to alter or remove their repressor function on mecA gene transcription, which may lead to constitutive production of PBP-2a. Therefore, it is important to analyse the genomic diversity found in mec regulator genes of staphylococci in order to understand the molecular basis for methicillin resistance. Several studies reported for the past three decades have shown genomic variation in mec regulator genes among MRSA and MR-CNS strains isolated from different countries and at different times [17-19]. However, the extent of diversity in mec regulator genes has not been studied among staphylococcal isolates from a limited area or from a single hospital.

In the present study, we investigated the distribution of *mec* regulator genes and the presence of the previously reported mutations in *mecI* among *S*. *aureus* and *S*. *epidermidis* strains derived from a single hospital, using polymerase chain reaction (PCR) and restriction fragment length polymorphism (RFLP) analysis. Results obtained revealed the present situation of *mec* regulator gene distribution in staphylococcal clinical isolates and suggested the occurrence of greater genomic variation in these genes than has been reported previously.

MATERIALS AND METHODS

Bacterial strains

A total of 176 S. aureus and 33 S. epidermis strains were employed in this study. The S. aureus strains consisted of 122 and 28 clinical isolates of MRSA and methicillin-susceptible S. aureus (MSSA), respectively, and of 26 environmental isolates of MRSA. All S. epidermidis isolates were derived from clinical specimens and 31 strains were identified as MRSE. Clinical isolates of S. aureus and S. epidermidis were isolated at Sapporo Medical University Hospital, Sapporo, Japan, in 1993 (between January and March), 1994 (between January and April), and 1995 (between February and May). The S. epidermidis isolates were derived from 33 different patients, whereas 122 MRSA were isolated from 97 patients. Multiple isolates of MRSA were obtained from different sites of 22 patients over a period of 2 weeks from a half of these patients, while almost simultaneously from the other patients. Environmental MRSA were isolated from swabs from floors and sinks of wards in three clinical departments of the hospital in January, March and April in 1994; an overall detection rate throughout the study period was 59.4%. Identification of bacterial species and antimicrobial susceptibility tests were performed with MicroScan WalkAwayTM-96 (Baxter Diagnostics, Inc., West Sacramento, USA). Differentiation of S. aureus used in this study was made by coagulase typing with coagulase type-specific antisera. Coagulase types of MRSA clinical isolates consisted of type II (91.8%), VII (5.7%), and IV (2.5%); all the environmental MRSA isolates belonged to type II. In contrast MSSA clinical isolates consisted of type VII (46·4%), II (14·3%), V (14·3%), III (7·1%), IV (7.1%), and untypable strains (10.7%).

Preparation of bacterial DNA and PCR

DNA samples for PCR were prepared as described previously [20]. Three sets of primers indicated in Table 1 were used for PCR to detect mec regulator genes. As shown in Figure 1, primers mecI1 and mecI2 amplified the whole open reading frame (ORF) of mecI gene, whereas primer pairs mecRA1 and mecRA2, and mecRB1 and mecRB2 amplified the regions located on the half portion of 5' end (mecR1A) and 3' end (mecR1B) of mecR1 gene, respectively. In addition, mecA gene fragment was also amplified by PCR as described in our previous study [20], in which detection of each mecA gene was performed with isolates obtained in 1993. Methicillin resistance of staphylococci was essentially defined by the presence of mecA gene encoding PBP-2a irrespective of the degree of phenotypic methicillin susceptibility [21, 22]. Therefore, in the present study, S. aureus and S. epidermidis possessing mecA gene were designated MRSA and MRSE, respectively.

DNA amplification was performed on a thermal cycler in 100 μ l of reaction mixture containing 200 μ M each of dATP, dCTP, dGTP, and dTTP; 100 pM each of a pair of primers; 2.5 U of AmpliTaq DNA polyermase (Perkin Elmer); 10 mM Tris-HCl (pH 8.3); 50 mM KCl; and 1.5 mM MgCl₂. DNA fragments were amplified for 25 cycles, with each cycle consisting of 1 min at 97 °C for denaturation, 1 min at 57 °C for

