Nasal colonization and bacterial contamination of mobile phones carried by medical staff in the operating room

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Abstract

Background

Mobile phones (MPs) have been an essential part of the lives of healthcare professionals and have improved communication, collaboration, and sharing of information. Nonetheless, the widespread use of MPs in hospitals has raised concerns of nosocomial infections, especially in areas requiring the highest hygienic standards such as operating rooms (ORs). This study evaluated the incidence of bacterial contamination of the MPs carried by medical staff working in the OR and determined its association with bacterial colonization of this personnel.

Methods

This is an observational cohort study. Medical staffs working in the OR were asked to take bacterial cultures from their MPs, anterior nares, and dominant hands. To identify the relation between MP contamination and bacterial colonization of the medical staff, genotyping of Staphylococcus aureus (SA) was done via Staphylococcus protein A gene (spa) typing and pulsed-field gel electrophoresis (PFGE).

Results

A total of 216 swab samples taken from 72 medical-staff members were analyzed. The culture-positive rate was 98.1% (212/216). In 59 (27.3%) samples, the bacteria were possible clinical pathogens. The anterior nares were the most common site of colonization by clinical pathogens (58.3%, 42/72), followed by MPs (13.9%, 10/72) and the dominant hand (9.7%, 7/72). SA was the most commonly isolated clinical pathogen and was found in 43 (19.9%)
samples. In 66 (94.3%) of the 70 staff members for whom bacteria were detected on their MPs, the same bacteria were detected in nares or hand. Among 31 medical staff who were carriers of SA in the anterior nares or dominant hand, 8 (25.8%) were found to have SA on their MPs, and genotyping confirmed the same SA strain in 7 (87.5%) of them.

**Conclusion**

A high rate of bacterial nasal colonization and MPs contamination were found among the OR medical staff. An MP may be a reservoir for pathogen contamination in the OR.

**Introduction**

Hospital-acquired infection (HAI) is a serious problem in hospitals and may cause increased morbidity and mortality [1]. It is estimated that between 5% and 10% of hospitalized patients have HAI, and recent data suggest that this figure is on the rise [2]. In the United States, the reported cost of HAI in 2002 was $6.7 billion [3], and increased to 2 million cases/year, 100,000 deaths, and costs $20 billion yearly [4]. As early as 1861, Semmelweis showed that bacteria are transmitted to patients via the contaminated hand of healthcare workers [5]. The transmission of HAI occurs through hospital staff and via the animate hospital environment, equipment, and devices that the hospital staff use [6].

Mobile phones (MPs) have become an integral part of the lives of healthcare professionals and have improved communication, collaboration, and sharing of information. Widespread use of MPs in hospitals has raised concerns of nosocomial infections, especially in areas requiring the highest hygienic standards such as the operating room (OR). In other studies, prevalence of MPs contaminated with bacteria was reported to range from 62% to 99% [7–10], and potential clinical pathogens were detected up to 14.3–75% MPs [7, 8, 10–13]. Brady et al. [11] found that 14.3% of MPs are colonized by bacteria known to cause nosocomial infections. Khivsara et al. even found that there is a 40% rate of MSSA (6.7% MRSA) colonization of MPs [14]. Borer et al. [15] found MPs to be an important cause of HAI.

*Staphylococcus aureus* (SA) is a common pathogen in osteomyelitis [16] and periprosthetic joint infection [17]. Patients who are colonized with SA are at a 2- to 9-fold higher risk of SA infection [18]. The most common area for SA colonization in the human body is nasal nares [19], and high prevalence of contamination of MPs was also reported [7–11, 14]. In public opinion, hospital OR should be the workplace with the highest hygienic standards. The same high hygienic requirements also hold for the personnel working there and the equipment used by them. Nonetheless, MPs are not forbidden in the OR [20].

