

Rapid Detection of Methicillin-Resistant *Staphylococcus aureus* Directly from Sterile or Nonsterile Clinical Samples by a New Molecular Assay

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A rapid procedure was developed for detection and identification of methicillin-resistant *Staphylococcus aureus* (MRSA) directly from sterile sites or mixed flora samples (e.g., nose or inguinal swabs). After a rapid conditioning of samples, the method consists of two main steps: (i) immunomagnetic enrichment in *S. aureus* and (ii) amplification-detection profile on DNA extracts using multiplex quantitative PCR (5'-exonuclease qPCR, TaqMan). The triplex qPCR assay measures simultaneously the following targets: (i) *mecA* gene, conferring methicillin resistance, common to both *S. aureus* and *Staphylococcus epidermidis*; (ii) *femA* gene from *S. aureus*; and (iii) *femA* gene from *S. epidermidis*. This quantitative approach allows discrimination of the origin of the measured *mecA* signal. qPCR data were calibrated using two reference strains (MRSA and methicillin-resistant *S. epidermidis*) processed in parallel to clinical samples. This 96-well format assay allowed analysis of 30 swab samples per run and detection of the presence of MRSA with exquisite sensitivity compared to optimal culture-based techniques. The complete protocol may provide results in less than 6 h (while standard procedure needs 2 to 3 days), thus allowing prompt and cost-effective implementation of contact precautions.

Staphylococcus aureus is a major pathogen responsible for nosocomial and community-acquired infections. Since 1960, the emergence of multiresistant strains of *S. aureus* carrying resistance to methicillin (MRSA) and to most currently available antibiotics (4, 23) has dramatically narrowed the therapeutic arsenal to the exclusive use of glycopeptides (such as vancomycin and teicoplanin) as the mainstay of MRSA treatment. Unfortunately, vancomycin overuse has led to the emergence of MRSA strains with decreased susceptibilities to glycopeptides (22, 37).

The presence of MRSA in an institution is clearly paralleled by an increased number of bacteremias due to MRSA (41), which carry a threefold attributable cost and a threefold excess length of hospital stay when compared with methicillin-susceptible *S. aureus* (MSSA) bacteremia (1). These data, together with successful containment effort programs (6, 7, 9, 16, 18, 34, 36), prompt screening of high-risk patients even in a setting of high endemicity (43). Screening for MRSA is a key component of successful infection control strategies (11, 38, 43) aiming to identify hidden reservoirs of MRSA patients and appropriately apply isolation precautions (17, 40). To this end, infection control benefits from automated alerts upon admission in order to identify patients who have been previously colonized by MRSA (39). Early detection of MRSA carriers is crucial not only for infection control (11) but also for therapeutic decision

with last-line antibiotics against MRSA, e.g., glycopeptides and oxazolidinones (49).

Rapid detection of MRSA by standard clinical microbiological procedures is tedious and time consuming, since it first requires identification of isolated *S. aureus* colonies within mixed flora samples before assessing their level of methicillin resistance. The development of selective media containing antibiotics and phenol red has provided better sensitivity than conventional agar-based cultures after 48 h of incubation, but at the expense of a longer turnaround time (10, 46, 51).

Direct or indirect particle agglutination assays using antibody-coated beads offer a rapid alternative to oxacillin susceptibility testing. For example, MRSA-Screen (Denka Seiken, Tokyo, Japan) provides sensitive and specific immunodetection of MRSA in a pure culture by using anti-PBP2' antibodies, which is similar to standard oxacillin disk diffusion or oxacillin-salt agar screening (5, 24, 42). However, the specific immunodetection of MRSA based on PBP2' cannot be performed in the presence of methicillin-resistant *Staphylococcus epidermidis* (MRSE), which is a frequent commensal contaminant of mixed flora samples (5). Indeed, the high level of sequence homology of the *mecA* gene present in *S. aureus*, *S. epidermidis*, and potentially other coagulase-negative staphylococci (CNS) species (52) precludes discrimination of methicillin-resistant strains of *S. aureus* from those of *S. epidermidis* (24).

