Prevalence of methicillin-resistant Staphylococcus aureus (MRSA) in retail food in Singapore

Kyaw Thu Aung1,5, Li Yang Hsu2, Tse Hsien Koh3, Hapuarachchie Chanditha Hapuarachchi1, Man Ling Chau1, Ramona Alikiteega Gutiérrez1* and Lee Ching Ng1,4

Abstract

We characterised 227 Staphylococcus aureus isolates from retail food and food handlers’ gloves samples obtained through food surveillance and risk assessment studies between 2011 and 2014. Of 227 isolates, five (2.2%) were methicillin-resistant and belonged to sequence types ST80 (n = 3) and ST6 (n = 2). All five isolates belonged to SCCmec type IV, were Panton-Valentine leukocidin (pvl)-negative and staphylococcal enterotoxin genes-positive. Resistance to azithromycin was found in ST80 isolates, in addition to resistance to beta-lactams. Our finding of two clinically relevant methicillin-resistant St. aureus (MRSA) strains (ST80 and ST6) in ready-to-eat food and food contact surfaces at retail in Singapore suggests food and food contact surfaces as potential environmental sources of MRSA in the community.

Keywords: Methicillin-resistant Staphylococcus aureus (MRSA), Retail food, Food contact surface, Antibiotic resistance, Enterotoxin genes, Panton-valentine leukocidin (pvl) gene

Dear Editor,

Methicillin-resistant Staphylococcus aureus (MRSA) has been recognised as an important nosocomial pathogen and has reportedly been associated with foodborne illnesses [1–3]. MRSA has also recently been listed as one of the high-priority antibiotic-resistant pathogens as ranked by the World Health Organisation. In Singapore, surveillance and control programmes of MRSA have been established in various hospitals [4, 5]. However, there is a substantial lack of information on the prevalence of MRSA in food, including at retail. Such information would be useful to better understand the risk of exposure to MRSA through food, particularly ready-to-eat food, in contrast with the more typical known transmission route via contact. Our findings provide preliminary insights on the extent of the spread of MRSA in ready-to-eat food in Singapore.

In this study, we characterised 227 coagulase-positive Staphylococcus aureus strains isolated from retail food and swabs taken from food handlers’ gloves. These isolates and samples were obtained from food surveillance and risk assessment studies conducted between 2011 and 2014. The isolates were confirmed to be S. aureus using coagulase rabbit plasma (Remel), and conventional PCR for the detection of femA gene with modified primers (F- GATCATTTATGGAAGATACGTCAG, R- GATAAAGAAGAAACCAGCAGAGATAG) and PCR conditions from previous studies [6, 7]. To understand the occurrence of methicillin resistance among these S. aureus isolates, we used mecA-PCR for the detection of MRSA, followed by PBP2 latex agglutination test (Oxoid) and disc diffusion with Cefoxitin 30 μg for confirmation. MRSA strains were further characterised by Multi-Locus Sequence Typing (MLST), staphylococcal Protein A (spa) typing, and staphylococcal cassette chromosome (SCCmec) typing. Strains were also analysed for the presence of virulence genes: staphylococcal enterotoxin (sea, seb, sec, sed, see, seg, seh, sei, sek, sel, sem, seo and sep), Panton-Valentine leukocidin (pvl), exfoliative toxin (eta, etb and etd), and toxic shock syndrome toxin (tsst-1) genes [7, 8]. Antibiotic susceptibility testing was performed and interpreted according to the Clinical and Laboratory Standards Institute (CLSI) guideline [9].
Table 1: Phenotypic and genotypic characteristics of methicillin resistant *Staphylococcus aureus* (MRSA) isolated from retail food and contact surfaces in Singapore. MLST multi locus sequence typing, *spa* staphylococcal protein A, *pvl* panton-valentine leukocidin, *se* staphylococcal enterotoxin, *et* exfoliative toxin, *tsst-1* toxic shock syndrome toxin-1, *AMC* amoxycillin-clavulanic acid, *AMP* ampicillin, *FOX* cefoxitin, *CRO* Ceftriaxone, *P* penicillin, *AK* amikacin, *AZM* azithromycin, *CIP* ciprofloxacin, *C* chloramphenicol, *CN* gentamicin, *NOR* norfloxacin, TE tetracycline, SXT trimethoprim-sulphamethoxazole