Nowadays almost everyone has at least one mobile device and carry them anytime and virtually everywhere. In the OR, medical staffs are asked to change surgical gowns and wear a surgical mask at all times to prevent from spreading bacteria to the environment. The use of personal MPs, however, has not been restricted in the OR mostly. Contamination of MPs revealed a potential bacteria reservation and a potential source of bacterial spreading in the OR. The colonization of medical staff’s nasal nares may be transmitted to MPs by hands. The relation among the rates of colonization of medial staff’s nares, hands, and MPs is currently unclear. Are hands randomly contaminated via MPs, or contaminated from a user’s nasal colonization? The purposes of this study were to evaluate the colonization status of orthopedic medical staff working in an OR in terms of nasal nares, hands, and MPs, and to identify the bacteria involved. The SA that was isolated was chosen for further genotyping to clarify the
bacterial transmission among nares, hands, and MPs. We hypothesized that nasal colonization by SA may be transmitted to hands and MPs.

**Materials and methods**

**Sample collection and processing and isolation of microorganisms**

This is an observational cohort study. After approval by the Institutional Review Board of Chang Gung Memorial Hospital (IRB approval No. 102-4862B), the study was initiated in January 2014 and ended in March 2014. The medical staff working in the OR was enrolled after written informed consent was obtained. In the Department of Orthopedic Surgery, there were totally 112 medical-staff who worked in the OR, including 60 OR nurses and 62 surgeons. We excluded 21 staffs who were in night shifts, 9 who were on vacation and 10 who were not willing to participate the study, leaving 72 in the analysis. The characteristics were listed in Table 1.

During daily work in the OR, samples were taken between every operation, 30 minutes after removing surgical groove at the end of operation. Samples for bacterial culture from the anterior nares, dominant hands, and MPs were collected with a cotton swab moistened with distilled water. The MPs belonged to the medical staffs and were used only by owners. Culture samples from anterior nasal nares were collected by rubbing the swab against the anterior 1 cm of the nasal vestibular wall of both nares. After swabbing around anterior nasal nares, dominant hand, or mobile phone respectively, a culture plate was streaked with the cotton swab immediately in the OR. The cotton swab of every sample (anterior nares, dominant hands, or MPs) was cultured on BP/EMB (blood agar plate/eosin methylene blue agar plate) and CNA (Columbia-colistin and nalidixic acid agar) on one fourth of the plate, and then we snapped the tip of the cotton swab into Thioglycollate broth without contamination of surrounding objects.

During sampling, the MP was swabbed all around. The dominant hand was swabbed both dorsally and ventrally and between the fingers. The nasal nares were swabbed the swab against the anterior 1 cm of the nasal vestibular wall of both nares. The culture plate and broth were transported to microbiology laboratory after the above procedure immediately. After the plate was delivered to the microbiology laboratory, the plate was streaked with a sterilized iron stick over the remaining three-fourths of the plate in a cell culture hood. Then, the culture plate and broth were incubated at 37˚C for 48 h. In every time slot (every 2 h) during sampling in the OR, a negative control was set up as follows: the tip of a cotton swab moistened with distilled water was snapped into Thioglycollate broth.

**Identification of the microorganisms**

Isolated microorganisms were identified using the Gram stain, pigmentation, colony morphology, catalase assay, motility, esculin hydrolysis, Staphaurex Plus (Remel Europa Ltd., Dartford, 

| Participate | Total |
|-------------|-------|
| Age         | 37.60 | 38.95 |
| Sex M/F     | 46/26 | 64/36 |
| Nurse       | 19    | 41    |
| Nurse partisan | 6    | 9     |
| Resident    | 22    | 24    |
| Attending doctor | 25   | 38    |

72   112

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The bacteria isolated as coagulation-negative *Staphylococcus* (CNS), gram-positive bacillus (GPB), *Streptococcus* spp. (*Strep*), *Moraxella* spp., and gram-negative *Bacillus* (GNB) were considered as normal flora [21]. The isolated bacteria were subjected to matrix-assisted laser desorption ionization time-of-flight mass spectrometry (MALDI-TOF MS) [22] for confirmation of the identification. If the culture revealed SA, further genotyping by means of surface protein A (*spa*) and pulsed-field gel electrophoresis (PFGE) were performed to identify the strain.

**Genotyping**

**DNA extraction.** Whole-cell (genomic) DNA, used as a template for PCR amplification, was prepared from single colonies using the Wizard® Genomic DNA Purification Kit (Promega, Dübendorf, Switzerland). The PCR products were purified for sequencing using the PureLink™ Quick Gel Extraction and PCR Purification Combo Kit (Invitrogen, Zug, Switzerland).

