Increased Community-Associated Infections Caused by Panton-Valentine Leukocidin–Negative MRSA, Shanghai, 2005–2014

Min Li, Yanan Wang, Yuanjun Zhu, Yingxin Dai, Xufen Hong, Qian Liu, Tianming Li, Juanxu Qin, Xiaowei Ma, Huiying Lu, Jie Xu, Michael Otto

During 2005–2014, community-associated methicillin-resistant Staphylococcus aureus infections increased in Shanghai, China. Most infections were caused by sequence type 59 S. aureus that lacked Panton-Valentine leukocidin. This finding challenges the notion that Panton-Valentine leukocidin is necessary for epidemiologic success of community-associated methicillin-resistant S. aureus.

In the United States, community-associated (CA) methicillin-resistant Staphylococcus aureus (MRSA) infections in otherwise healthy persons in the community, first reported in the late 1990s (1), have reached epidemic dimensions (2). Despite considerable research efforts, the molecular underpinnings of the epidemiologic success of CA-MRSA are still not completely understood. Most typically connected with CA-MRSA is Panton-Valentine leukocidin (PVL). However, the role of PVL in CA-MRSA infection is controversial, primarily because of contradictory results from studies of animal infection models (3). A common belief is that if a clone from a patient with CA–S. aureus infection is positive for PVL, the S. aureus is probably a more dangerous clone and the patient would require specific care (4).

In the United States, virtually all CA-MRSA infections are caused by a PVL-positive clone of pulsed-field type USA300 (5). CA-MRSA infections with USA300 have also occasionally occurred outside the United States and adjacent regions. However, according to a recent study, they are derived from multiple importation events, suggesting that further spread in those locations is unlikely (6). Rather, global CA-MRSA infections are caused by geographically divergent clones that are unrelated to USA300. Like USA300, most of them contain PVL genes (2), although PVL is extremely rare in hospital-associated MRSA clones. This epidemiologic correlation is the predominant basis of the notion that PVL is causally associated with the enhanced virulence potential of CA-MRSA clones (7). Of note, despite generally enhanced virulence in animal models at levels similar to that of USA300 (8), the number of infections caused by global CA-MRSA clones remains limited (9,10). This situation raises the question of whether non-USA300 global CA-MRSA lineages have the potential to further intensify infection frequency and severity, and if so, whether PVL would be a necessary factor in such a scenario.

The CA-MRSA lineage that predominates in China and many other parts of Asia, thus threatening the largest global population, is sequence type (ST) 59 (10). Recent studies performed in Taiwan and northern Vietnam found a correlation between a PVL-positive subset of ST59 (Taiwan clone) and infection, but PVL-negative ST59 (Asia–Pacific clone) was found to be a largely noninfectious colonizer (11,12). Therefore, a causal relationship between PVL and infection has also been proposed for that CA-MRSA lineage.

The Study
We studied S. aureus isolates collected over 10 years (2005–2014) at Shanghai Renji Hospital, Shanghai, China, a large teaching hospital at which >10,000 patients from the entire Shanghai metropolitan area are admitted each day. We obtained 2,048 infectious S. aureus isolates and characterized them by multilocus sequence and spa typing, antibiotic resistance profiling, determination of the staphylococcal cassette chromosome (SCC) mec type (encoding methicillin resistance), and analytical PCR to determine presence of the lukSF genes encoding PVL. For isolates obtained during 2005–2010, we investigated randomly selected subsets (100 isolates/year); for isolates obtained during 2011, 2012, and 2014, we investigated all isolates. No isolates were collected in 2013. CA–S. aureus was defined as an isolate obtained from either an outpatient or an inpatient (including from general and urgent care and emergency rooms) ≤24 h after hospital admission, who lacked risk factors (contact with the hospital environment in the 6 months preceding the culture, S. aureus infection history, residence in a long-term care facility in the 12 months before culture, presence of a central vascular catheter at the

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time of infection, or recent use of antimicrobial drugs. These data were obtained by a review of medical records. The study was approved by the ethics committee of Renji Hospital, School of Medicine, Shanghai Jiaotong University, Shanghai (protocol RJ-H-2015–0221).

