

Evaluation of a Simple Protein Extraction Method for Species Identification of Clinically Relevant Staphylococci by Matrix-Assisted Laser Desorption Ionization–Time of Flight Mass Spectrometry

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In clinical microbiology, bacterial identification is labor-intensive and time-consuming. A solution for this problem is the use of matrix-assisted laser desorption ionization—time of flight mass spectrometry (MALDI-TOF MS). In this study, we evaluated a modified protein extraction method of identification performed on target plates (on-plate extraction method) with MALDI-TOF (Bruker Microflex LT with Biotyper version 3.0) and compared it to 2 previously described methods: the direct colony method and a standard protein extraction method (standard extraction method). We evaluated the species of 273 clinical strains and 14 reference strains of staphylococci. All isolates were characterized using the superoxide dismutase A sequence as a reference. For the species identification, the on-plate, standard extraction, and direct colony methods identified 257 isolates (89.5%), 232 isolates (80.8%), and 173 isolates (60.2%), respectively, with statistically significant differences among the three methods (P < 0.05). In conclusion, the on-plate extraction method is at least as good as standard extraction in identification rate and has the advantage of a shorter processing time.

atrix-assisted laser desorption ionization-time of flight mass spectrometry (MALDI-TOF MS) has been used to identify various microorganisms in clinical laboratories, including Gram-positive bacteria, Gram-negative bacteria, yeasts, and even filamentous fungi (1, 8, 10, 12, 14, 16-19, 21, 22, 24). It is a rapid and inexpensive alternative to molecular identification that offers equivalent accuracy (6). Additionally, complete bacterial identification by MALDI-TOF MS is more cost-effective than by conventional methods (9, 15, 22). Two preparatory methods have been reported for MALDI-TOF MS-based identification (1, 4, 5): the direct colony method and the standard protein extraction method (standard extraction method). The direct colony method is performed by picking up colonies from a culture plate and placing them, as a thin layer of sample, onto the target plate and then subjecting the matrix to MALDI-TOF MS analysis (1). This method is simple and can be performed rapidly, but it is inferior in accuracy to the standard extraction method. This deficiency is particularly evident for Gram-positive organisms because of insufficient cell wall disruption (1). The standard extraction method, in contrast, employs formic acid and acetonitrile to disrupt the cells before they are placed onto the target plate for MALDI-TOF MS analysis. The standard extraction method involves 2 centrifugation steps and requires approximately 6-foldmore processing time than does the direct colony method.

We evaluated an on-plate extraction method that could be performed on the target plates without any centrifugation steps. Our study compared the on-plate extraction method with both the direct colony and the standard extraction methods for MALDI-TOF MS-based identification of staphylococcal clinical isolates.

MATERIALS AND METHODS

Bacterial isolates. This study used 273 staphylococci of nonduplicate clinical isolates and 14 type and reference strains (*Staphylococcus aureus*)

NCTC 8325, S. epidermidis ATCC 14490^T, Staphylococcus capitis subsp. capitis CCUG 7326^T, Staphylococcus capitis subsp. ureolyticus ATCC 49326, Staphylococcus haemolyticus ATCC 29970^T, Staphylococcus lugdunensis ATCC 43809^T, Staphylococcus saprophyticus JCM 2427^T, Staphylococcus cohnii ATCC 29974^T, Staphylococcus hominis ATCC 27844^T, Staphylococcus pettenkoferi CCUG 51279^T, Staphylococcus schleiferi subsp. schleiferi ATCC 43808^T, Staphylococcus schleiferi subsp. coagulans JCM 7470, Staphylococcus warneri ATCC 27836^T, and Staphylococcus caprae ATCC 33538^T). All strains were stored at -80° C until use, precultured for 12 to 24 h, and cultured aerobically overnight on 5% sheep blood agar at 37°C.

Species identification of staphylococci. Isolated species were identified by partial sequencing of superoxide dismutase A (*sodA*) by using primers d1 and d2, as previously described (20, 23). All isolates were considered correctly identified when the *sodA* sequence yielded \geq 98% sequence similarity with the closest species sequence in the GenBank database. The *sodA* sequence profiles and the origin of the clinical isolates are presented in the supplemental material.

