Development of an Immunochromatographic Strip for Simple Detection of Penicillin-Binding Protein 2^\textsuperscript{\*V}

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Infections with methicillin-resistant Staphylococcus aureus (MRSA) and methicillin-resistant coagulase-negative Staphylococcus (MR-CNS) are a serious problem in hospitals because these bacteria produce penicillin-binding protein 2’ (PBP2’ or PBP2a), which shows low affinity to β-lactam antibiotics. Furthermore, the bacteria show resistance to a variety of antibiotics. Identification of these pathogens has been carried out mainly by the oxacillin susceptibility test, which takes several days to produce a reliable result. We developed a simple immunochromatographic test that enabled the detection of PBP2’ within about 20 min. Anti-PBP2’ monoclonal antibodies were produced by a hybridoma of recombinant PBP2’ (rPBP2’)-immunized mouse spleen cells and myeloma cells. The monoclonal antibodies reacted only with PBP2’ of whole-cell extracts and showed no detectable cross-reactivity with extracts from other bacterial species tested so far. One of the monoclonal antibodies was conjugated with gold colloid particles, which react with PBP2’, and another antibody was immobilized on a nitrocellulose membrane, which captures the PBP2’-gold colloid particle complex on a nitrocellulose strip. This strip was able to detect 1.0 ng of rPBP2’ or 2.8 × 10⁶ to 1.7 × 10⁷ CFU of MRSA cells. The cross-reactivity test using 15 bacterial species and a Candida albicans strain showed no detectable false-positive results. The accuracy of this method in the detection of MRSA and MR-CNS appeared to be 100%, compared with the results obtained by PCR amplification of the PBP2’ gene, mecA. This newly developed immunochromatographic test can be used for simple and accurate detection of PBP2’-producing cells in clinical laboratories.

Methicillin-resistant Staphylococcus aureus (MRSA) and methicillin-resistant coagulase-negative Staphylococcus (MR-CNS) are the major pathogens causing nosocomial infections and have been increasingly isolated in recent years from patients with community-acquired infections (23, 26, 29). These pathogens often show resistance to multiple classes of antibiotics, including β-lactams, macrolides, tetracyclines, aminoglycosides, and more (19). Therefore, the infections caused by MRSA and MR-CNS are difficult to eradicate. Studies have reported a high mortality rate among patients with MRSA septicemia compared with the mortality caused by methicillin-susceptible S. aureus (MSSA) (4, 25). A characteristic feature of MRSA and MR-CNS is the production of penicillin-binding protein 2’ (PBP2’ or PBP2a), an enzyme involved in the final step of peptidoglycan synthesis which consists of 668 amino acid residues and has a molecular mass of 76 kDa (5, 23a). PBP2’ is not inhibited by most β-lactam antibiotics, and thus, PBP2’-producing isolates show resistance to most, if not all, β-lactam antibiotics. Since the infection progresses very rapidly, a simple and accurate method to detect the PBP2’-producing pathogens is needed to facilitate appropriate chemotherapy and infection control.

Conventionally, methicillin-resistant Staphylococcus (MRS) has been detected by the oxacillin or cefoxitin susceptibility assay based on the disk diffusion method or a method to determine the MIC. However, these methods require several days to produce a reliable result. Alternatively, PCR amplification of the mecA gene encoding PBP2’ has been developed (8, 22). Although this method fulfills requirements for high speed, sensitivity, and specificity, it is costly, requires skilled personnel, and needs advanced equipment. Therefore, it is not practical for routine testing in clinical laboratories. Accordingly, a simple and accurate method of MRS detection has been long awaited.

This study reports on the development of an immunochromatographic test (ICT) that can fulfill the requirements for the detection of PBP2’ using novel monoclonal antibodies. The reliability of this method was assessed using clinical isolates of S. aureus and CNS.

