Molecular cloning and nucleotide sequence determination of the regulator region of mecA gene in methicillin-resistant Staphylococcus aureus (MRSA)

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Molecular cloning and nucleotide sequence analysis were performed for the identification of the regulator genes of methicillin resistance in the genome of a MRSA strain N315. Two open reading frames (orfs) were identified in the 5'-flanking region of the mecA gene. Predicted amino acid sequences of these orfs showed extensive homology to the co-inducer and the repressor protein of the penicillinase (PCase) production in Staphylococcus aureus as well as in Bacillus licheniformis. These orfs are considered to encode putative co-inducer and repressor proteins specific for the regulation of methicillin resistance in MRSA.

mecA; Repressor; Co-inducer; Methicillin resistance; Staphylococcus aureus

1. INTRODUCTION

MRSA expresses methicillin resistance by producing the specific penicillin-binding protein, PBP2', that has a decreased binding affinity to β -lactam antibiotics [1-3]. Although the structure gene, mecA, coding for PBP2' is located on the chromosome [4], it is shown to be regulated by the PCase plamid which is present in most of the MRSA strains [5]. Elimination of the PCase plasmid from the cell changes the inducible mode of PBP2' production to the constitutive one in those MRSA strains [5]. Recently, however, Tesch et al. have isolated from a clinical strain of S. epidermidis a genomic clone carrying the mecA gene, and have identified the genomic region responsible for down-regulation of PBP2' towards the upstream of the mecA gene [6]. On the other hand, we have identified a group of MRSA strains, represented by a strain N315, which retains inducible production of PBP2' even after elimination of the PCase plasmid from the cell [7]. In this study, we looked for the regulator genes of the mecA gene using the chromosomal DNA of N315 as a source for cloning as well as for direct genomic nucleotide sequencing.

2. EXPERIMENTAL

2.1. Bacterial strains

Three strains were used for nucleotide sequence determination. The stain N315 is a MRSA strain isolated in Japan in 1982, and was

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described previously [7]. N315P-ZR is a highly methicillin-resistant mutant of N315 [7], PBP2' production of which is derepressed (unpublished data). MR108 is a MRSA strain whose production of PBP2' is inducible but becomes constitutive when its PCase plamid is eliminated [8].

2.2. Molecular cloning

The N315 genomic DNA library was constructed as follows. The DNA of the strain N315 was extracted as described previously [8]. DNA was partially digested with Sau3Al, and subjected to electrophoresis in an 0.8% agarose gel. The fragments larger than 4-kb were eluted from the gel, and ligated into the BanHI-cleaved vector plasmid pACYC177. Resultant recombinant plasmids were used to transform E. coli MC1061. The library was screened by colony hybridization using the ³²P-labelled 4.0-kb HindIII insert of pMR111 as a probe. The insert of the plasmid pMR111 is a genomic DNA fragment derived from MR108, and contains the whole mecA gene [8]. One of the obtained clones, pES1, contained a 6.0-kb genomic fragment whose restriction map is presented in Fig. 1A.

2.3. DNA sequencing

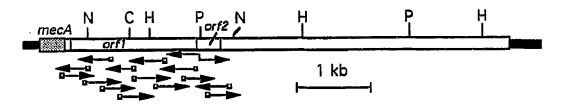
The nucleotide sequences of the 5'-flanking region of the mecA gene carried by the plasmids pES1 and pMR111, and the corresponding regions of genomic DNAs of N315, N315P-ZR and MR108 were determined by the dideoxynucleotide-termination method of Sanger et al. [9] using synthesized oligonucleotides as primers (Fig. 1A). In order to eliminate possible cloning artifacts represented on the plasmid clones, direct sequencing of the genomic DNAs was carried out using the PCR-amplified DNA fragments as templates for the sequence reaction after elution and purification of these by agarose gel electrophoresis.