	Oligodeoxyı	nucleotide primer			
Amplified gene*	Pair Sequence (5' to 3') [†]		Position ⁺	Size of PCR product (bp)	
mecR1 (mecR1A)	mecRA1	+ GTCTCCACGTTAATTCCATT	133–152	310	
	mecRA2	- GTCGTTCATTAAGATATGACG	422-442		
mecRl (mecRlB)	mecRB1	+AAGCACCGTTACTATCTGCACA	986-1007	236	
	mecRB2	– GAGTAAATTTTGGTCGAATGCC	1200-1221		
mecI	mecIl	+AATGGCGAAAAAGCACAACA	1699-1718	481	
	mecI2	- GACTTGATTGTTTCCTCTGTT	2159-2179		

Table 1. Primers used for PCR

* mecR1A and mecR1B represent half portion of the 5' and 3' end of mecR1 gene, respectively. Amplified regions of mecR1 and mecI genes are shown in Fig. 1.

+ +, sense primer; -, antisense primer.

[‡] Positions are expressed as the nucleotide number from the first ORF of *mec* regulator gene (*mecR1*) based on the sequence data reported previously [14].

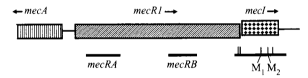


Fig. 1. Schematic representation of mecA and mec regulator genes, and amplified regions by PCR (mecR1A, mecR1B, and mecI), which are shown as horizontal bars. Vertical lines on the amplified region of mecI denote cleavage sites of restriction enzyme Mse I. Two Mse I sites indicated as M_1 and M_2 are not present in normal mecI gene but appear as a result of point mutations previously reported, a nucleotide substitution at positions 202 and 260, of mecI gene, respectively [18]. Arrows indicate directions of transcription.

annealing, and 2 min at 72 °C for primer extension. The amplified PCR product $(10 \ \mu l)$ was analysed by electrophoresis on 1% agarose gel and visualized with u.v. light after staining with ethidium bromide.

RFLP analysis of mecI gene

RFLP analysis was carried out for mecI gene in order to detect point mutations which had been described previously by Suzuki and colleagues [18]. They identified three point mutations in mecI gene of the 6 clinical MRSA strains derived from 5 different countries; nucleotide substitution (C to T) at position 202 (mutation 1), nucleotide substitution (T to A) at position 260 (mutation 2), and deletion of nucleotide (T) at position 143 (mutation 3). Mutations 1 and 2 were detected in 5 of the 6 MRSA strains examined, suggesting these to be dominant types of mutation. The mutations 1 and 2 result in a new nucleotide sequence (5'-TTAA-3') which is recognized by restriction endonuclease Mse I (5'-T \downarrow TAA-3'). Hence RFLP analysis was done in this study using the PCR product of mecI gene and Mse I. Whereas the PCR

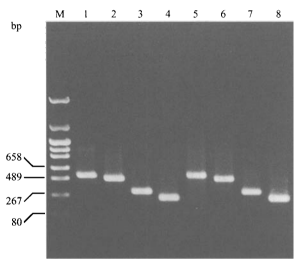


Fig. 2. PCR products obtained from clinical MRSA isolates (strain SH162 for lanes 1–4, strain SH163 for lanes 5–8). Amplified genes are as follows: lanes 1 and 5, *mecA* (533 bp); lanes 2 and 6, *mecI* (481 bp); lanes 3 and 7, *mecR1A* (310 bp); lanes 4 and 8, *mecR1B* (236 bp). M, molecular weight marker.

product for *mecI* gene originally contained 3 Mse I sites as shown in Figure 1, generation of fragmentation patterns distinct from that of normal *mecI* gene implied the emergence of novel Mse I sites due to mutation 1 or 2. After PCR product was digested with Mse I (New England Biolabs) according to the manufacturer's instructions, DNA fragments were separated by electrophoresis on a 1% agarose gel.

RESULTS

DNA fragments with the size of 310 bp, 236 bp, and 481 bp were amplified by PCR from *mecR1A*, *mecR1B*, and *mecI* gene of MRSA, respectively, as shown in Figure 2. On the basis of detection pattern