| Food stall | Year of isolation | Source | Molecular typing | pvl gene | Multi locus sequence typing (ST) | SCCmec | Staphylococcal enterotoxin genes | Exfoliative toxin genes | tsst-1 | Beta lactams | Non beta lactams |
|------------|------------------|--------|------------------|----------|---------------------------------|--------|---------------------------------|------------------------|--------|----------------|-----------------|
| A 2011     | Sliced Onion     | ST80 t1198 IV | -              | +        | -                              | IV     | -                              | -                      | +      | R              | R               |
| A 2011     | Prawn fritters   | ST80 t1198 IV | -              | +        | -                              | IV     | -                              | -                      | +      | R              | R               |
| A 2011     | Fried egg        | ST80 t1198 IV | -              | +        | -                              | IV     | -                              | -                      | +      | R              | R               |
| B 2013     | Gloves           | ST6 t304 IV | +              | -        | -                              | IV     | -                              | -                      | -      | R              | R               |
| B 2013     | Gloves           | ST6 t304 IV | +              | -        | -                              | IV     | -                              | -                      | -      | R              | R               |
Of the 227 *S. aureus* isolates, five (2.2%) from two food stalls were methicillin-resistant (Table 1) and belonged to Sequence Types (ST) 80 (spa type t1198) (isolated from sliced onion, prawn fritters, fried egg) and ST6 (spa type t304) (from swabs of food handlers’ gloves) [7]. All five isolates belonged to SCCmec type IV, and were *pvl*-negative. Unlike SCCmec types I-III strains which are often associated with nosocomial infections, strains belonging to SCCmec type IV are capable of colonising healthy individuals and may be community-associated. Some SCCmec type IV strains may be associated with livestock, suggesting a possible transmission between human and food-producing animals [10–12]. In Singapore, community-associated (CA) MRSA strains previously reported were isolated from clinical samples and belonged mainly to ST30, ST59 and ST772 [13]. To our knowledge, no ST6 strain has been reported in Singapore, whereas a ST80 CA-MRSA strain was previously reported in a local hospital in 2003, from a patient with a chin abscess [8]. *S. aureus* ST80, belonging to Clonal Complex 80 (CC80), was first reported in Denmark in 1993 and has been recognised as a major clone of CA-MRSA, widely spread across Europe [14–16]. Most ST80 strains reported elsewhere belong to SCCmec type IV, are *pvl*-positive, and are usually associated with severe skin/soft tissue infections and necrotising pneumonia [17]. Nonetheless, *pvl*-negative ST80 strains have also been reported in clinical cases overseas [18, 19]. ST6 MRSA is a double-locus variant of CC5, which is one of the five CCs from which major epidemic MRISA isolates are believed to have emerged [20]. ST6 MRSA was previously reported in human cases and carriers in Australia, Oman and United Arab Emirates (UAE) [16, 21, 22]. In particular, ST6 SCCmec type IV (t304) *pvl*-negative MRSA, a strain susceptible to non-beta-lactams, was previously isolated from patients overseas, suggesting that the ST6 strain isolated in our study may be capable of causing human infection [23]. In addition, CC6-ST6-t304 strains have previously been isolated from human (MSSA and MRSA), feral cat (MRSA) and camel (MSSA) suggesting that the strain may be transmissible across different host species [19, 24–26].