MATERIALS AND METHODS

Expression and purification of His-tagged recombinant PBP2’ (rPBP2’). DNA was purified from Staphylococcus aureus (MRSA) strain 92-1191, isolated from the blood of a septicemia patient using the DNeasy blood and tissue kit (Qiagen K.K., Tokyo, Japan). The mecA gene was amplified by PCR using a pair of primers, 5’-TAATCTCAGCCCTTTTAAAGAGATTAAGT-3’ (nucleotides 70 to 89) and 5’-GTAATACCTGGGAATATCTATGTA-3’ (nucleotides 1990 to 2006), which contain the restriction sites NcoI and HindIII, respectively. PCR was carried out with PrimeSTAR HS DNA polymerase (Takara Bio Inc., Shiga, Japan) under the following conditions. After initial denaturation of DNA at 94°C for 3 min with a Gene Amp PCR System 9700 thermal cycler (Applied Biosystems Japan, Tokyo, Japan), a program was set at 94°C for 30 s, 57°C for 15 s, and 72°C for 15 s for a total of 30 cycles and then 72°C for 5 min for the termination. Amplified mecA DNA of 1,963 bp, corresponding to the mature PBP2’ protein with deletion of the N-terminal membrane-spanning segment (amino acid residues 24 to 668), was treated with NcoI...
and HindIII and was ligated to the pETBlue2 vector (Novagen, Madison, WI). *Escherichia coli* Novabac (DE3) (Novagen) cells were transformed with the plasmid by the heat shock method, and blue-white colonies were selected on Luria-Bertani (LB) agar (Nippon Becton, Dickinson and Company, Tokyo, Japan) containing 50 μg/ml of carbenicillin, 12.5 μg/ml of tetracycline, 80 μM isopropyl-β-D-thiogalactopyranoside (IPTG), and 70 μg/ml of 5-bromo-4-chloro-3-indolyl-β-D-galactopyranoside (X-Gal). E. coli strain Tuner(DE3)pLac (Novagen) was transformed with the pETBlue2-ne4 plasmid prepared with QIAprep Miniprep (Qiagen) from the white-colony cells.

The transformants were cultivated in LB broth containing 50 μg/ml of carbenicillin and 34 μg/ml of chloramphenicol at 37°C until absorbance at 578 nm reached 0.6. Expression of His-tagged rPB2 was induced by adding 0.5 mM IPTG, and the culture was then incubated at 22°C for an additional 24 h. Cells were harvested by centrifugation at 5,000 × g for 20 min and washed twice with a solution containing 50 mM Tris-HCl and 0.3 M NaCl (pH 8.0). The pellets were resuspended in a solution of 50 mM Tris-HCl, 0.3 M NaCl, and 10 mM imidazole (pH 8.0); subjected to ultrasonic oscillation for a total 10 min by 20-s exposure and 30-s intermittent cooling using an Astaron3000 sonicator (Misonix, Melville, NY) at a dial setting of 5, and then centrifuged at 8,000 × g for 20 min at 4°C. The clear soluble fraction was applied to a nickel-ion immobilized metal affinity chromatography (IMAC) resin column (Bio-Rad Laboratories, K.K., Tokyo, Japan) equilibrated with a solution of 50 mM Tris-HCl, 0.3 M NaCl, and 10 mM imidazole (pH 8.0) and then washed with the same solution; the column was eluted with a solution of 50 mM Tris-HCl, 0.3 M NaCl, and 250 mM imidazole (pH 8.0). Neighboring fractions containing the rPB2 protein were pooled and dialyzed against a large excess of phosphate-buffered saline (PBS) supplemented with 20 mM imidazole. The protein concentration was quantified by the Lowry method with bovine serum albumin (BSA) as the standard (Thermo Fisher Scientific K.K., Kanagawa, Japan).

**Electrophoresis and Western blotting.** Crude extracts of the transformant cells and purified rPB2 were subjected to sodium dodecyl sulphate-polyacrylamide gel (10%) electrophoresis (SDS-PAGE). Protein bands were visualized by staining with Coomassie brilliant blue (CBB) or Western blotting using anti-His tag antibody (Qiagen K.K., Japan). For Western blotting, proteins were electrotransferred onto a polyvinylidene difluoride (PVDF) membrane (Alto Corporation, Tokyo, Japan). The membrane was treated as follows: it was blocked with PBS containing 4.0% Block Ace (DS Pharma Biomedical Co., Ltd., Osaka, Japan) for 0.1% Tween 20 for 1 h at 24°C, washed 4 times with PBS containing 0.1% Tween 20, incubated with 100 ng/ml of anti-His tag antibody at 4°C overnight, washed 4 times with PBS containing 0.1% Tween 20, incubated with 0.1% Tween 20, washed twice with PBS, blocked with 100 μl of PBS containing 2% skim milk for 1 h at 24°C, and washed with washing buffer; 100 μl of washing buffer containing 10% normal rabbit serum was added, and then the protein bands were blotted on a PVDF membrane. The membrane was treated by the following steps: it was blocked with PBS containing 4.0% Block Ace and 0.1% Tween 20 for 1 h at 24°C, incubated with PBS containing 0.4% Block Ace, 0.1% Tween 20, and 10% normal rabbit serum at 4°C overnight; washed 4 times with PBS containing 0.1% Tween 20; and incubated with 1,000-fold-diluted ascites fluid (clones 10G2 and 1G2) at 4°C for 0.5 h. Following washing and color development, the steps were similar to those described above.