The percentage of methicillin resistance in the S. aureus infectious isolates was high, as is generally reported for China (13), and remained stable (at ≈70%) over the past 10 years (Table; Figure, panel A). In contrast, during the same time, resistance among CA–S. aureus infections was lower and fluctuated over time (Figure, panels B, C).

### Table. Characteristics of Staphylococcus aureus isolated at Shanghai Renji Hospital, Shanghai, China, 2005–2014*

<table>
<thead>
<tr>
<th>Year</th>
<th>Total†</th>
<th>HA-MSSA</th>
<th>HA-MRSA</th>
<th>CA-MSSA</th>
<th>CA-MRSA</th>
<th>Invasive among CA-MRSA</th>
<th>CA</th>
<th>HA-MSSA</th>
<th>MRSA</th>
<th>ST59</th>
<th>ST59</th>
<th>CA-MRSA</th>
<th>CA-MRSA</th>
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<tr>
<td>2010</td>
<td>600</td>
<td>(21)</td>
<td>(68)</td>
<td>(8.6)</td>
<td>(2.4)</td>
<td>(11)</td>
<td>(11)</td>
<td>(89)</td>
<td>(30)</td>
<td>(70)</td>
<td>(3.0)</td>
<td>(0.8)</td>
<td>(50)</td>
</tr>
<tr>
<td>2011</td>
<td>478</td>
<td>(21)</td>
<td>(66)</td>
<td>(9.5)</td>
<td>(3.5)</td>
<td>(67)</td>
<td>(13)</td>
<td>(87)</td>
<td>(30)</td>
<td>(70)</td>
<td>(4.0)</td>
<td>(1.5)</td>
<td>(67)</td>
</tr>
<tr>
<td>2012</td>
<td>470</td>
<td>(26)</td>
<td>(60)</td>
<td>(7.9)</td>
<td>(5.6)</td>
<td>(67)</td>
<td>(14)</td>
<td>(86)</td>
<td>(34)</td>
<td>(66)</td>
<td>(8.2)</td>
<td>(3.8)</td>
<td>(78)</td>
</tr>
<tr>
<td>2013</td>
<td>470</td>
<td>(24)</td>
<td>(59)</td>
<td>(9.8)</td>
<td>(7.4)</td>
<td>(51)</td>
<td>(17)</td>
<td>(83)</td>
<td>(34)</td>
<td>(66)</td>
<td>(9.6)</td>
<td>(5.3)</td>
<td>(52)</td>
</tr>
</tbody>
</table>

*CA, community-associated; HA, hospital-associated; MRSA, methicillin-resistant S. aureus; MSSA, methicillin-sensitive S. aureus; ST, sequence type.
†For 2005–2010, of a total of 2,681 isolates collected, 100 randomly selected isolates/year selected by using a random sample of data (SPSS, Chicago, IL, USA). For 2011, 2012, and 2014, all isolates obtained at the hospital were tested.
rose considerably, from 21% to 43% (p = 0.0108, Fisher exact test). While the percentage of HA-MRSA infections remained stable, CA-MRSA infections increased significantly, from 2.4% (12/500 total S. aureus infections) during 2005–2010 to 7.4% (35/470) in 2014 (p = 0.0003, Fisher exact test) (Table; Figure, panel A). ST59 dominated among CA-MRSA infections; increasing relative frequency reached a level of 71.4% in 2014 (Table; Figure, panel A). This finding links the observed surge in the CA-MRSA infection rate nearly exclusively to the ST59 lineage.