Since the *mecA* gene, encoding the low-affinity penicillin-binding protein (PBP2') (44), represents the "gold standard" for detecting methicillin resistance (32), several assays based on the direct detection of the *mecA* gene have been described, using chemiluminescent probes (53), amplification methods such as PCR (42), or cycling probe technology (12). A PCR

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TABLE 1. List and characteristics of oligonucleotides used in the triplex qPCR assay

Gene and primer or probe name ^a	Sequence (5'→3')	Length	5' dye ^b	GenBank accession no. (reference)
<i>mecA</i>				X52593 (44)
F <i>mecA</i>	CATTGATCGCAACGTTCAATTT	22		
R <i>mecA</i>	TGGTCTTCTGCATTCCTGGA	21		
P <i>mecA</i>	TGGAAGTTAGATTGGGATCATAGCGTCAT	29	JOE	
<i>femA-SA</i>				X17688 (3)
F <i>femA-SA</i>	TGCCTTTACAGATAGCATGCCA	22		
R <i>femA-SA</i>	AGTAAGTAAGCAAGCTGCAATGACC	25		
P <i>femA-SA</i>	TCATTTACGCAAACCTGTTGGCCACTATG	29	FAM	
<i>femA-SE</i>				U23713 (2)
F <i>femA-SE</i>	CAACTCGATGCAAATCAGCAA	21		
R <i>femA-SE</i>	GAACCGCATAGCTCCCTGC	19		
P <i>femA-SE</i>	TACTACGCTGGTGGAACTTCAAATCGTTATCG	32	TET	

^a *femA-SA*, *S. aureus femA*; *femA-SE*, *S. epidermidis femA*.

^b All probes are quenched by TAMRA bound to the 3' end.

immunoassay was reported by Towner et al., based on the amplification and immunodetection of *mecA* and *femB* amplicons, and performed after overnight culture enrichment of clinical samples. This technique outperforms conventional detection methods (48) by providing rapid immunodetection and avoiding the use of acrylamide gel. In another approach, PCR-amplified products were detected with low-density oligoarrays, providing detection of several targets during the same hybridization step (13). Other more elaborated multiplexed PCR, using different targets (*coagulase* and *femA* genes) (25, 50) or *S. aureus* toxins (47), yielded promising results with good specificity. Promising results were also obtained by using either triplex PCR assays based on the detection of *S. aureus* rRNA, *mecA*, and *nuc* (27, 29) or *clfA* genes (30) or by in situ hybridization performed on blood culture bottles (35).

The recent introduction of multiplexed real-time qPCR techniques (19, 26) combining accurate identification with limits of detection close to a single gene copy/sample provided a significant technological advantage for the rapid and large-scale identification of various microorganisms (8, 28, 31). This report describes a novel immuno-qPCR procedure allowing rapid detection of MRSA from mixed flora samples. The procedure consists in a direct one-step enrichment of MRSA present in either nasal or inguinal swabs, followed by DNA extraction of immunocaptured bacteria and their identification by a triplex qPCR. The specificity of MRSA molecular identification is based on the presence of the *mecA* gene and the presence of an *S. aureus*-specific *femA* signal that does not cross-react with other bacterial species, including *S. epidermidis*. This novel qPCR assay allows detection and identification of MRSA in less than 6 h after sample collection and may provide substantial benefits for infection control by allowing prompt and cost-effective implementation of contact precautions.

MATERIALS AND METHODS

Materials. Biotinylated mouse monoclonal antibodies raised against *S. aureus* protein A (anti-spa, mouse immunoglobulin G1 clone spa-27) were obtained from Fluka (Sigma Chemie, Buchs, Switzerland), and streptavidin-coated paramagnetic beads were obtained from Merck (Basel, Switzerland). qPCR kits,

primers, and TaqMan probes were purchased from Eurogentec (Seraing, Belgium).

Strains. Reference strains were ATCC33591 (MRSA), ATCC25923 or NCTC8530 (MSSA), and ATCC12228 (methicillin-susceptible *S. epidermidis* [MSSE]). The reference MRSE and a panel of various bacterial species were clinical isolates identified using NCCLS procedures at the Clinical Microbiology Laboratory (University Hospitals of Geneva, Geneva, Switzerland). For these assays, overnight cultures were washed in saline and quantified by turbidimetry (ATB 1550; API bioMérieux). Titters were adjusted using 10-fold dilutions in saline and checked by plating on agar.