MALDI-TOF MS. For the direct colony method, bacteria were applied, using sterile toothpicks, as thin films on 96-spot, polished, stainless steel target plates (Bruker Daltonik GmbH, Leipzig, Germany). The bacteria were then left to dry at room temperature for 1 min. Subsequently, 1.5 μ l of the matrix solution, comprising a saturated α -cyano-4-hydrocinnamic acid (Bruker Daltonik) in 50% acetonitrile (Wako Pure Chemical Industries, Osaka, Japan) and 2.5% trifluoroacetic acid (Wako), was

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Organism	No. of isolates	No. of isolates (% correct) identified by ^{<i>a</i>} :						
		Standard extraction		On-plate extraction		Direct colony		No. of spectra
		Genus level	Species level	Genus level	Species level	Genus level	Species level	in database
Staphylococcus aureus	64	64 (100)	63 (98.4)	64 (100)	63 (98.4)	64 (100)	56 (87.5)	12
Staphylococcus epidermidis	93	93 (100)	87 (93.5)	93 (100)	89 (95.7)	93 (100)	64 (68.8)	9
Staphylococcus capitis	20	20 (100)	20 (100)	20 (100)	18 (90)	20 (100)	16 (80)	6
Staphylococcus caprae	24	22 (91.6)	1 (4.2)	24 (100)	18 (75)	21 (87.5)	6 (25)	2
Staphylococcus haemolyticus	19	19 (100)	18 (94.7)	18 (94.7)	17 (89.4)	16 (88.9)	7 (36.8)	8
Staphylococcus lugdunensis	21	21 (100)	21 (100)	20 (95.2)	19 (90.4)	21 (100)	17 (80.9)	6
Staphylococcus saprophyticus	19	17 (89.4)	1 (5.2)	19 (100)	15 (78.9)	5 (26.3)	0(0)	8
Staphylococcus cohnii	3	3 (100)	0 (0)	1 (33.3)	0 (0)	0(0)	0 (0)	5
Staphylococcus hominis	7	7 (100)	7 (100)	7 (100)	7 (100)	7 (100)	5 (71.4)	6
Staphylococcus pettenkoferi	5	5 (100)	2 (40)	5 (100)	5 (100)	5 (100)	1 (20)	5
Staphylococcus schleiferi	6	6 (100)	6 (100)	6 (100)	4 (66.7)	2 (33.3)	1 (16.6)	6
Staphylococcus warneri	6	6 (100)	6 (100)	6 (100)	2 (33.3)	2 (33.3)	0 (0)	4
Total	287	283 (98.6)	232 (80.8)*	283 (98.6)	257 (89.5)**	256 (89.2)	173 (60.2)	

TABLE 1 Comparison of three methods for identification of Staphylococcus to the genus and species levels

^{*a*}*, *P* < 0.05 against direct colony for species-level identification; **, *P* < 0.05 against both direct colony and standard extraction for species-level identification.

applied to the samples and cocrystallized with them at room temperature for 10 min.

For the on-plate extraction method, each strain was applied and dried on the target plate as in the direct colony method. Following this, $0.5 \,\mu$ l of 70% formic acid (Wako) was mixed with the sample on the plate by pipetting, followed by $0.5 \,\mu$ l of acetonitrile, and the resultant mixture was dried at room temperature for approximately 10 min. Finally, $1.5 \,\mu$ l of the matrix solution was applied onto the spot as in the direct colony method.

For the standard extraction method, a small sample of each colony was suspended in 300 μ l of distilled water and adjusted at McFarland 2 standard, and 900 μ l of absolute ethanol was added. The suspension was vortexed vigorously and centrifuged at 20,000 \times *g* for 2 min. The supernatant was then discarded, and the pellet was dried at 55°C for at least 30 min. Fifty microliters of 70% formic acid was then added and thoroughly mixed by pipetting. Next, 50 μ l of acetonitrile was added, and the sample was centrifuged again at 20,000 \times *g* for 2 min. Subsequently, 1 μ l of supernatant was placed onto the target plate and left to dry for approximately 10 min at room temperature. Finally, 1.5 μ l of matrix solution was applied onto the spot, as in the direct colony procedure.

