Is wet swab superior to dry swab as an intranasal screening test?

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Abstract

Methicillin-resistant Staphylococcus aureus is still a great concern, and recognition of the carrier is essential for appropriate infection control in intensive care units. The utility of wet swab compared to dry swab as an intranasal screening test has not been well assessed yet. A comparative study of the wet and dry swab in its ability to detect the organism was performed against critically ill patients, and it was found that there were no statistically significant differences between the two different methods. The wet swab did not show increased sensitivity compared to dry one.

Keywords: Intensive care unit, Methicillin-resistant Staphylococcus aureus, Screening test, Swab

Correspondence/findings

Nosocomial infection by methicillin-resistant Staphylococcus aureus (MRSA) is a great concern in the intensive care unit (ICU) where critically ill patients are gathered. However, controlling MRSA infection is still a hard matter [1]. Since bacterial culture of nasal sample is inexpensive, easy, and available, it is a standard method for the screening test for MRSA carrier. An earlier study comparing the sensitivities of dry and wet swab sampling has shown that these methods led to the same results regarding the detection of intranasal MRSA [2]. However, it is possible that the effectiveness of swab-screening tests may be differed depending on the sampling technique, condition, and the circumstances of each ICU. We therefore conducted a prospective study in purpose of reassessing the validity of wet swab as an intranasal screening test for MRSA carrier in our clinical setting.

The study was performed at an ICU of Tsuyama Central Hospital (Okayama, Japan) from March to May in 2012. Only those patients who were provided with informed consent were eligible for inclusion. For each patient, ICU nurse obtained two anterior nares samples (one dry and one wet), using rayon swab (CultureSwab Plus: Becton, Dickinson and Company, BBL). A naris for sampling was chosen in a random manner for wet swab and the other for dry. Wet swabs were manually moistened with sterile saline just before sampling. After sampling, the specimen was immediately transferred to the own microbiology division. The samples were plated on mannitol salt agar plate after washing the swabs with 1 mL of sterile saline. After incubating 24 h at room temperature, the number of colonies was counted, and the identification of the organism was performed using Microscan WalkAway® (Siemens, Tokyo, Japan). The comparison between the number of colonies of MRSA, methicillin-sensitive S. aureus (MSSA), and all bacteria grown on the plate was performed. Statistical analysis was performed using Kaleida Graph 4.1 (Synergy Software, Reading, PA, USA), and Wilcoxon signed rank test was applied. The present study protocol (No. 122) was approved by ethics committee of Tsuyama Central Hospital.

The total number of subjects was 141. MRSA was isolated from eight samples with dry swab and nine samples with wet swab, and MSSA was isolated from 18 samples with dry swab and 17 samples with wet swab, respectively. Comparison of dry and wet swab was performed in those MRSA positive (A), MSSA positive (B), and all bacteria (C) (Figure 1). There were no statistically significant differences between dry swab
and wet swab in each group ((A) $P = 0.23$, (B) $P = 0.26$, and (C) $P = 0.11$).

MRSA can be easily transferred by healthcare workers [3] and often causes life-threatening infections in ICU [4]. Isolating or cohort of those carrier patients is generally recommended [5], and a recent study reported that the rate of MRSA infections was reduced by 62% in ICU and 45% in the other ward with an introduction of MRSA prevention bundle [6]. On the other hand, Huskins et al. reported that surveillance for MRSA colonization combined with subsequently expanded barrier precautions was not effective in reducing the transmission of MRSA [7]. The recognition of MRSA carrier in ICU is, however, considered essential for the infection control, and for that, a high-quality screening method is indispensable [8,9].

The efficacy of progressive screening test on nasal swab for MRSA with using polymerase chain reaction (PCR) has been reported [10]. PCR screening outperforms the classical bacterial culture in its high sensitivity and specificity, but it cannot be introduced to majority of medical institutions because of its high cost and equipment investments.

Recent studies showed that universal decolonization using chlorhexidine was more effective than ‘screening and isolation’ strategy in reducing MRSA infection rate [11,12]. However, such a methodology is still not appreciated as a general way to control nosocomial MRSA infection, and we consider that establishment of a reliable screening method is fundamental at present.

Provided that the wet swab yielded more sensitivity in detecting the pathogen, it would be appreciated since it is easy and available method, and does not cost. However, according to our result, the wet swab did not show increased sensitivity compared to the dry one. This result was same as the previous study [2]. The sampling protocol in our study, inserting the wet and dry swabs into each nostril separately, could have influenced the result. According to Kildow et al., healthy adults are more likely to carry $S. aureus$ in one nostril than in both [13]. Or, since the distinctive sampling method was not defined precisely in our protocol, therefore the depth or degree of swab insertion into nasal cavities could be different in each subject, which could lead to the sampling error. Small sample size could also be responsible to the result. In any cases, our result indicates that the validity of preparing wet swab rather than dry one is not warranted as an intranasal screening test for MRSA carrier in ICU.

**Abbreviations**

ICU: Intensive care unit; MRSA: Methicillin-resistant *Staphylococcus aureus*; MSSA: Methicillin-sensitive *S. aureus*; PCR: Polymerase chain reaction.

**Competing interests**

The authors declare that they have no competing interests.
Authors’ contributions
HH, TM, NM, and SS conceived of the study and participated in its design. HH drafted the manuscript. KE and YK helped for coordination for sampling. TU carried out the colony count. MM performed the statistical analysis. FO helped to draft the manuscript. All authors read and approved the final manuscript.

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