Specimen collection. Samples consisting in nasal, inguinal, and wound swabs (Copanswabs; Copan Italia S.p.a, Brescia, Italy) were collected directly from patients admitted to University Hospitals of Geneva according to a previously defined infection control strategy (16, 39). Samples were simultaneously analyzed by molecular and conventional culture-based techniques. Swabs were suspended in 2 ml of colistin-salt broth (CS broth: brain-heart infusion with 10 µg of colistin/ml and 2.5% NaCl) and then divided equally and processed in parallel.

Identification of MRSA by standard microbiological procedure. One milliliter of CS broth was incubated at 35°C for 24 h and inoculated on phenylethyl alcohol agar plates. Suspect colonies were identified as MRSA based on Pastorex agglutination (Bio-Rad, Reinach, Switzerland), positive reaction on DNase agar, and growth on Mueller-Hinton (MH) oxacillin agar (6 µg of oxacillin/ml, according to NCCLS [33]). The presence of MRSA was confirmed using the Vitek 2 identification and susceptibility testing cards for gram-positives (bioMérieux, Marcy l'Etoile, France).

Immunomagnetic enrichment. The remaining 1 ml of CS broth was immediately processed by adding an optimized titer of biotinylated anti-spa antibody in the presence of 1% human serum albumin (HSA) (injectable grade; Swiss Red Cross, Bern, Switzerland). After a 30-min incubation at room temperature on a rotary shaker, the bacterial suspension was centrifuged for 10 min at 5,000 × g. The pellet was resuspended in 200 µl of phosphate-buffered saline (PBS) (Invitrogen; Basel, Switzerland) containing 1% human serum albumin (PBS-HSA) and supplemented with 20 µl of streptavidin-coated paramagnetic beads. After a 30-min incubation on a rotary shaker, the magnetic beads were rinsed twice in PBS-HSA on the magnetic holder. The efficiency of the immunomagnetic capture was assessed by plating diluted portions on MH plates and determining the number of CFU after 24 h of incubation.

Optimization of anti-spa antibody titer. To optimize the antibody titer, 10³ CFU of Cowan I (NCTC8530) was incubated with increasing antibody dilutions (1:166 to 1:5'400). The percentage of immunocaptured bacteria recovered from the original spiking was determined by viable counts on MH agar.

Efficiency of the recovery rate for various MRSA titers. The recovery rate as a function of the number of spiked bacteria was evaluated by incubating a range of MRSA suspensions (ATCC 33591) in the presence of the optimized anti-spa titer in 1 ml of CS broth. The yield was determined by viable counts.

Immunocapture of MRSA from mixed cultures. Various numbers of MRSA (ATCC 33591) were mixed with increasing titers of MRSE, yielding ratios ranging from 1:1 to 1:1,000. Immunocapture was evaluated by viable CFU counts

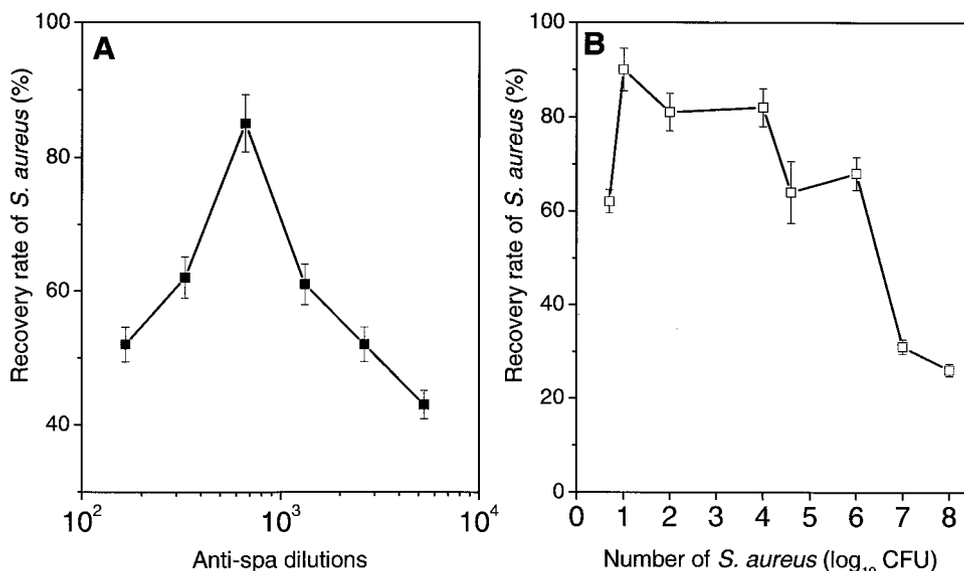


FIG. 1. Effect of antibody concentration on *S. aureus* recovery rate. Cowan I strain (NCTC8530; 10³ CFU) was used to evaluate the optimal concentration of anti-spa antibodies required for the enrichment step. (A) Recovery was determined by CFU counts of immunocaptured colonies. The curve showed maximal recovery at an antibody dilution of 1:666. (B). Using this optimized anti-spa dilution, recovery was assessed across >7 orders of magnitude in inoculum size. Average values \pm standard errors of the means for four experiments performed in duplicate are given.

performed on MH agar. Strains were visually discriminated by their pigmentation.