The samples prepared by each method were applied to a MicroFlex LT mass spectrometer (Bruker Daltonik), and the results were analyzed by MALDI Biotyper 3.0 software (Bruker Daltonik). Each measurement was performed only once for each culture. *Escherichia coli* DH5 α was used as a quality control as recommended by the manufacturer on each experiment.

Data analysis. The manufacturer's recommended log score identification criteria were used as follows: a score of 2.000 to 3.000 indicated species-level identification, a score of 1.700 to 1.999 indicated genus-level identification, and a score of <1.700 indicated an unreliable identification. Duplicate experiments were performed. For genus- and species-level identification, a less stringent identification criterion was used for analysis. If 1 strain attained genus-level identification in the first experiment and species-level identification in the second experiment, genus-level identification was used for analysis. If 2 experiments resulted in different results, such as genus-level identification and unreliable identification, the unreliable-identification result was used. Additionally, the rates for different criteria (genus-level, species-level, or unreliable identification) were calculated by the number of different criteria in first and second experiments divided by the total number of isolates. Different results between MALDI-TOF MS and molecular identification were categorized as erroneous identifications. Additionally, when the Biotyper ascertained only genus-level identification (log score, 1.70 to 2.00), probable species identification was estimated in accordance with the log score order. Different species results between the highest log score candidate and molecular identification were categorized as discordant results. In the case of such results, the protein signature profile was analyzed. A dendrogram was constructed using the correlation distance measure and the averagelinkage algorithm settings of the Biotyper 3.0 software. All processes related to MALDI-TOF MS identification were performed by trained personnel only.

Statistical analysis. Comparisons of genus- and species-level identifications among the 3 preparatory methods were performed using chisquare tests. A *P* value of < 0.05 was considered statistically significant.

RESULTS

Identification rates for the three different methods. The identification rates obtained using the 3 different methods are shown in Table 1. The on-plate method identified 283 (98.6%) isolates at the genus level and 257 (89.5%) isolates at the species level. By comparison, the standard extraction method identified 283 (98.6%) isolates at the genus level and 232 (80.8%) isolates at the species level, while the direct colony method identified 256 (89.2%) isolates at the genus level and 173 (60.2%) isolates at the species level. There were no statistically significant differences in genus-level identifications between the on-plate method and the standard extraction method (P = 0.1545), but a significantly higher identification rate was achieved by the on-plate method at the species level than by the standard extraction method (P <0.0001). On the other hand, the direct colony method yielded lower identification rates at the species level than did both the standard extraction (P < 0.0001) and on-plate extraction methods (P = 0.0450). The rates for different identification criteria in the first and second experiments were 30 (10.4%), 23 (8%), and 79 (27.5%) for standard extraction, on-plate extraction, and the direct method, respectively (data not shown). In species identification, there was a large difference between on-plate extraction and standard extraction for S. caprae (75% and 4.2%, respectively) and S. saprophyticus (78.9% and 5.2%, respectively). Other strains have smaller identification rate differences or only a small number of isolates. The species identification rates for coagulase-negative staphylococci (CoNS) were 169 (75.8%), 194 (87.0%), and 117 (52.5%) for standard extraction, on-plate extraction, and the direct colony method, respectively.