**Spa typing.** The polymorphic repeat region (region x) of the SA protein A encoded by the *spa* gene was amplified as described previously [23] using primers listed below. Using a software package (Ridom StaphType software version 1.4, Ridom GmbH, Würzburg, Germany), the *spa* types of SA strains were analyzed and the new *spa* type assignment was provided automatically through the Ridom SpaServer (http://spa.ridom.de/index/shtml). The BURP (Based Upon Repeat Patterns) can be used to assign *spa* clonal complexes (CCs).

**SA protein A primers.**

1095F
AGACGATCCTTCGGTGAGC  
1517R
GCTTTTGCAATGTCATT TACTG

**Pulsed-field gel electrophoresis (PFGE).** The DNA fingerprints generated by PFGE were analyzed by both the manual method according to the criteria proposed by Tenover et al. [24] and by a digitized method using BioNumerics Fingerprint types and Cluster Analysis software (Applied Maths, Austin, TX). Percent similarities were identified on a dendrogram built using arithmetic averages and Dice coefficients. Band position tolerance and optimization were set to 1.25 and 0.5%, respectively.

**Results**

There were 216 swab samples taken from 72 medical-staff members included in the study, with 3 samples (nares, hands, MPs) per person. The overall bacteria-positive rate was 98.1% (212/216), the highest in nasal nares (100%, 72/72), followed by dominant hands (97.2%, 70/72) and MPs (97.2%, 70/72). The isolated microorganism was a possible clinical pathogen in 27.3% (59/216) of the samples, and was most frequently found in nasal nares (58.3%, 42/72), followed by hands (13.9%, 10/72) and MPs (9.7%, 7/72; Table 2). The most common clinical pathogen was SA (19.9%, 43/216), with 27 methicillin-sensitive strains (MSSA, 12.5%, 27/216) and 16 methicillin-resistant strains (MRSA, 7.4%, 16/216), followed by *Enterobacter* spp. (5.6%, 12/216) and *Citrobacter koseri* (4.6%, 10/216; Table 3).

In 72 medical-staff members, there were 70 people having a positive culture from their MPs (97.2%, 70/72). Among that, 66 (94.3%, 66/70) were found to have the same microorganism in the nares or hands (nares only, 16.7%; hands only, 4.5%; both, 78.8%). Among the 12 participants for whom a clinical pathogen was isolated from their MPs, the same clinical pathogen was also found in 10 participants (83.3%) in their nares or dominant hands (nares, 70%; hands, 10%; both, 20%).

The distribution of SA discovered at the 3 sites is shown in Table 4. SA was found in 31 (31/72; 43%) swab samples from anterior nares, 8 samples (8/72; 11.1%) from MPs, and 4 (4/72;
5.6%) from the hands. Among these samples, methicillin-resistant strains were found in 10 (32.3%), 3 (37.5%), and 3 (75%) samples from the anterior nares, MPs, and hands, respectively. It should be noted that among 31 SA carriers, 8 (25.8%) had growth of SA in cultures from their MPs, and all 8 medical staff who had SA in cultures from their MPs were SA carriers (6 in the anterior nares and 2 in the anterior nares and hand). Genotyping using Spa and PFGE confirmed the same strains of SA in 7 of them (87.5%, 7/8).

Table 2. Isolation proportion in different culture.

| Culture     | Nasal nares | Hands | MPs |
|-------------|-------------|-------|-----|
| Culture positive | 100%(72/72) | 97.20%(70/72) | 97.20%(70/72) |
| Culture negative | 0          | 2.8%(2/72)    | 2.8%(2/72)    |

Table 3. Distribution of isolated bacteria.

| Isolation rate | Nasal Nares | Hands | MPs |
|----------------|-------------|-------|-----|
| Possible clinical pathogen | 100% (72/72) | 97.20% (70/72) | 97.20% (70/72) |