Of the 56 ST59 CA-MRSA infections, most (34 [61%]) were skin and soft tissue infections (i.e., spontaneous pyogenic skin abscesses), but a considerable number were respiratory (10 [18%]) or blood (9 [16%]) infections. The percentage of invasive infections (as defined by isolation from an otherwise sterile site, such as primary skin and soft tissue infection, with subsequent isolation from the blood, lung, or other otherwise sterile body fluids) among ST59 CA-MRSA patients was high at 62.5% (35/56). Fatality rate was 14% (5/35) among patients with invasive infections. Multidrug resistance was frequent. In addition to being resistant to β-lactams, most of the recently (2014) isolated ST59 CA-MRSA was also resistant to erythromycin (23/25; 92%), clindamycin (23/25; 92%), gentamicin (2/25; 8%), levofloxacin (3/25; 12%), trimethoprim/sulfamethoxazole (4/25; 16%), fosfomycin (3/25; 12%), or rifampin (5/25; 20%). None was resistant to tetracycline, linezolid, or vancomycin.

The ST59 CA-MRSA isolates were genetically heterogeneous and belonged to 7 spa types, predominantly t437 (31/56; 55%), t216 (12/56; 21%), and t441 (6/56; 11%). This finding is in contrast to the scenario described for USA300 CA-MRSA isolates, which are closely related (14), and suggests independent acquisition of SCCmec elements by genetically divergent parental ST59 methicillin-sensitive S. aureus strains.

Of note, only 20 (36%) of the 56 ST59 CA-MRSA isolates that we obtained contained the lukSF genes encoding PVL (Figure, panel B), and presence of the PVL genes was not correlated with more severe (i.e., invasive) infection (Figure, panel C). The PVL-positive Taiwan clone (spa types t437/t441, lukSF+, SCCmec V) was responsible for only 20% of cases (Figure, panel D). Moreover, while the percentage of PVL-positive ST59 CA-MRSA isolates and those belonging to the Taiwan clone increased in 2012, probably because of dissemination of the Taiwan clone into China, those numbers recently declined, indicating that PVL and the Taiwan clone are not main driving forces explaining the increase and current high percentage of CA-MRSA infections in Shanghai (Figure, panel B). Also, these subsets were not correlated with a specific infection type (Figure, panels C,D). Last, the Taiwan clone was not more frequently involved with invasive infections than were other ST59 CA-MRSA isolates (Figure, panel D).

Conclusions

CA-MRSA infections caused by a non-USA300 clone increased significantly in a highly populated area in China. Whether our findings are representative of all of China and adjacent countries remains to be addressed. Our findings do not support the previously indicated correlation of the PVL-positive ST59 subset (Taiwan clone) with infection (11,12). Thus, our study provides epidemiologic evidence challenging the widespread notion about a significant role of PVL in CA-MRSA dissemination in the ST59 lineage and in general. Inasmuch as our findings underscore the idea that the development of CA-MRSA clones is less limited to specific genetic backgrounds than previously thought, they underscore that novel successful CA-MRSA clones will probably continue to emerge.

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**CDC PROVIDES INFORMATION ABOUT MRSA SKIN INFECTIONS. CHANCES ARE, YOU’LL NEED IT.**

Recent data suggest that MRSA in the community is increasing. CDC encourages you to consider MRSA in the differential diagnosis for patients presenting with signs and symptoms of skin infections (red, swollen, painful, may be referred to as a spider bite by patient) especially those that are purulent (fluctuant or palpable fluid-filled cavity, yellow or white center, central point or “head,” draining pus, or possible to aspirate pus with needle or syringe).

Incision and drainage constitutes the primary therapy for purulent skin infections, including those caused by MRSA. Based on clinical assessment, empiric antimicrobial coverage for MRSA may be warranted in addition to incision and drainage. Obtaining specimens for culture and susceptibility testing are useful to guide therapy, particularly for those who fail to respond adequately to initial management.

MRSA skin infections can develop into more serious infections. It is important to discuss a follow-up plan with your patients in case they develop systemic symptoms or worsening local symptoms, or if symptoms do not improve within 48 hours.

**For more information, please call 1-800-CDC-INFO or visit www.cdc.gov/MRSA.**

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