In this study, we detected enterotoxin genes *seb*, *sek*, and exfoliative toxin gene *etd* in ST80 strains; and enterotoxin gene *sea* in ST6 strains [7]. No other toxin genes were detected. Enterotoxins *sea* and *seb* are known to cause approximately 90% of staphylococcal food poisoning worldwide [27]. Enterotoxin *sek* is considered a staphylococcal enterotoxin-like protein, though its ability to cause food poisoning has yet to be demonstrated [28]. *etd* is an exfoliative toxin serotype known to be associated with mild forms of cutaneous infections such as abscesses and furuncles [29]. The presence of enterotoxin genes (*sea* and *seb*) in these MRSA isolates suggests the isolates’ potential to produce toxins, and cause staphylococcal food poisoning, if allowed to grow in sufficient numbers in food. Antimicrobial susceptibility results showed that all 5 MRSA isolates were phenotypically susceptible to amikacin, ciprofloxacin, chloramphenicol, gentamicin, norfloxacin, tetracycline, and trimethoprim/sulfamethoxazole. Susceptibility to at least 3 non-beta-lactams in MRSA is used as a proxy to define community-associated strains [30]. We found ST80 isolates resistant to azithromycin in addition to the beta-lactams tested (Table 1). An emerging trend of macrolide resistance has been reported in CA-MRSA from the human and livestock sectors [11, 31]. MRSA with resistance to additional antibiotic classes is a concern, and may reflect the increasing use of these antibiotics in the local clinical practice, which warrants further investigation.

In conclusion, we report two clinically relevant MRSA strains (ST80 and ST6) in ready-to-eat food and food contact surfaces at retail. Humans (food handlers), rather than animals, were likely the sources of contamination. Our limited findings suggest ready-to-eat food and food contact surfaces as potential environmental sources for colonisation and spread of MRSA in the community. To date, little is known about the transmission of MRSA infections through food and food contact surfaces, however their possible roles in the dissemination of specific MRSA lineages cannot be ruled out. The data warrant a more comprehensive and integrated (farm-to-hospital approach) surveillance of MRSA in Singapore and elsewhere.

**Abbreviations**

AK: Amikacin; AMC: Amoxycillin-clavulanic acid; AMP: Ampicillin; AZM: Azithromycin; C: Chloramphenicol; CA-MRSA: Community-associated methicillin-resistant *Staphylococcus aureus*; CIP: Ciprofloxacin; CN: Gentamicin; CRO: Ceftriaxone; E: Exfoliative toxin; FOX: Cefoxitin; mecA: Gene encoding methicillin-resistant *S. aureus*-specific penicillin-binding protein; MLST: Multi-Locus Sequence Typing; MRSA: Methicillin-resistant *Staphylococcus aureus*; NOR: Norfloxacin; P: Penicillin; PBP2: Penicillin-binding protein 2; Pvl: Panton-valentine leukocidin; SCCmec: Staphylococcal cassette chromosome; Se: Staphylococcal enterotoxin; Spa: Staphylococcal Protein A; ST: Sequence type; SXT: Trimethoprim-Sulphamethoxazole; TE: Tetracycline; Tss: Toxic shock syndrome toxin

**Acknowledgements**

Not applicable.

**Funding**

This study was funded by the National Environment Agency, Singapore.

**Availability of data and materials**

Please contact corresponding author for data requests.

**Authors’ contributions**

KTA, LYH, HCH and RAG were involved in the conception and design of the study. KTA, MLC and THK performed the identification and characterisation of isolates. KTA, LYH, THK, HCH, RAG and LCN were involved in the analysis and interpretation of the data. All authors read and approved the final manuscript.
Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Publisher’s Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details
1 Environmental Health Institute, National Environment Agency, 11 Biopolis Way, #04-03/04, Helios Block, Singapore 138667, Singapore. 2 Department of Infectious Diseases, Tan Tock Seng Hospital, 11 Jalan Tan Tock Seng, Singapore 304833, Singapore. 3 Diagnostic Bacteriology, Pathology Department, Singapore General Hospital, 20 College Road, Academia, Level 7, Diagnostic Tower, Singapore 169856, Singapore. 4 School of Biological Sciences, College of Science, Nanyang Technological University, 60 Nanyang Drive, Singapore 637551, Singapore. 5 School of Chemical and Biomedical Engineering, College of Engineering, Nanyang Technological University, 62 Nanyang Drive, Singapore 637459, Singapore.