**Cross-reactivity of the monoclonal antibody.** Cross-reactivity of the monoclonal antibody against microorganisms was assayed by indirect ELISA. Strains used were *Staphylococcus aureus* 744 (MSSA), *Streptococcus pyogenes* 92-1191, *Klebsiella pneumoniae* ATCC 9341, *Enterococcus faecalis* ATCC 29212, *Enterococcus faecium* NCTC12204, *Escherichia coli* Novih JC-2, *Klebsiella pneumoniae* NCTC8632, Servattus marcenes ATCC12648, Enterobacter cloacae IFO13535, Enterobacter aerogenes NCTC10006, Pseudomonas aeruginosa E-2, Acinetobacter calcoaceticus IFO12552, and Candida albicans ATCC 10231. These strains were grown on Trypticase soy agar supplemented with 5% sheep blood (Nippon Becton, Dickinson and Company) for 24 h. *Streptococcus pyogenes* gogens GT262, *Streptococcus agalactiae* GT1C1234, and group C *Streptococcus* clinical isolate no. 1 were cultivated on Trypticase soy agar with 5% sheep blood in the presence of 5% carbon dioxide at 35°C for 18 h. Cells were suspended in 1.0 ml of sterile deionized water. Absorbance at 578 nm was adjusted to 2.0, and the suspension was mixed with an equal volume of 0.2 M NaOH, boiled for 5 min, and neutralized with 0.5 ml of 0.5 M KH₂PO₄. Microtiter plates were coated with 50 μl of coating buffer containing 10% sheep serum for 20 min at 4°C, washed twice with PBS, blocked with 100 μl of PBS containing 2% skim milk for 1 h at 24°C, and washed with washing buffer; 100 μl of washing buffer containing 10% normal rabbit serum was added, and then the plates were incubated at 24°C for 1 h and then washed with washing buffer, after which 50 μl of 1,000 ng/ml anti-rPB2 monoclonal antibody was added and the plates were incubated at 24°C for 0.5 h. This material was subjected to the indirect ELISA described above.

**Preparation of the immunochromatographic strip.** Anti-rPB2 monoclonal IgM (clone 1G12) and anti-mouse IgG antibody (Vector Laboratories, Inc., Burlingame, CA) were immobilized on a 2.5- by 0.5-cm nitrocellulose membrane (Schleicher & Schuell, Keene, NH) at 1.0 and 1.5 cm, respectively, from the proximal end (see Fig. 4). These two lines served as a test line and a control line, respectively. The membrane was blocked with 0.5% casein solution for 20 min. After washing, it was soaked in 5% sucrose solution and air dried. To prepare the gold colloid-conjugated IgG, 1.0 ml of 0.06% gold colloid suspension (40 nm) (Tanaka Kikinzoku Kogyo, Tokyo, Japan) and 0.1 ml of 60 μg/ml anti-rPB2 monoclonal IgG (clone 10G2) were mixed, incubated for 20 min, and then blocked by addition of 1.0% sodium casein. The mixture was centrifuged at 14,000 × g for 15 min to remove free IgG, and the pellet was resuspended in 0.5 ml of 10 mM Tris-HCl buffer containing 0.1% sodium casein and 10% (v/v)-trehalose dihydrate, pH 8.2. Preparation of the immunochromatographic strip has been described in the supplemental material of reference 15.