**Gram-positive**
- MSSA: 21, 1, 5
- MRSA: 10, 3, 3

**Gram-negative**
- *Enterobacter aerogenes*: 10, 1, 1
- *Escherichia coli*: 1
- *Acinetobacter baumannii*: 1, 3, 1
- *Klebsiella pneumoniae*: 1
- *Pseudomonas putida*: 2
- *Citrobacter koseri*: 10

**Normal flora**
- CNS: 70, 60, 65
- GPB: 35, 24, 14
- *Streptococcus spp.*: 5, 13, 10
- *Moraxella spp.*: 3, 2
- GNB: 2

MSSA: Methicillin-sensitive *Staphylococcus aureus*
MRSA: Methicillin-resistant *Staphylococcus aureus*
CNS: Coagulase-negative *Staphylococcus aureus*
GPB: Gram-positive *Bacillus*
GNB: Gram-negative *Bacillus*
Discussion

MPs contaminated with bacteria had been studied well [7–14], but the relation between nasal colonization and contamination of MPs is not well understood. For further evaluation of this relation, we initiated this study by doing culture analysis of nares, hand, and MP. Further SA genotyping with *spa* analysis and PFGE was also conducted to clarify the relation among nasal, hands, and MPs colonization proportion.

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Table 4. Distribution and genotyping of *Staphylococcus aureus*.

| Staff No | Drug sensitivity | Nasal Nares | Hands | MPs |
|----------|------------------|-------------|-------|-----|
|          |                  | Spa | PFGE | Spa | PFGE | Spa | PFGE |
| *1*      | S                | S1  | I    |     |      | S1  | I    |
| 6        | S                | S2  | A    |     |      |     |      |
| *8*      | R                | t4911| E    | t4911| E    | t4911| E    |
| 9        | S                | t1209| A    |     |      |     |      |
| *10*     | R                | t026 | B    | t026 | B    |     |      |
| 11       | S                | t4911| D    |     |      |     |      |
| 13       | S                | S3  | H    |     |      |     |      |
| *15*     | S                | S4  | F    | S4  | F    |     |      |
| 19       | S                | S5  | F    |     |      |     |      |
| *21*     | R/S              | t026| C    | S6  | J    |     |      |
| 24       | R                | S7  | L    |     |      |     |      |
| *29*     | R                | t1751| G    |     |      | t1751| G    |
| 32       | S                | S8  | A1   |     |      |     |      |
| 33       | S                | S9  | K    |     |      |     |      |
| 37       | S                | t213| Q    |     |      |     |      |
| 39       | R                | t4911| X   |     |      |     |      |
| 41       | S                | t2379| R   |     |      |     |      |
| 42       | S                | t3535| W   |     |      |     |      |
| *48*     | S                | S10 | M    | S10 | M    |     |      |
| 49       | S                | t338| N    |     |      |     |      |
| 51       | S                | t5693| O   |     |      |     |      |
| 52       | S                | t189 | P   |     |      |     |      |
| *53*     | R                | t441 | S   | t441| S    | t441| S    |
| 55       | S                | t5693| T   |     |      |     |      |
| 58       | S                | t111 | U    |     |      |     |      |
| *61*     | S                | S12 | V    | S12 | V    |     |      |
| 62       | R                | t4911| Y   |     |      |     |      |
| 66       | S                | t737 | AA  |     |      |     |      |
| 67       | R                | t437 | AB  |     |      |     |      |
| 68       | R                | t437 | AC  |     |      |     |      |
| *71*     | S                | t2868| AD  |     |      | t2868| AD  |

* Staphylococcus aureus isolation from both nares and either dominant hand or mobile phone
S1–S12: Repeat succession showed in S1 File
MPs: Mobile phones
*spa*: Staphylococcus protein A gene
PFGE: Pulsed-field gel electrophoresis
S: sensitive
R: resistant