Received: 7 June 2017 Accepted: 4 September 2017

Published online: 08 September 2017

References
1. Alexander F, Bittta K, Gladys K, Beatriz G-R, Katja A, Jens-Andre H, Annemarie K, Juliane B, Bernd A, Bernd-Akiko T. Methicillin susceptible and resistant Staphylococcus aureus from farm to fork impact on food safety. Technologija mes. 2011;160–3.
2. Jones TF, Kellum ME, Porter SS, Bell M, Schaffner W. An outbreak of community-acquired foodborne illness caused by methicillin-resistant Staphylococcus aureus. Emerg Infect Dis. 2002;8(1):82–4.
3. Klymans J, van Leeuwen W, Goessens W, Hollis R, Messer S, Heflandt V, Bruining H, Beck M, Rost J, Van Leeuwen N. Food-initiated outbreak of methicillin-resistant Staphylococcus aureus analysed by pheno- and genotyping. J Clin Microbiol. 1995;33(5):1211–8.
4. Hsu LY, Koh TH, Tan TY, Ito T, Ma XX, Lin RT, et al. Emergence of community-associated methicillin-resistant Staphylococcus aureus in Singapore: a further six cases. Singap Med J. 2006;47(1):20–6.
5. Tambayla PH, Kumarrasinghe G. Methicillin-resistant Staphylococcus aureus control at the National University Hospital, Singapore: a historical perspective. Ann Med Assoc Singap. 2008;3(10):855–60.
6. Veras JP, Camto LSD, Tong LC, Shupp JW, Cummings C, Santos OD. Cerqueira MMOP, Cantini A, Nicolli JR, Jett M. A study of the enterotoxigenicity of coagulase-negative and coagulase-positive staphylococcal isolates from food poisoning outbreaks in Minas Gerais, Brazil. Int J Infect Dis. 2008;12:410–5.
7. Aung KT, Lo JACY, Chau ML, Kang JSL, Choy J, Skow R, De Lencastre H. Epidemiology of emerging methicillin-resistant Staphylococcus aureus (MRSA) in Denmark: a nationwide study in a country with low prevalence of MRSA infection. J Clin Microbiol. 2005;43:1836–42.
8. Faria NA, Oliveira DC, Westh H, Monnet DL, Larsen AR, Skov R. De Lencastre H. Spread of a methicillin-resistant Staphylococcus aureus ST80 strain in the community of the northern-Netherlands. Eur J Clin Microbiol Infect Dis. 2007;26:723–7.
9. Deurenberg RH, Stobberingh EE. The evolution of Staphylococcus aureus. Infect Genet Evol. 2008;8:747–63.
10. Budimir A, Deurenberg RH, Bosnjak Z, Stobberingh EE, Cerqueira MMOP, Cantini A, Nicoli JR, Jett M. A study of the enterotoxigenicity of methicillin-resistant Staphylococcus aureus in Singapore. Infect Genet Evol. 2008;8:976–82.
11. Monecke S, Slukni K, Hasan R, Ruppert A, Ghazal SS, Hakavai A, Slickers P, Ehricht R. Characterisation of MRSA strains isolated from patients in a hospital in Riyadh, Kingdom of Saudi Arabia. BMC Microbiol. 2012;12:146–54.
12. Weber S, Ehricht R, Slickers P, Abdel-Wareth L, Donnelly G, Pitout M, Stefan M. Genetic fingerprinting of MRSA from Abu Dhabi. Vienna. Eur J Clin Microbiol Infect Dis. 2010;2010.
13. Udo EE, Al-Lawati BA-H, Al-Muharri Z, Thukral SS. Genotyping of methicillin-resistant Staphylococcus aureus in the Sultan Qaboos University Hospital, Oman reveals the dominance of Panton-Valentine leukocidin-negative ST6-IV/t304 clone. New Microbes New Infect. 2014;2(4):100–5.
14. Sonnevend A, Blair I, Alkaabi M, Jumaa P, Al Haj M, Ghazavi A, Akawi N, Jouhar FS, Hamadeh MB, et al. Change in methicillin resistant Staphylococcus aureus clones at a tertiary care hospital in the United Arab Emirates over a 5-year period. J Clin Pathol. 2012;65:178–82.
15. Harastani HH, Tokajian ST. Community-associated methicillin-resistant Staphylococcus aureus clonal complex 80 type IV (CC80-MRSA-IV) isolated from the Middle East: a heterogeneous expanding clonal lineage. PLoS One. 2014;9(1):11.
16. Bierowiec K, Ploneczka-Janeczko K, Rypula K. Is the colonisation of Staphylococcus aureus in food animals, humans and the environment, Aix-en-Provence. France. Springer Nature Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.