Bacterial lysis and DNA extraction. Immunocaptured bacteria were suspended in 400 μ l of chaotropic buffer (DNeasy kit; Qiagen) with 200 mg of glass beads (diameter, 100 to 200 μ m). Bacteria were lysed at 4°C in a bead-beater (Mixer Mill; Qiagen) using two homogenization cycles of 45 s each at a frequency of 30 Hertz. The liquid phase was cleared from beads and debris by a 10-min centrifugation at 5,000 \times g and then transferred into clean tubes containing 200 μ l of ethanol. The nucleic acids were purified according to the manufacturer's instructions (Qiagen), eluted in water, dried in an evaporator, and finally resuspended in 20 μ l of water.

DNA extraction from reference strains. Genomic DNA was extracted after a 10-min treatment at 37°C in TE containing 100 μ g of lysostaphin (Ambicin; Applied Microbiology, Tarrytown, N.Y.)/ml. DNA concentration and purity were assessed by spectrophotometry (45) using an Uvikon 942 (Kontron; Basel, Switzerland).

Oligonucleotide design and sequences. Sequences of primers and TaqMan probes are listed in Table 1. The design was performed using the software Primer Express version 1.0 (PE Biosystems, Foster City, Calif.). Since the sequences of both *S. aureus* and *S. epidermidis mecA* genes showed 100% homology, primers and TaqMan probe were determined based on the *S. aureus* gene (GenBank accession no. X52593). On the opposite, whereas *femA* nucleotide sequences are phylogenetically conserved among the staphylococci (13), *S. aureus femA* and *S. epidermidis femA* displayed only 78% homology as determined by LALVIEW (<http://www.expasy.ch/>). Primers and TaqMan probes were selected in regions displaying low homology with at least four mismatches between both *femA* oligonucleotide sequences (2, 3). Selected primers and probes were checked against GenBank to exclude potential cross-reacting sequences. Sequence identity between *S. aureus femA* and the *Streptococcus milleri* gene for millericin B (GenBank accession no. AF243359) was detected for both primer sequences as well as a homology of 25 of 29 nucleotides in the probe. However, the length of the generated amplicon (703 bp) was found to be excessive for reliable qPCR detection (not shown).

Nucleic acid detection by qPCR. For robustness issues, each analysis was performed in triplicate. Nucleic acids from reference MRSA, MSSA, MRSE, and MSSE strains (100 pg of genomic DNA in each well) were simultaneously analyzed in each run and used as standards to adjust threshold values (C_t). Amplification procedure on the SDS 7700 (PE Biosystems) was the following: t_1 , 2 min at 50°C; t_2 , 10 min at 95°C; t_3 , 15 s at 95°C; t_4 , 1 min at 60°C (t_3 and t_4 , repeated 50 times). The final volume of the PCR mixture was 20 μ l and contained all primers and TaqMan probes at 100 nM concentrations except the *mecA* probe, which was adjusted to 75 nM, following assay optimization (data not

shown). The cycling procedure was immediately started upon addition of 6 μ l of sample. The specificity of qPCR identification was assessed using the following pathogens (number of different strains): *Escherichia coli* (10), *Campylobacter fetus* (4), *Proteus vulgaris* (4), *Enterococcus faecalis* (11), *Enterococcus faecium* (4), *Enterobacter cloacae* (3), *Klebsiella pneumoniae* (3), *Streptococcus pneumoniae* (4), *Streptococcus alpha-haemolyticus* (4), *S. agalactiae* (2), *S. milleri* (3), *Pseudomonas aeruginosa* (10), *Stenotrophomonas maltophilia* (1), *Lactococcus cremoris* (1), *Neisseria* sp. (1), and *Staphylococcus haemolyticus* (10); also *Candida glabrata* (2) and *Candida albicans* (2). None of these different species yielded any false-positive qPCR signal.