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A high identification proportion, up to 98.1% (212/216), was observed in our study. Review of other studies suggests that the contamination rates of MPs vary from 62% to 90% [7–10], and detection of potential clinical pathogens also varies: from 14.3% to 75% of samples [7, 8, 10–13]. The reason for the large variation may be the sampling from different types of MPs owners (healthcare workers, anesthesiologists, or doctors at a teaching hospital), sampling methods (cotton swab was moistened or not before sampling, immediate streaking of the plate or not, final placement of the cotton swab tip into culture broth or not), and where the culture plate was streaked with the cotton swab (in room air, in a cell culture hood after transportation to a microbiology laboratory, or in the OR under laminar flow conditions). There are only a few papers with a high identification rate, up to 90% [9, 10], and those authors used moistened cotton swabs to obtain the samples. We tried to optimize the culture rate and lower the possible contamination by using moistened cotton swabs, immediate streaking on the culture plate under OR laminar flow conditions (transported the material to a microbiology laboratory to streak the rest of the culture plate), and placed the cotton swab tip into culture broth for enriched culture finally. The sampling was done in nares, dominant hands, and MPs at the same time for us to trace the possible transmission. The distribution of bacteria among nares, hands, and MPs was reasonable, with the biggest presence in nares, followed by MPs and hands (Tables 2 and 3). The nares are a moist and warm environment for bacterial colonization; this situation probably contributed to the highest isolation rate. MPs are rarely cleaned up, with a little bit higher isolation rate than from hands, which are frequently washed in the OR. It was warning that nasal colonization might be passed on to MPs by hands.

After we reviewed the results of bacterial isolation, the majority of bacteria were CNS. The CNS colonization rate was up to 97.2% (70/72) in nasal nares, followed by MPs (90.2%, 65/72) and hands (83.3%, 60/72; Table 3), comparable with data from other studies—by Ulger et al. [9] and Pal et al. [25]—who showed that the most common isolated bacterium from hands and MPs is CNS. The potential clinical pathogens identified in this study are a cause for concern. Among the medical staff, 58.3% members (42/72) carried potential clinical pathogens, and most of these bacteria were isolated from nares (Table 3). Among possible clinical pathogens, the most common species was SA. There were 43% (31/72) SA and 13.9% (10/72) MRSA cases among swab samples from nares (Table 3). According to other studies, the nasal colonization rate among hospital janitors is 15.3% [26], and the nasal colonization rate among residents in a nursing home is 18.3% [27]. The highest colonization rate was reported among patients in an Indian hospital: up to 37% [28]. In Taiwan, studies revealed that nasal MRSA colonization rates among healthcare workers (5.0–7.8%) [29] and pediatricians (6.8%) [30] are significantly higher than the rate in the general population (3.8%) [29–31]. If we compare our results with data from other studies, the SA nasal colonization among our medical staff was much higher, even higher than among patients. The possible reason is the methods of sample collection as already mentioned, and our method allowed us to trace the transmission on the basis of the high SA isolation rate. Nevertheless, not only was the nasal colonization rate of SA high, the prevalence of MRSA was high (32.3% of SA-positive samples) too. Even in comparison with a recent study involving sampling from pediatricians and polymerase chain reaction (PCR) analysis [30], our MRSA isolation rate is still higher (13.9% versus 6.8%). Because of the high colonization rate in nares with high prevalence of resistant strains, additional measures for intraoperative sterilization and avoidance of contamination can be recommended.

As the most popular nonmedical electronic equipment, MPs are used in close contact with the body and often without cleaning guidelines that meet hospital standards. The bacterium most frequently isolated from MPs in our study is CNS, followed by Streptococcus spp. and SA. There was also a gram-negative bacterium (11.1%, 8/72). The nasal colonization of medical staff may be transmitted to patients via hands, but there is no evidence to prove that yet. For all
bacteria, isolation from MPs coincided with isolation from nares or hands in 94.3% of such cases (66/70). Among 31 medical staff for whom SA was isolated from their MPs, anterior nares, or dominant hands, 8 (25.8%) were found to have SA on MPs. As the most common pathogen in human bodies causing tissue infections, pneumonia, septicemia, and device-associated infection, especially in the orthopedic field including osteomyelitis [16] and periprosthetic joint infection [17], SA was chosen for genotyping to confirm the transmission pattern. For 8 SA-positive MPs, the same SA strain was found in 7 samples from nares (87.5%); this finding was proved by \textit{spa} and PFGE typing (Table 4). Among those 7 SA-positive cultures, 3 were MRSA, and 4 were MSSA. MRSA seemed to be transmitted to MPs more easily than MSSA did (33.3% vs. 19.0%, 3/9 vs. 4/21; Table 4).