3. RESULTS

3.1. Nucleotide sequence of the 5'-flanking region of mecA gene

Fig. 1A shows the sequencing strategy of plasmid pES1. Nucleotide sequence of a total of 2471 bases

Α



B

-35 -10 6D TTCATCANTATCCTCCTTATATA<u>AGACTACATTIGTAGTATA</u>T*TACAAATGTAGTAT*TTATGTCAAAATAAATGTATATATTTTCGTGATATGGAGGTGTAGAAGGTGTTATCATCTTTTT 240 ACAGTT AGGA GG TCATAT Mot LeuSerSerPheL GTA SD -10 -35 eumetleuSorIloIleSorSorLouLouThrfleCysVolIlePhoLouVolArgMotLouTyrlleLysTyrThrClnAsnfleMotSorHisLysIleTrpLouLouVolLouVolS erThrleuileProleuileProPheTyrlysileSorAsnPheThrPheSerlysAspMetMetAsnArgAsnVolSerAspThrThrSerSerVolSerHisMetLeuAspClyClnC AATCATCTGTTACGAAAGACTTAGCAATTAATGTTAATCAGTTTCACACCTCAAATATAACGTATATCTTTTGATTATGGGTATTTTGGTAGTTTGTTGTCTTATTATTATGATTA 600 inserSerValThrLysAsplouAlaileAsnValAsnClnPhcCluThrSerAsnIleThrTyrMetIleLeuLeuIleTrpValPhcClySerLeuLeuCysLeuPhcTyrMetIleL AGGCATTCCGACAAATTGATGTTATTAAAAGTTCGTCATTGGAATCGTCATATCTTAATGAACGACTTAAAGTATCTCAAAAGTATCACAAAGTATCACAAAGTATCACAAAGTATCACAAAGTACAATTGATCT ysAlaPheArgCinIleAspValIleLysSerSerSerLeuGluSerSerTyrLeuAsnGluArgLeuLysValCysGlnSerLysMetGlnPheTyrLysLysHisIleThrIleSerT yrSerSerAsnIleAspAsnProMetValPheGlyLeuValLysSerGlnIleValLeuProThrValValUalUaThrMetAsnAspLysGluIleGluTyrIleIleLeuHisGluL euSerHisYalLysSerHisAspLeu1lePheAsnGlnLeuTyrYalYalPheLysMetIlePheTrpPheAsnPrcAlaLeuTyr1leSerLysThrMetMetAspAsnAspCysGluL <u>AAGTATGTGATAGAAACGTTTTAAAAATTTTGAATCGCCATGAACATATACGTTATGGTGAACAATGCTATATAAAATGGTCTATTTTAAAAATGTCACGACACATAAAAATGCCCACAAT</u> 1080 yaValCysAspArgAsnValLeuLysIleLeuAsnArgHisGluHisIleArgTyrGlyCluSerIleLeuLysCysSerIleLeuLysSerGinHisIleAsnAsnValAlaAlaGinT yrLeulcuClyPheAsnSerAsnIlaLysGluArgvalLysTyrIleAlaLeuTyrAspSerMetProLysProAsnArgAsnLysArgIleValAlaTyrIleValCysSerIleSerL GCTTCACATGAAACAGCTAAAGAAGCTT erPheThr*** culculleGinAlaProleuteuSerAlaHisValGinGinAspLysTyrGluThrAsnValSerTyrLysLysLouAsnGlnLeuAlaProTyrPheLysGlyPheAspGlySorPheV alleutyrnanGluArqGluGlnAlatyrSerlletyrnanGluProGluSerLyaGlnArqTyrSerProAanSerThrTyrLyaIletyrLeuAlaLeuMotAlaPheAapGlnAanL euLeuSerLeuAsnHisThrCluCinCinTrpAspLysHisCinTyrProPheLysGiuTrpAsnClnAspClnAsnLeuAsnSerSerMetLysTyrSerValAsnTrpTyrTyrCluA snleuAsnlysHisLeuArqClnAspCluYallysSerTyrLeuAspleuIleCluTyrClyAsnCluCluIleSerClyAsnCluAsnTyrTrpAsnCluSerSerLeuLysIleSerA laileGluGlnVqlAsnLeuLeuLysAsnMetLysGlnHisAsnMetHisPheAspAsnLysAlaileGluLysValGluAsnSerMetThrLeuLysGlnLysAspThrTyrLysTyrV alGlyLysThrGlyThrGlyTleValAsnHisLysGluAlaAsnGlyTrpPheValGlyTyrValGluThrLysAspAsnThrTyrTyrPheAlaThrHisLeuLysGlyGluAspAsnA ΩĐ off2 CONATCCCGNANAGCNCNACNATTTCTGGCCTNTTTTAANAGNNTGGNGTTNATNTNNTATCGNTNATAANACGTNTGANTATCNTCTGCAGNATGGGAAGTTATGANTNTCATTTC 2040 laAsnGlyGluLysAlaGlnGlnIleSerGluArgIleLeuLysGluMetGluLeuIle*** MetAspAsnLysThrTyrCluIleSerSerAlaCluTrpCluValMetAsnIleIloTr pMetLysLysTyrAlaSerAlaAsnAsn1leIleGluGluIleGlnMetGlnLysAspTrpSerProLysThrIleArqThrLeuIleThrArgLeuTyrLysLysGlyPheileAspAr Hi TANAAAAGACAAMAATTTTTCAATATTACTCTCTTGTAGAAGAAGTGATATAAAATATAAAACTCTAAAAACTTTATCAATAAAGTATACAAAGGCGGTTTCAATTCAATTCACTTGTCTT 2280 gLysLysAspAsnLysIlePheGlnTyrTyrSerLeuValGluGluSerAspIleLysTyrLysThrSerLysAsnPheIleAsnLysValTyrLysGlyGlyPheAsnSerLeuValLe AAACTTTGTAGAAAAAGATCTATCACAAGATGAAATAGAAATTGAGAAATTATATTGAATAAAATAAAATTGTATGTTTTACAACAATACA<u>TAGAAACAGAGAAACAATGA</u>A uAsnPheValCluLysCluAspLeuSerClnAspCluIleCluCluLeuArgAsnIleLeuAsnLysLys***