TABLE 2 Strains with discordant identification by each method

Organism		Expt	ID and score of MALDI-TOF MS ^a						
			Standard extraction		On-plate extraction		Direct colony		
	Strain		ID	Score	ID	Score	ID	Score	
Staphylococcus caprae	54709	1	Staphylococcus caprae	2.051	Staphylococcus caprae	2.188	Staphylococcus caprae	1.797*	
		2	Staphylococcus pasteuri	1.909**	Staphylococcus caprae	2.015	Staphylococcus caprae	1.802*	
	PG1037	1	Staphylococcus caprae	2.105	Staphylococcus caprae	2.162	Staphylococcus caprae	1.876*	
		2	Staphylococcus epidermidis	1.741**	Staphylococcus caprae	2.137	Staphylococcus caprae	1.751*	
	PG2043	1	Staphylococcus caprae	2.014	Staphylococcus caprae	2.16	Staphylococcus caprae	1.888^{*}	
		2	Staphylococcus pasteuri	1.882**	Staphylococcus caprae	2.123	Staphylococcus caprae	2.094	
	PG2075	1	Staphylococcus caprae	1.749*	Staphylococcus caprae	2.22	Staphylococcus caprae	1.909*	
		2	Staphylococcus pasteuri	1.703**	Staphylococcus caprae	2.076	Staphylococcus caprae	1.999*	
Staphylococcus warneri	54826	1	Staphylococcus warneri	2.04	Staphylococcus pasteuri	1.841**	Staphylococcus warneri	1.812*	
		2	Staphylococcus warneri	2.084	Staphylococcus warneri	1.889	Unreliable identification	1.64	

^a*, genus-level identification with correct species identification (ID); **, genus-level ID with discordant species ID.

Time required for each method. The mean times required for 48 duplicate samples (24 strains) in each procedure were 30 min for the direct colony method, 60 min for on-plate extraction, and 180 min for the standard extraction method.

Erroneous or discordant identification. No erroneous identifications were made, but 5 discordant results were obtained (Table 2).

Dendrogram. A dendrogram was created with the strains that showed discordant results and related reference strains that were installed in Biotyper 3.0 (Fig. 1). The protein signature profiles of *S. caprae* strains created by the standard extraction method comprised a cluster with *S. epidermidis* CCM4505 and *S. pasteuri* DSM 10656^T. In contrast, the profiles obtained by the direct colony and on-plate extraction methods were closely related to the profile of *S. caprae* DSM 20608^T (Fig. 1). The profiles of *S. warneri* 54826 as determined by the 3 methods were clustered into the same group as the profile of *S. warneri* DSM 20316^T.

DISCUSSION

Since MALDI-TOF MS was first applied for the identification of microorganisms as early as 1975 (2), this system has been developed for clinical microbiology laboratories as a high-throughput apparatus (3, 4, 24). The direct identification of positive blood cultures by MALDI-TOF MS was recently reported with a species identification rate of 91% (13). Improvements in the Biotyper 3.0 software and its database may change the clinical microbiology laboratory workflow (5, 6).

For MALDI-TOF MS-based bacterial identification, 2 preparatory methods have been introduced by the manufacturer. One of these is the direct colony method, which is suited for routine workflows in the modern clinical microbiology laboratory, requiring a shorter period (less than 30 min for 48 samples) and an easier procedure. However, the method presents the problem of a low identification rate, especially for Gram-positive bacteria (1, 5). Bizzini et al. reported species identification rates by the direct colony method for *Streptococcus agalactiae* as 58%, *Streptococcus pneumoniae* as 73%, *S. aureus* as 79%, and *S. epidermidis* as 58%, using MALDI Biotyper 2.0 software (4). In this study, we used MALDI Biotyper 3.0 and achieved only slightly higher rates, i.e., 87.5% and 68.8%, respectively, for *S. aureus* and *S. epidermidis* by the direct colony method. Our higher identification rates may be due to the improvement of the Biotyper version.

Another method, the standard extraction method, has been used in the construction of a database for Biotyper and has been recommended by the manufacturer as a reference method for identifications (1, 4, 7). In this study, the identification rate of the standard extraction method was 80.8% at the species level, a better result than previously reported (1). The variability of identification rates among different studies may be explained by differences in growth conditions, sample preparation, number of reference strains, version of Biotyper software, and study design (3). In our protocol, the standard extraction method consisted of approximately 13 steps, including completion of 2 rounds of centrifugation for 2 min each, requiring approximately 180 min for 48 samples. Because this is difficult to automate, the routine use of the standard extraction method may not be suitable for primary preparation (1, 4). Standard extraction may be more suitable as a reference method for use when the on-plate method or direct colony method fails to identify the species.

The on-plate extraction method offers the advantage of a simple and easy-to-use procedure. It requires only 4 steps and 60 min for a complete identification of 48 samples. Bacterial identification rates achieved with the on-plate extraction method were at least equal to those achieved using the standard extraction method. We found that *S. aureus* had a higher identification rate than CoNS. Further study is needed to identify the cause of this difference.