**Sensitivity and cross-reactivity testing of ICT.** The sensitivity of ICT was evaluated using quantified rPB2 (see above) and extracts from three MRSA strains: MRSA ATCC 43300, MRSA 92-1191, and MRSA 70. The cells were grown on Trypticase soy agar supplemented with 5% sheep blood at 35°C for 18 h, suspended in sterile deionized water. The viable cell number was counted on Mueller-Hinton agar. The antigen was extracted by heating a mixture of 50 μl each of cell suspension (or rPB2 solution) and 0.2 M NaOH in a boiling water bath for 10 min, and the solution was neutralized by adding 25 μl of 0.5 M KH₂PO₄, containing 2.5% BSA. The rPB2 preparation was diluted with adjustment to 0.2 to 50 ng protein/50 μl/test, or the cell suspension was serially diluted in 10× increments to 10⁴ to 10⁶ CFU equivalent/test. A 50-μl aliquot of the extracts and 10 μl of gold colloid-conjugated antibody suspension were mixed in a well, and an immunochromatographic strip was left to stand in it. The
was reported previously (10).

S. aureus of ICT was tested using 62 and 53 clinical isolates of in the same manner as described above. For the evaluation of reproducibility of ICT, intra- and interassay comparisons were performed using 2 strains each of clinical isolates of MRSA and CNS. The reliability results only with the MRSA extracts, not with extracts

grown in the presence of IPTG, and the extracts were subjected to SDS-PAGE. A large amount of protein corresponding to about 75 kDa was confirmed (Fig. 1a, lane 2), and the affinity-purified material showed only a single protein band corresponding to 75 kDa (Fig. 1a, lane 3). Western blotting using anti-His tag antibody also showed a single band corresponding to 75 kDa (Fig. 1b, lanes 2 and 3).

Specificity and cross-reactivity of novel monoclonal antibodies. Mice were immunized with the purified rPBP2, and antibody titer against rPBP2 was measured by indirect ELISA. All the sera from the immunized mice showed high titers of anti-rPBP2 antibody, whereas the preimmune sera showed no detectable anti-rPBP2 antibody (data not shown). The mouse which produced the highest titer of anti-rPBP2 antibody was selected for hybridoma cell production. Hybridoma cells prepared from the spleen cells of this mouse were screened for the production of monoclonal antibodies. Two hybridoma cells, 10G2 and 1G12, were cloned after two limited dilutions and were then injected into the mouse peritoneal cavity.

To evaluate the specificity of the anti-rPBP2 antibody, the reactivity of the mouse ascites fluid was examined by using purified rPBP2 and the crude membrane fraction from MRSA and MSSA cells by the Western blotting method. Both ascites showed only a single protein band with the membrane fraction from the MRSA cells and rPBP2 at the position corresponding to a mass of 75 kDa, whereas no protein band was seen in the membrane fraction from the MSSA cells (Fig. 2). To test the cross-reactivity of the antibodies, 16 bacterial species and one fungal species were examined by indirect ELISA. The purified monoclonal antibodies 1G12 and 10G2 showed positive results only with the MRSA extracts, not with extracts from the remaining 15 bacterial species and a fungal species (Fig. 3). The antibody isotype and IgG subclass of 1G12 and

RESULTS

Preparation of His-tagged rPBP2. The mecA gene encoding amino acid residues 24 to 668 of PBP2 was cloned on pETBlue2 with the sequence for hexahistidine residues. E. coli strain Tuner(DE3)/pLacI harboring pETBlue2-meca plasmid in the presence and absence of IPTG. The rPBP2 protein was purified by Ni-resin affinity chromatography from cells cultured in the presence of IPTG. Materials were subjected to 10% SDS-PAGE. (a) The soluble fraction (20 μg) and purified rPBP2 (5 μg) were applied. The gel was stained with CBB. (b) The soluble fraction (500 ng) and purified rPBP2 (100 ng) were applied and transferred to a PVDF membrane after electrophoresis. The membrane was stained by the Western blotting method using anti-His tag antibody. Lanes M, molecular mass markers; lanes 1, soluble fraction from uninduced cells; lanes 2, soluble fraction from IPTG-induced cells; lanes 3, affinity purified rPBP2.

![Image 1](http://cvi.asm.org/Downloadedfrom)

![Image 2](http://cvi.asm.org/Downloadedfrom)
the cell density of 10^8 CFU/test and 10^7 CFU/test, respectively, from 15 bacterial species and one strain of MRSA ATCC 43300, MRSA 70, and MRSA 92-1191, respectively.

10G2 were found to be IgM and IgG1, respectively, as tested with specific antibodies by ELISA. 1G12 were found to be IgM and IgG1, respectively, as tested with specific antibodies by ELISA.