Fig. 1. (A) Restriction map of pES1 and sequencing strategies. N, Ndel; C, Clal; H, HindIII; P, Pst1. The positions of primers are designated by open boxes. (B) Nucleotide sequences of the 5'-flanking region of the mecA gene. The nucleotide and predicted amino acid sequences of N315 are fully spelled out. The different nucleotide sequence of MR108 and its amino acid translation is shown above the N315 sequence. Sequences of MR108 beyond the position 1228 have not been determined yet. Possible Shine-Dalgarno sequences and promoter sequences (-10, -35) are shown in italics. The positions of inverted repeats are underlined. These sequence data will appear in the EMBL Nucleotide Sequence Data Library under the accession number X63598.

 $\mathtt{GTCG}\underline{\mathsf{TGA}}\underline{\mathsf{ATA}}\underline{\mathsf{TTTCCTCTGTTTT}}\underline{\mathsf{TA}}\mathsf{GTTGA}\underline{\mathsf{AAAAAAATTAACCGAAAGCCTCAATGCAAGTCTTGATTAAA}$

A

- 1 MISSFIMISIISSLITICVIFLVRMLYIKYTQNIMSHKIWLLVLVSTLIPLIPFYKISNFTFSK-DMMNRIVSDTTSSVSHMLDGQQSSVTKDLAINVNQFETSNITYMILLIWYFGSLL

 1 MAKLLIMSIVSFCFIFLLIFFRYILKRYFNYMLNYKVWYLTLLAGLIPFIPI-KFSLFKFNINNNQAPTVESKSHDLNHNINT--TKPIQEFATDIHKFNMDSIDNISTVIWIVLVII

 120 CLFYMIKAFRQIDVIKSSSLESSYLNERLKVCQSKMQF---YKKHITISYSSNIDNPMVFGLVKSQIVLPTVVVETMIDKELEYILLHELSHVKSHDLIFNQLYVVFKMIFMFNPALYIS

 117 LSFKFLKALLYLKYLKKQSL---YLNENEKNKIDTILFNHQYKKNIVIRKAETIQSPITFWYGKYILLIPSSYFKSVIDKRLKYILHEYAHAKNROTTHLIIFNIFSIIMSYNPIVHIV

 237 KTMMDNDCEKVCDRNVLKILNRHEHIRYGESILKCSILKSQHINNVAAQYLLGFNSNIKERVKYIALYDSMPKPNRNKRIVAYIVCSISLLIQAP-LLSAHVQQDKYETNV--SYKKLNQ