We chose 2 types of genotyping to confirm the transmission of SA: PFGE and SA \textit{A} gene (\textit{spa}). PFGE is the gold standard of SA strain typing although it is time-consuming and the interlaboratory comparability of results requires extensive effort by harmonization of protocols. To make genotyping easier (sequence data can be transferred between laboratories via the Internet), DNA sequence-based approaches are becoming more popular. There is evidence for recombination in SA, where single-locus DNA sequencing of repeat regions of the coagulase gene (\textit{coa}) and the \textit{spa} gene can be used for reliable and accurate typing of SA [32, 33]. The \textit{spa} typing is especially valuable for rapid typing of SA in hospital settings because it offers higher resolution than \textit{coa} typing. The \textit{spa} typing was not commonly used in the past because of problems with a consensus on assignment of new \textit{spa} repeats and types. Nevertheless, software Ridom StaphType was developed which now synchronizes data either directly via the http protocol or via a file-based transfer (e.g., by e-mail) with an accompanying SpaServer that functions as an operative source for all new \textit{spa} repeats and type codes [34]. We chose PFGE and \textit{spa} typing for every strain of SA to reliably identify SA transmission among nares, hands, and MPs.

We tried to evaluate the colonization status of orthopedic medical staff working in an OR by identifying the bacterial flora with further genotyping of SA to clarify the relation among nasal, hands, and MPs colonization rates. However, there are some limitations to this study. First, we performed genotyping of only SA to trace and confirm the transmission from nares to hands and MPs. The transmission of other bacteria may not be the same as that for SA. Nevertheless, SA was found to be the most common isolated bacterium among possible clinical pathogens for us to trace. Second, normal flora in this study was defined according to a definition of specific types of infections [21]. On the other hand, normal flora as defined here may be potentially infective. For example, CNS here was identified as normal flora. Nowadays, CNS is classified into several strains [35], some of which were reported to cause a disease, such as \textit{Staphylococcus epidermidis} [36] and \textit{Staphylococcus capitis} [37] reported as an opportunistic human pathogen. The majority of the bacterial isolates in this study were CNS: a cause for concern. Third, although we confirmed the transmission of nasal colonization to hand and MP by genotyping, we could not identify a direct relation between colonization of medical staff and surgical-site infections (SSIs). In our hospital, the definition of SSI is that infection was noticed by symptoms (ecchymosis, swelling, local heat, or tenderness) or pus formation proved by tapping or imaging study within 1 year after surgery. And The incidence of SSIs in our hospital in recent 4 years was around 0.3–0.8%, with 30.2% of MSSA, 12.7% of MRSA, 4.8% CoNS, and 15.9% Gram-negative bacteria. Although the pathogens of SSIs were similar to the bacteria isolated in this study, further studies on bacteria isolated from infection sites and from colonized medical staff or patients are needed to clarify this issue.

In conclusion, at a high isolation rate of bacteria, the same bacteria in MPs and nares or hands were often detected here (94.3%, 66/70). The high rate of SA nasal colonization (43.1%, 31/72) was observed, and in 22.6% of cases (7/31), this colonization may be passed on to MPs.
according to spa and PFGE genotyping confirmation. The same SA strain on MPs and nares or hands was also proved in most cases (87.5%, 7/8). As for SA contamination, MRSA was transmitted more easily from nares to hands or MPs than MSSA was (33.3% vs. 19.0%). Therefore, MPs may serve as a reservoir of potential pathogens in the OR from a user’s nasal colonization because the same strain was confirmed by spa and PFGE genotyping. As the pathogen of SSIs was similar to contamination pathogens on MPs, and transmission of bacteria was probed by genotyping, further research about the relationship between SSIs and MPs are needed. With the evidence revealed in this study, we suggested that the use of personal MPs should be regulated in the ORs. If possible, regular clean-up is suggested for decreasing MPs contamination.

**Supporting information**

S1 File. Repeat succession of *Staphylococcus aureus*. (DOCX)

**Author Contributions**

Conceptualization: PHH.

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