Performance of ICT. The ICT system was prepared using two monoclonal antibodies, 10G2 and 1G12. The 10G2 antibody was combined with a colloidal gold particle in a complex that served as the detector of rPBP2', and the 1G12 antibody was immobilized on a nitrocellulose membrane that captured the 10G2-gold colloid-rPBP2' complex, thereby forming an antigen sandwich with two monoclonal antibodies. To evaluate the sensitivity of the ICT, homogeneously purified rPBP2' protein was used first. The test strip was able to detect levels as low as 1.0 ng/50 μl/test of rPBP2' but was unable to detect protein at levels lower than 0.5 ng/test, macroscopically (Fig. 4). In the next experiment, whole-cell extracts from three MRSA strains were subjected to ICT. The results showed that the ICT was able to detect 1.7 x 10^5, 4.7 x 10^5, and 2.8 x 10^5 CFU/test of MRSA ATCC 43300, MRSA 70, and MRSA 92-1191, respectively.

To test for cross-reactivity of the present ICT, 16 strains from 15 bacterial species and one strain of Candida albicans at the cell density of 10^8 CFU/test and 10^7 CFU/test, respectively, were examined. The results clearly showed that none of these cells yielded false-positive reactions.

The thermostability of monoclonal antibody immobilized on the membrane strip was evaluated by an accelerated stability test at 60°C. The monoclonal antibody-immobilized strips were kept for 5, 10, 21, and 30 days, and the sensitivity was tested. We found that the sensitivity in detection of the MRSA 92-1191 cells was totally unchanged even after storage for 30 days (data not shown). To evaluate the reproducibility of the ICT, intra- and interassay comparisons were performed using two strains each of MRSA and MSSA from clinical sources. Five repeated experiments using two MRSA and two MSSA strains simultaneously showed identical positive and negative results, respectively. In addition, repeated experiments on 5 consecutive days also showed identical positive and negative results in 2 MRSA and 2 MSSA strains, respectively. It was firmly established that the present ICT has high reproducibility and is specific to the PBP2'-producing cells.

Reliability of the present ICT and comparison with PCR and LAT. To test the reliability of the newly developed ICT, 62 clinical isolates of S. aureus and 53 clinical isolates of CNS were examined. One loopful (1 μl) of cells was subjected to ICT, and the results were compared with those of PCR and LAT (Table 1). As mentioned above, PCR detects the mecA gene and LAT detects the PBP2' protein. Among 62 strains of S. aureus tested, PCR detected the mecA gene from 37 strains, and 25 strains were mecA negative. The ICT and LAT showed results consistent with those of PCR without exception. Thus, it was established that ICT, PCR, and LAT are equally good methods for differentiating MRSA from MSSA. Among 53 strains of CNS, the present ICT method yielded PBP2' protein positivity in 38 strains and negativity in 15 strains, which matched perfectly with the results obtained by PCR. However, LAT was able to detect PBP2' in only 28 CNS strains and the remaining 25 strains appeared to be negative for PBP2'. Thus, LAT failed to detect 10 mecA-positive- and PBP2'-positive CNS strains, resulting in false-negative results.

DISCUSSION

Identification of MRSA and MR-CNS mainly relies on the β-lactam susceptibility test for oxacillin or cefoxitin (3). The biggest drawback of the culture method is that it is time-consuming, requiring more than 1 day to obtain a result from isolated colonies. PCR amplification of the marker gene(s) was adopted for rapid and accurate detection of MRSA and MR-CNS. Recently, a new real-time PCR method that amplifies the right-extremity sequences of the staphylococcal cassette chromosome mec (SCCmec) element and the 3' end of the S. aureus orfX gene has been developed (11). This method has enabled MRSA to be detected directly from clinical specimens. Though the PCR method is highly sensitive, accurate, and rapid, only a limited number of clinical laboratories have the opportunity to use and run the expensive equipment. More-

![FIG. 3. Cross-reactivity of monoclonal antibodies tested by indirect ELISA. The antigen of the test cells was immobilized in the microplate wells and subjected to ELISA using the monoclonal antibodies 10G2 and 1G12. Black columns, 1G12; gray columns, 10G2.](image1)

![FIG. 4. Sensitivity of ICT to the purified rPBP2'. Purified rPBP2' protein was diluted, with adjustment of the protein concentration from 0.2 to 50 ng/50 μl/test, and was subjected to ICT for 10 min.](image2)

![TABLE 1. Comparison of results obtained by PCR, LAT, and ICT using clinically isolated strains *(a)*](table1)
over, the method requires highly trained technical experts using costly reagents. The alternative method employed currently is to detect the PBP2\(^{\star}\) protein by the latex aggregation method (17). Though this is a time-saving method, the greatest drawback is that the target microorganism is limited to MRSA. Therefore, it is questionable whether the latex aggregation method can be applied to the detection of MR-CNS.