 234 KRKIIHDNEVEADRFVLNNINNNEFKTYAESIMDSVLNVPFFNKNILSHSFNGKKSLLKRRLINIKE-ANLKKQSKLILIFICIFTFLLMVIQSQFLMGQSITDYNYKKPLHNDYQILDK

 354 LAPYFKGFDGSFVLYNEREQAYSIYNEPESKQRYSPNSTYKIYLALMAFDQNLLSLNHTEQQWDKHQYFFKEWNQDQNLNSSMKYSVNKYYENLNKHLRQDEVKSYLDLIEYGNEEISGN

 353 -SKIFGSNSGSFVMYSMKKDKYYIYNEKESRKRYSPNSTYKIYLAMFGLDRHIINDENSRMSWNHKHYFFDAWNKEQDLNTAMQNSVNWYFERISDQIPKNYTATQLKQLMYGNKNLGSY

 474 ENYWNESSLKISAIEQVNLLKNMKQHNMHFDNKAIEKVENSMTLKQKDTYKYVGKTGTGIVNHKEANGWFVGYVETKDNTYYFATHLKGEDNANGEKAQQISERILKEMELI

 472 KSYMMEDSLKISNIEQVNLLKNMKQHNMHFDNKAIEKVENSMTLKQKDTYKYVGKTGTGIVNHKEANGWFVGYVITNHDKYYFATHL-SDGKØSGKNAELISEKILKEMOVINGQ
- B
- 1' MONKTYEISSAEWEVMNIINMKKYASANNIIEEIQMOKOWSPKTIRTLITRLYKKGFIDRKKONKIFQYYSLVEESDIKYKTSKNFINKVYKGGFNSLVLNFVEKEDLSQDEIEELRNIL
 1" MANKQVEISMAEWDVMNIINGKKSVSANEIVVEIQKYKEVSDKTIRTLITRLYKKEIIKRYKSENIYFYSSNIKEDDIKMKTAKTFLNKLYGGDMKSLVLNFAKNEELNNKEIEELRDIL
- 121' NKK
- 121" NDISKK

Fig. 2. Comparison of amino acid sequences. (A) Predicted amino acid sequence of orf1 vs. BlaR1. Arrowheads indicate the presumed penicillin-binding motif. (B) Predicted amino acid sequence of orf2 vs. BlaI. The upper sequences are the predicted ones of N315 in both panels. Asterisks show identical amino acids. Dots show the substitution of amino acids by those of the same nature.

containing 125 bases of the 5' end of mecA gene and its 5'-flanking region was determined (Fig. 1B). The nucleotide sequence was free from cloning artifacts because nucleotide sequencing of the corresponding region of the N315 genomic DNA showed complete agreement with that of pES1. Two orfs were found in the direction of transcription which is opposite to that of the mecA gene (shown as orf1 and orf2 in Fig. 1B). The nucleotide sequencing of N315P-ZR, however, revealed one base substitution from guanine to adenine at position 2160 in orf2 of N315 which caused amino acid substitution from arginine to histidine in the predicted protein (Fig. 1B). In the case of MR 108, a more drastic difference was observed; the nucleotide sequence was identical with that of N315 until position 1199 in the orfl, beyond that point, however, it was replaced by a totally different sequence, causing the orfl of MR108 terminate prematurely at position 1208 (Fig. 1B). Moreover, the DNA region corresponding to the orf2 was not present in the genome of MR108 which was shown by Southern hybridization using a PCR-amplified orf2 DNA as a probe (data not shown).

3.2. Predicted proteins encoded by the orf1

The predicted protein encoded by orfl consists of 585 amino acids (predicted molecular weight = 68,501.92). The protein is 34% homologous to BlaRl of S. aureus [10], and is 31% homologous to PenJ (or BlaRl) of B. licheniformis [11,12], both of which are known as the co-inducer protein involved in the regulation of penicillinase production in the respective bacterial species (Fig. 2). Especially, the predicted amino acids corre-

sponding to the penicillin-binding site of BlaR1 with a motif of [serine-X-X-lysine] (indicated by arrowheads) shows strong homology. Hydropathy profile further supported similarity of the predicted protein to BlaR1 (Fig. 3). The predicted protein shared the characteristic five hydrophobic peaks (indicated by arrows) and an increased hydrophylicity towards the carboxyl terminus.