Analysis of multiplex qPCR signals. After fluorescence background subtraction, we calibrated each run using the signals of the reference strains MRSA and MRSE. Detection thresholds were adjusted so that the *mecA* signal matched the corresponding *femA* signal for each reference strain. MRSA was considered present only when average C_t values (triplicates) met the following conditions: (i) average C_t for *mecA* of <50; (ii) average C_t for *S. aureus femA* of <50; and (iii) C_t for *mecA* that is less than the C_t for *S. epidermidis femA*.

RESULTS

Optimization of *S. aureus* immunocapture. The enrichment procedure for *S. aureus* was based on the ubiquitous and specific presence of protein A on *S. aureus* bacterial cells (either MRSA or MSSA). To optimize the conditions for *S. aureus* recovery, we incubated bacterial suspensions with various anti-spa antibody dilutions, ranging from 1:166 to 1:5,400. An optimal recovery of 85% from a suspension of 10³ Cowan I CFU/ml was obtained with a titer of 1:666, equivalent to 1.5 μ l of anti-spa antibody per milliliter of CS broth (Fig. 1A). Higher antibody concentrations led to lower bacterial recovery, presumably due to saturation of streptavidin-coated beads. Using this optimal antibody titer, we assessed the recovery rates from *S. aureus* suspensions over >7 orders of magnitude. Figure 1B shows that the highest recovery rates (>60%) were consistently obtained from bacterial suspensions of 0.8 to 6 log₁₀ CFU/ml. This good recovery of diluted bacterial suspensions is

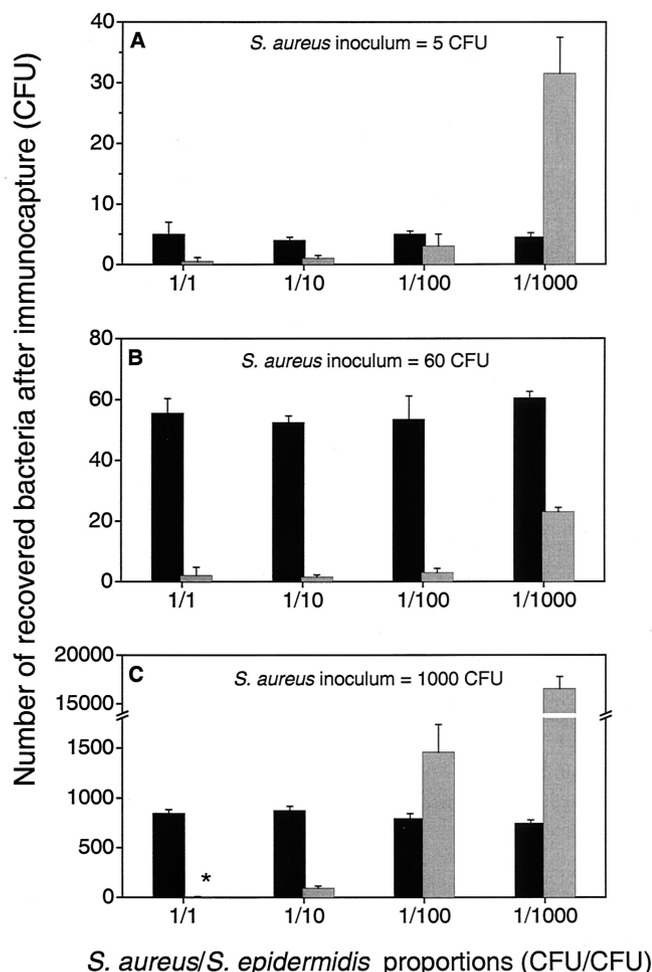


FIG. 2. Immunocapture of MRSA from mixed cultures. 5 (A), 60 (B), or 1,000 (C) CFU of MRSA (ATCC 33591) was mixed with increasing titers of MRSE, yielding ratios ranging from 1:1 to 1:1,000 for *S. aureus* (black bars) versus *S. epidermidis* (grey bars). Immunocapture was evaluated by viable CFU counts performed on MH agar. Means \pm ranges for two experiments performed in duplicate are shown. *, <10 CFU of MRSE.

a prerequisite for sensitive molecular detection and molecular identification of MRSA.