In this paper, we have reported a simple, time-saving method of ICT for detecting PBP2\(^{\star}\) using the extracts from pure culture cells. Since our ICT method employed monoclonal antibodies against rPBP2\(^{\star}\), the specificity was extremely high. In fact, the whole-cell extracts of MSSA and MRSA showed no detectable protein band except for PBP2\(^{\star}\).

The ICT method using the 1G12 and 10G2 monoclonal antibodies was able to detect levels from 2.8 \(\times 10^5\) to 1.7 \(\times 10^7\) CFU/test of MRSA. We found an approximately 60-fold difference in the level of PBP2\(^{\star}\) expression among the MRSA strains tested so far. In fact, the reference strain ATCC 43300 contains a heterogeneous population of cells in regard to oxacillin susceptibility (14). On the other hand, the clinically isolated strains 92-1191 and 70 showed stable, high oxacillin resistance, with an MIC of \(\geq 128\) \(\mu\)g/ml.

Cross-reactivity testing of the present ICT revealed that there was absolutely no cross-reaction when 15 different species of bacteria and a Candida albicans strain were tested. One important point to be considered is that Staphylococcus aureus produces protein A (6), which reacts with the Fc region of IgG, potentially causing false-positive results in immunoassay systems (18, 20). We used the monoclonal antibodies to IgG1 and IgM for the present ICT. Fortunately, it is generally accepted that the binding affinity of protein A to mouse IgG1 and IgM is lower than that to other subclasses of mouse IgG (7).

Another point to be clarified would be the stability of IgM antibody on the membrane strip because it is said that IgM is generally less stable than IgG. Thus, we tested the stability of our IgM monoclonal antibody (clone 1G12) on the immobilized membrane strip by the accelerated stability test at 60°C up to 30 days. We found no detectable deterioration of our IgM antibody in the reaction with the PBP2\(^{\star}\)-producing cells. It was generally accepted that stability at 60°C for 3 weeks is equivalent to that for 2 years of storage at room temperature (2, 21). It was reported that a sugar additive such as sucrose or trehalose protects proteins from deterioration under the dried condition (21). Our membrane strips were soaked in 3% sucrose solution before being dried.

The performance of ICT was first tested using a total of 62 strains of clinically isolated S. aureus, and the results were compared with those of PCR and commercially available LAT. ICT, LAT, and PCR all detected PBP2\(^{\star}\) positivity in 37 strains, thereby indicating that all three methods were equally effective for detecting PBP2\(^{\star}\)-positive S. aureus. The present result confirmed previous reports that the sensitivity and specificity of LAT were 96.7 to 100% and 99.2 to 100%, respectively, in S. aureus (1, 24, 28). Since LAT is recommended by the manufacturer for use in detecting MRSA, the performance of this kit may be acceptable.

Similarly, 53 strains of CNS were tested and ICT detected PBP2\(^{\star}\) in 38 strains with sensitivity and specificity 100% identical to those obtained by PCR. On the other hand, LAT showed only 73.6% sensitivity and 100% specificity. MR-CNS strains often show heterogeneous levels of mecA gene expression, and therefore, the use of an increased amount of cells for the LAT was evaluated (9, 13). In fact, it has been reported that the use of 2.5-times-more cells and extension of incubation to 10 min improved the sensitivity of LAT for CNS from 68.4 to 95.7% (27). However, the application of this modified method to S. aureus caused false-positive results (16), which were most likely attributable to reaction with protein A. Our ICT method was able to detect PBP2\(^{\star}\) from MRSA and MR-CNS correctly, because of the improvement in the detection limits compared with those of LAT.

In conclusion, the findings of the present study show that the newly developed ICT is a highly sensitive, accurate, and rapid method for the detection of PBP2\(^{\star}\)-producing Staphylococcus spp. from pure cultures. Therefore, this method is suitable for use in routine tests at clinical laboratories.

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REFERENCES