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Bacterial species*	Number of isolates					Detection patterns of PCR product		
		PCR product					Number of isolates	Number of oxacillin- susceptible
		mec A	mecR1A	mecR1B	mecI	Type	(%)	strain
MRSA (clinical isolates)	122	+	+	+	+	I	115 (94.3)	0
		+	+	+	_	II	3 (2.5)	0
		+	+	-	_	III	1 (0.8)	1
		+	_	-	_	IV	3 (2.5)	2
MRSA	26	+	+	+	+	Ι	23 (88.5)	0
(environmental strains)		+	-	-	—	IV	3 (11.5)	0
MSSA	28	_		-	-	V	28 (100)	28
MRSE	31	+	+	+	+	Ι	26 (83.9)	1
		+	+	+	_	II	1 (3.2)	0
		+	+	_	-	III	3 (9.7)	1
		+	_	-	-	IV	1 (3.2)	0
MSSE	2	_	_	-		V	2 (100)	2

Table 2. Detection of mecR1 and mecI genes in S. aureus and S. epidermidis

* MRSA and MSSA represent methicillin-resistant and methicillin-susceptible *S. aureus*, respectively. Similarly, MRSE and MSSE indicate methicillin-resistant and methicillin-susceptible *S. epidermidis*, respectively.

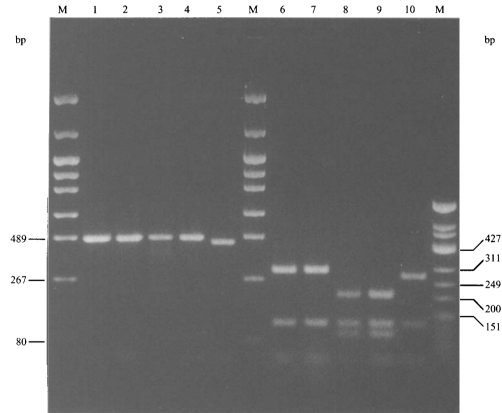


Fig. 3. RFLP analysis of *mec1* gene with *Mse* I restriction endonuclease. Lanes 6–10 represent RFLP patterns derived from *mec1*-PCR product arranged in the same order in lanes 1–5. Lanes 6 and 7, RFLP pattern 1; lanes 8 and 9, pattern 2; lane 10, pattern 3. *mec1*-PCR products in lanes 1 to 5 are obtained from MRSA strains SH21795, SH22058, SH19651, SH212, and SH13, respectively. M, molecular weight markers.

of mecA gene and mec regulator genes, S. aureus and S. epidermidis strains were classified into five types (I-V), as shown in Table 2. In 28 MSSA and 2 methicillin-susceptible S. epidermidis (MSSE), neither mecA nor mec regulator genes was detected (type V). The mecR1 (mecR1A and mecR1B) and mecI-positive pattern (type I) were detected in 94.3% of MRSA clinical isolates, 88.5% of MRSA environmental isolates, and 83.9% of MRSE. A few strains were assigned to type II lacking mecI, type III possessing only mecR1A, or type IV in which no mec regulator genes were detectable. In clinical isolates of MRSA and MRSE, all the four types were observed, whereas environmental MRSA strains were classified as either type I or IV. Two MRSA clinical strains that belonged to type II were isolated from different specimens (pus and sputum) derived from the same patient, and two environmental MRSA strains that belonged to type IV were isolated from different sites in the same department. Besides these, all MRSA and MRSE strains grouped into types II, III and IV were isolated from different patients.

None of MSSA and MSSE strains possessed mecA and they were all susceptible to methicillin. However, a few MRSA and MRSE strains also showed phenotypic oxacillin-susceptibility and they belonged to type III or IV in *S. aureus*, and type I or III in *S. epidermidis* (Table 2).

RFLP analysis with Mse I was performed for 164 strains in which PCR product of mecI was obtained. Most of the staphylococcal strains showed RFLP pattern 1 (160 strains, 97.6%) which represented the absence of novel MseI site, and yielded fragments of 303 bp and 122 bp, and smaller fragments (37 bp and 13 bp) (pattern 1) (Fig. 3, lanes 6 and 7). Three MRSA clinical strains isolated from different patients in 1993 and 1994 showed RFLP pattern 2 which suggested mutation 1, consisted of fragments of 206 bp, 122 bp an 95 bp, and smaller fragments (Fig. 3, lanes 8 and 9). From a single MRSA (a clinical isolate in 1993), a slightly smaller sized mecI gene-PCR product was obtained (Fig. 3, lane 5). Restriction fragments of the PCR product digested by Mse I contained a DNA fragment with the approximate size of 280 bp (pattern 3) (Fig. 3, lane 10) instead of 303 bp-fragment which was seen in pattern 1. All the strains that showed RLFP patterns 2 and 3 possessed both mecR1A and mecR1B.