The identification rate of *S. caprae* and *S. saprophyticus* by the on-plate method was shown to be higher than that by the standard extraction method. The other species showed smaller identification differences between the on-plate method and standard extraction methods. This indicates that for some species of staphylococci, the on-plate method may have a better identification rate than the standard extraction method. Our protocol of complete drying up of samples for standard extraction at 55°C may provide an explanation. This method may cause the degradation of proteins, shifting the protein profile used for identification. If we had chosen a different protocol, such as a dry-up time of 10 min at room temperature, the total time required for protein extraction would be shortened, potentially producing a better identification rate for standard extraction.

Furthermore, the on-plate method may be improved as Haigh et al. reported (16). They introduced a method that did not use

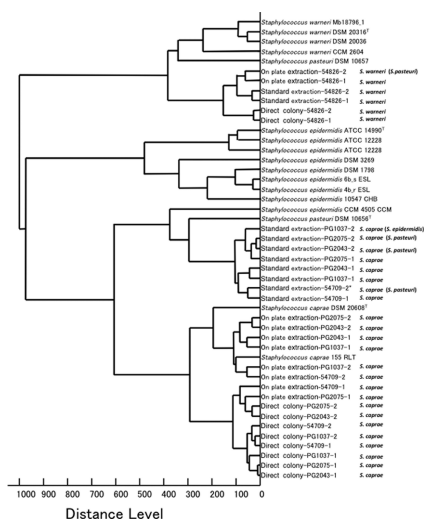


FIG 1 Dendrogram derived from the MALDI-TOF MS-specific protein signatures for strains with discordant identification results, including the profiles of *S. warneri*, *S. pasteuri*, *S. epidermidis*, and *S. caprae* from the MALDI Biotyper 3.0 database. Species names in this study defined by *sodA* sequence are shown after sample names. Additional species in parentheses are the discordant species identified by MALDI-TOF-based identification. Distance values were relative and normalized to a maximal value of 1,000.

acetonitrile, demonstrating a 10.9% improvement in the genuslevel identification rates of various clinical strains (16). A shorter method will be a great help in the installation of MALDI-TOFbased bacterial identification.

There were 5 discordant results in the data set. *S. caprae* was identified as *S. pasteuri* and *S. epidermidis* in experiment 2 using the standard extraction method (Table 2). *S. warneri* was identified as *S. pasteuri* in experiment 1 using the on-plate extraction method. According to results from the MALDI Biotyper 3.0 database, the protein profile dendrograms for standard extractions of 4 strains of *S. caprae* showed clustering with *S. epidermidis* CCM4045 and *S. pasteuri* DSM10656^T and separation from the cluster with *S. caprae* DSM 20608^T (Fig. 1). The protein profile of *S. warneri* DSM 20316^T was closely related to that of *S. pasteuri* DSM10656^T. This discrepancy of phylogeny between the MALDI-TOF MS profile and the *sodA* sequence may be correlated with the discordant results obtained in our experiment. Similarly, the close genetic and protein relatedness

between *S. pasteuri* and *S. warneri* might have caused the discordant results with *S. warneri* 54826.

The MALDI Biotyper 3.0 included only 2 strains of *S. caprae* as references, compared with 4 strains of *S. warneri* and 9 of *S. epi-dermidis* (Table 1). This small number of reference strains might have limited the accuracy in identifying *S. caprae*. Indeed, Seng et al. previously described the correlation between the accuracy of identification by MALDI-TOF MS and the number of reference strains (22). Similar observations have been reported by several other researchers (1, 18). Further, Lista et al. showed that additional reference strains should more accurately represent the genetic diversity of the strain (19). Therefore, proper representatives from various genetic backgrounds for each species should improve the species-level identification rates. Future studies using an improved and extended database should overcome this issue.

In conclusion, we demonstrated that the on-plate extraction method offers species identification rates at least equivalent to the results of the standard extraction method, with advantages in convenience and faster processing time, and a significantly better species identification rate than the direct colony method in *Staphylococcus* analysis.

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