Optimized recovery conditions in mixed cultures of *S. aureus* and *S. epidermidis*. To validate the immunocapture procedure, three different titers of MRSA (5, 60, and 1,000 CFU/ml) were incubated with increasing amounts of MRSE (Fig. 2A, B, and C). The recovery of MRSA was excellent and not significantly affected by adding MRSE in ratios ranging from 1:1 to 1:1,000. Constant recovery rates (>80%) were recorded for MRSA concentrations ranging from 5 to 1,000 CFU/ml, even in the presence of a 1,000-fold excess amount of MRSE. Equivalent data were recorded with two Pastorex-negative MRSA isolates (not shown). This suggests that the minimal amount of protein A expressed by Pastorex-negative strains can be reliably immunocaptured.

Multiplex PCR assay. The limits of detection as well as the linearity of the qPCR assay were calibrated using increasing amounts (2 fg to 10 ng) of purified genomic DNA from MRSA

(ATCC 33591). The C_t values for *mecA* (Fig. 3) and *S. aureus femA* (not shown) were linear across 6 orders of magnitude of DNA input. The upper limit of linearity was >5 ng of template DNA, equivalent to 10^6 genome copies. The lower limit of MRSA DNA detection was reached with 5 fg of genomic DNA, which is nearly equivalent to the genome of 1 to 2 *S. aureus* bacterial cells (3.5 fg/cell). As expected, the qPCR signals were no longer consistently detected when using less than 10 fg of genomic DNA. On the other hand, the *mecA* and *S. aureus femA* signals detected from an input genomic DNA amount of 100 pg (100 pg of genomic DNA corresponds to >4 \log_{10} CFU) yielded similar C_t values without generating a nonspecific *S. epidermidis femA* signal. The specificity of the *femA* signals was also verified with other reference strains (Table 2).

Impact of the immunocapture procedure on the qPCR assay. When 10^3 CFU of MRSA was incubated with MRSE at ratios ranging from 1:1 to 1:1,000, detection of MRSA after immunocapture remained optimal even in the presence of a 100-fold excess amount of MRSE (Fig. 4). Equivalent MRSA detection efficacy was achieved using as few as 5 CFU of MRSA. Average C_t values for *S. aureus femA* and *mecA* were then constantly observed around 39.5 ± 0.4 . On the opposite, the immunocapture of a 100-fold excess amount of MRSE led to *S. epidermidis femA* signals >42. Such levels of residual MRSE contamination do not affect appropriate MRSA identification. As suggested by Fig. 2B, the qPCR assay identified correctly the presence of 60 CFU of MRSA even when it was incubated with a 1,000-fold excess amount of MRSE.

When the immunocapture procedure was omitted, the detection of MRSA by the *S. aureus femA* (Fig. 4) or *mecA* signals (not shown) was drastically affected by the presence of increasing amounts of MRSE. In the presence of a 1,000-fold excess amount of MRSE, nonimmunocaptured MRSA in suspension failed to be detected by the qPCR assay. Contamination of MRSA suspensions by other bacterial species, such as *E. coli* or *Pseudomonas aeruginosa*, in excess amounts yielded similar results (data not shown).

Evaluation of the qPCR assay with consecutive clinical swab samples. To test the specificity of the qPCR assay, 100 pg of genomic DNA from reference strains was analyzed (Table 2). Manual adjustment of detection thresholds led to an improved matching of species-specific *femA* signals with their corresponding *mecA* signal. The *mecA* signal was specifically detected in MRSA, MRSE, and a methicillin-resistant *S. haemolyticus* clinical isolate. Furthermore, *femA* signals were strictly species specific.

Table 3 shows the comparative results of immuno-qPCR and an optimal bacteriological procedure applied to 48 clinical swab samples. All the culture-positive samples were also detected by the immuno-qPCR assay ($n = 23$). Among the samples found to be MRSA negative by microbiological criteria ($n = 25$), 16 were also scored as negative by immuno-qPCR. In contrast, a subgroup of nine culture-negative samples was scored as positive by immuno-qPCR. Altogether, these data yielded sensitivity and negative predictive values of 100%. In contrast, the specificity and positive predictive values were 64 and 72%, respectively. A possible reason for the high proportion of false-positive cases might be the recent use of topical antimicrobial agents in those previously identified MRSA car-