3.3. Predicted protein encoded by the orf2

The second orf is predicted to encode a protein composed of 123 amino acids (predicted molecular weight = 14,789.74), which characteristically contains a high percentage of basic amino acids; it contains 18 lysine (14.63%) and 4 arginine residues (3.25%). The amino acid sequence of the predicted protein was very homologous to BlaI [10] of S. aureaus (61%) and to PenI [13] of B. licheniformis (43%); both of these are repressor proteins of PCase regulation. The amino acid compositions of these three proteins were also very similar to each other.

4. DISCUSSION

Fig. 4 summarizes the genomic organization of the *mec* region (containing the *mecA* gene and its 5'-flanking region) of N315 in comparison with that of the *bla* region of the *S. aureus* PCase plasmid pl258 [10]. It is noted that both the sizes and pattern of arrangement of the two orfs are evidently similar to those of *blaR1* and *bla1*. Based on the comparison of the nucleotide sequence of the *mecA* gene and its operator region with

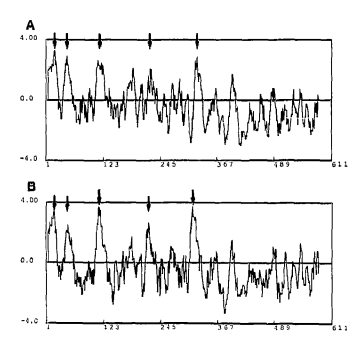


Fig. 3. Hydropathy profile of the predicted orfl-encoded protein (A) and of BlaR1 (B). The program of Kyte and Doolittle was used [15]. The mean hydropathy values of a moving segment of 9 amino acid residues are plotted at the midpoint of each segment. Arrows show conserved five hydrophobic peaks among two proteins.

those of E. coli PBP genes and a S. aureus PCase gene, Song et al. have proposed the idea that the 5' end and the operator region of the mecA structure gene is derived from the β -lactamase gene and the rest of the structure gene derived from the PBP gene [14]. This idea is further substantiated by the similarity of size, arrangement, and encoded proteins of the two orfs with those of the bla regulator genes (Fig. 4).

Although further studies on the gene products are required, it is very likely that the two orfs encode the co-inducer (MecR1) and the repressor (MecI) proteins which are specifically involved in the regulation of methicillin resistance. Premature termination of the orfl and the absence of orf2 in the genome of MR108 coincides well with this view as the expression of the mecA gene product is constitutive in MR108 after elimination of its PCase plasmid from the cell [8]. Most MRSA clinical isolates in recent years, represented by MR108, are constitutive producers of PBP2' in the absence of the PCase plasmid. In these cells, the genomic fragment containing orf2 seems to be absent. Tesch et al. also reported that the genomic region carrying an element for down-regulation of PBP2' production (probably identical with the orf2 according to the similarity of restriction enzyme map) is absent in the constitutive PBP2' producer strains [8]. The amino acid substitution identified in the orf2 of N315R-ZR may also be responsible for the observed de-repressed transcription of the mecA gene by partially inactivating the repressor function of the orf2-encoded protein.

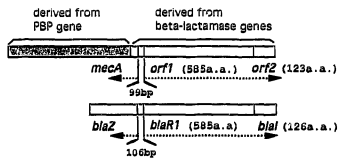


Fig. 4. Genomic organization of the *mec* region in comparison with the *bla* region. The stippled box shows the part of the *mecA* structure gene homologous to *E. coli* PBP genes [14]. Open boxes show DNA segments homologous to the corresponding segments of the *bla* region. The number of base pairs show the length of the operator of each region. Arrows indicate the direction of transcription.

It seems that N315 expresses the original sets of regulator proteins for methicillin resistance which are lost in the recent MRSA strains by deletion or other genetic alterations and are replaced by the regulator proteins of the PCase plasmid. It is conceivable that better inducibility and a higher resistance level towards various β -lactam antibiotics are achieved by the latter regulator system than the original one; thus, the selective expansion of the MRSA strains with the latter regulation system was brought about in the hospital environment.

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