DISCUSSION

It has been accepted that intact mec regulator genes strongly repress the transcription of mecA gene, whereas the repression activity is lost by mutation or deletion which occurred in mec regulator genes, resulting in constitutive production of PBP-2a [15, 17, 18]. Suzuki and colleagues [18] studied in detail the genomic diversity in the mec regulator gene and distinguished four different groups; group A, strains carried the whole mec regulator region including mecI which contained a point mutation; group B, the strain in which mecI only was deleted; group C, the strain in which mecI and nearly half of the 3' end of mecRI were deleted; group D, strains lacked almost all mecI and mecR1 genes. In 26 epidemic MRSA strains isolated from various countries, groups A (59.5%) and C (40.5%) were predominant, whereas groups B and D were not detected. In contrast, all the four groups were found in 27 MR-CNS isolated in Japan, although no S. epidermidis was assigned for group D. Our present study using recent isolates from a single hospital in Japan indicated that the major type of mec regulator gene variation was type I (corresponding to group A by Suzuki and colleagues [18]). Moreover, our results demonstrated that greater genomic variation existed in MRSA and MRSE than reported previously; types II and IV (groups B and D, respectively) were found in MRSA and type IV (group D) was detected in MRSE in our study.

Results in RFLP analysis of *mecI* revealed that most *mecI* gene tested in our study did not harbour the same point mutations as those identified previously [18]. This result suggested the presence of mutations in other sites of *mecI* which cannot be detected by the *Mse* I digestion. However, it was of note that the *mecI*-mutation 1 (RFLP pattern 2) found in 3 MRSA strains in our study had been identified in 3 MRSA isolates derived from different European countries. This might imply either that the mutation 1 readily occurs in MRSA strains worldwide, or that an MRSA clone with this mutation has disseminated worldwide.

The *Mse* I-RFLP pattern 3 which was observed in only one MRSA strain appears to represent a novel type of genomic diversity in *mecI*. The size of *mecI*-PCR product showing pattern 3 was slightly smaller than those of all other strains, and the smaller size of DNA fragment was generated instead of 303 bpfragment which was observed in RFLP pattern 1. This finding suggested that the *mecI* gene that showed pattern 3 had an internal deletion with approximate length of 20 bp, which may be confirmed by further sequence analysis.

Since the emergence of MRSA in England in 1961 [23], distribution of MRSA has been expanding globally. Several speculations have been presented for origin and evolution of mec locus including mecA and mec regulator genes. Kreiswirth and colleagues [24] hypothesized that all MRSA evolved vertically from a single progenitor clone which had acquired mecA. Other studies also supported the idea of a clonal origin of European MRSA strains [25-27]. However, occurrence of horizontal transfer of mecA has also been suggested by studies on diversity of mecA and chromosomal DNA [28]. Musser and Kapur [29], using multilocus enzyme electrophoresis, found that while MRSA derived from 1960s showed a single genotype, a fact favouring clonal dissemination, more recent isolates showed extensive diversity of chromosome, suggesting polyclonality. Hence they hypothesized that multiple episodes of horizontal transfer of mec locus and recombination have occurred recently.

Similar change in distribution of mec regulator genes has also been observed; most of the MRSA isolated from various countries in 1960s lacked the mec regulator genes, whereas these genes existed in most strains isolated since 1980 [17-19]. Consistent with these observations, the presence of mecI and mecR1 was demonstrated in most clinical and environmental MRSA in our present study. These findings may indicate that old strains had only mecA gene and acquired the mec regulator genes later. Alternatively, MRSA with mec regulator genes surpassed those with mecA only after the period when both types of MRSA existed. Archer and colleagues [19] recently presented a hypothesis that mec regulator genes including those with deletion or mutation have been transferred from CNS to S. aureus. However, it is still unclear why MRSA with dysfunctional mec regulator genes became predominant recently in place of old type MRSA that possessed only mecA. It was noted in our study that 3 MRSA and 1 MRSE strains lacking mecI (type III and IV) were phenotypically oxacillin-susceptible. Thus, the involvement of other unknown mechanism for regulation of PBP-2a production is suggested. Further investigation of the distribution, genetic divergence and function of mec regulator genes will be required to understand the evolution of MRSA and the basis for its global spread.

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