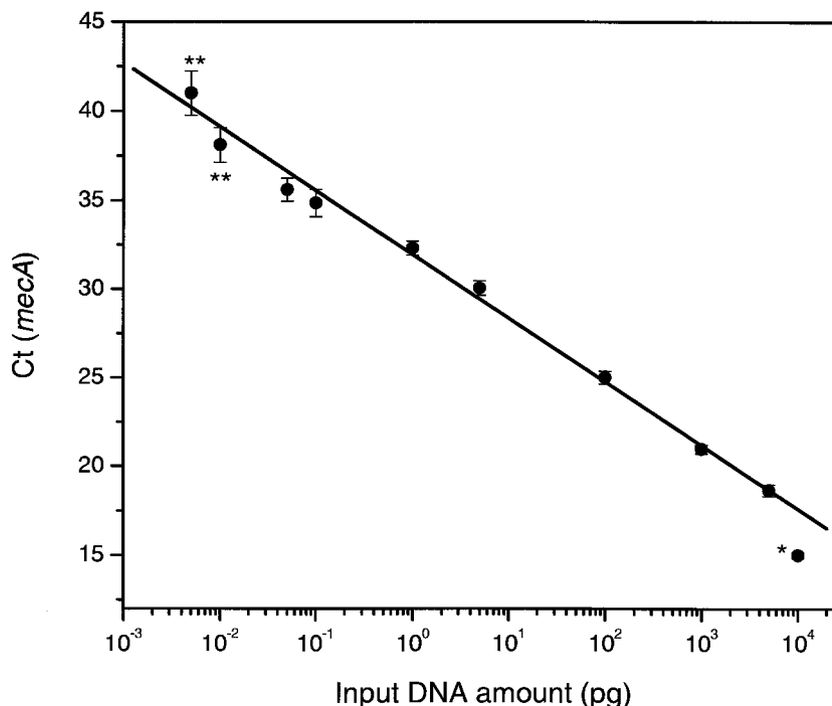


FIG. 3. Linearity and limits of detection of the qPCR assay. The linear response of the qPCR *mecA* assay as a function of input genomic DNA (2 fg to 10 ng) purified from strain ATCC33591 is shown. Correlation coefficient, >0.99; slope, -3.59. Values are means \pm standard errors of the means of 4 to 16 determinations. *, value excluded from linear regression. **, amounts of input DNA leading to irregular signal detection.

riers. This suggests that the immuno-qPCR procedure detected the presence of nonviable MRSA and yielded misleading results in this subgroup. Furthermore, all these recently decontaminated MRSA carriers relapsed shortly (<2 weeks) after the comparative analysis.

Altogether, the immuno-qPCR assay yields performance equivalent to that of optimal culture methods but has a dramatic impact on the delay for MRSA identification and implementation of infection control measures.

DISCUSSION

The identification of MRSA in biological samples is a time-consuming process relying on phenotypic or molecular analysis of isolated bacteria. We describe here a novel molecular method that does not require any bacterial culture, the time-limiting step. The major advantage of this approach, compared

to other published techniques, is same-day MRSA identification directly from swab samples or other potentially polymicrobial samples. This two-step molecular assay involves the immunocapture of *S. aureus*, followed by MRSA identification using a multiplex qPCR. This assay is based on the highly conserved *mecA* sequence within all methicillin-resistant strains and species of staphylococci, thus warranting the detection of any organism carrying this resistance determinant (52). To strictly discriminate MRSA from any other methicillin-resistant staphylococcal species, the qPCR assay unambiguously detects a species-specific *femA* region. The *femA* target was selected because of the following: (i) its presence in all *S. aureus* and *S. epidermidis* strains, (ii) the identification of species-specific oligonucleotides, and (iii) the successful development of a triplex qPCR assay. Other conserved sequences, such as *nuc* for *S. aureus* and *femB* for *S. epidermidis*, might represent alternate targets. Experiments using samples spiked with known amounts of *S. aureus* and *S. epidermidis* not only accurately discriminated either species but also maintained the lower limit of detection to one or two genome copies.

A critical innovative step of this method is the enrichment procedure that specifically immunocaptures virtually all strains of *S. aureus*. The target of our antibody, the protein A, is routinely used to confirm *S. aureus* identification because of its high prevalence and specificity. Control experiments verified the absence of significant immunocapture by a wide range of gram-positive and gram-negative species (data not shown). Recovery of MRSA was marginally influenced by the presence of *S. epidermidis* even at a 1,000-fold excess amount. This assumption was verified even for very low titers of *S. aureus*

TABLE 2. qPCR results for reference strains

Strain	C_t value for ^a :		
	<i>femA-SA</i>	<i>femA-SE</i>	<i>mecA</i>
MRSA (ATCC33591)	27.5	NA ^b	27.5
MSSA (ATCC25923)	28.5	NA	NA
MRSE ^c	NA	28.3	28.3
MSSE (ATCC12228)	NA	27.7	NA
<i>S. haemolyticus</i> ^c	NA	NA	25.1

^a Average C_t value for triplicate determinations with 100 pg of genomic DNA. *femA-SA*, *S. aureus femA*; *femA-SE*, *S. epidermidis femA*.

^b NA, not assessed; no signals detected within 50 cycles.

^c Clinical methicillin-resistant isolate.

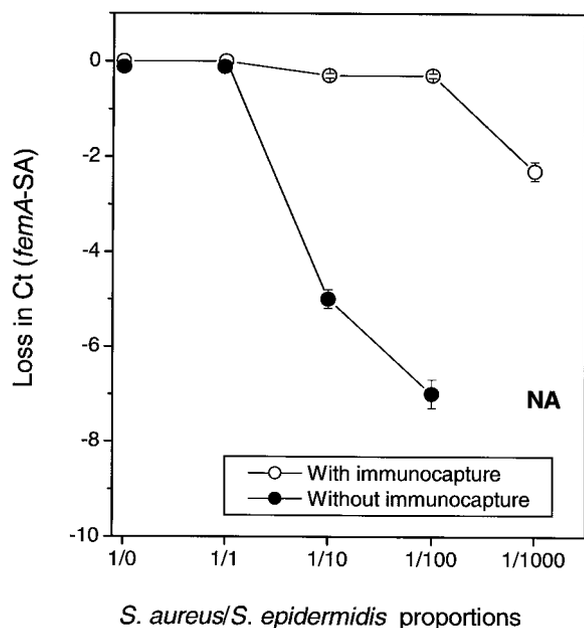


FIG. 4. Effect of the immunocapture procedure on the qPCR assay. *S. aureus* (ATCC 33591; 10^3 CFU) was mixed with various *S. epidermidis* titers (MRSE) and analyzed by qPCR following (○) or not following (●) immunocapture enrichment. MRSA in the presence of >100-fold-excess amounts of MRSE was not detected (NA). Values are means \pm standard errors of the means for three experiments performed in triplicate.

(<10 CFU) or for two exceptional Pastorex-negative MRSA clinical isolates.

Several groups have already reported on the use of multiplexed PCR assays on pure cultures (25, 47, 48). Two recently described procedures (27, 29) detected the presence of MRSA from positive blood culture bottles, with a limit of detection around 10^5 CFU/ml (29). The implementation of an immunocapture step allowed the use of qPCR directly on mixed flora samples.

The clinical performance of this immuno-qPCR assay was evaluated by testing 48 consecutive clinical swab samples. All culture-positive MRSA samples were successfully detected by immuno-qPCR. Among 25 culture-negative samples, a surprisingly high number of false-positive results were recorded ($n = 9$). A retrospective analysis revealed that these nine cases were previously known as MRSA carriers and relapsed, as evidenced by culture. However, in this high-risk population sample, 50% of the 16 cases identified as MRSA negative by both

techniques became MRSA positive during their hospital stay. Thus, this low specificity does not compromise the utility of the assay, since it mostly reflects situations of transient decolonization (14, 15, 20, 21).

Larger numbers of samples are warranted to assess the sensitivity and specificity of the immuno-qPCR assay, especially in low-risk patients. A large-scale prospective comparative study was recently initiated in our institution to evaluate the impact of single-day MRSA identification on patient management and infection control decisions.

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TABLE 3. Comparison between culture-based and immuno-qPCR methods in the detection of MRSA in clinical swab samples^a

Method	No. true positive	No. false negative	No. false positive	No. true negative	Sensitivity (%)	Specificity (%)
Culture ^b	23	0	0	25	100	100
qPCR	23	0	9	16	100	64

^a Forty-eight clinical swab samples were collected from different body sites: nares (16), inguinal (17), pooled swabs (4), or wound (11). Sensitivity (23 of 23), 100%; specificity (16 of 25), 64%.

^b Method used as the reference technique.

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