Staphylococcus aureus (MRSA and CA-MRSA strains) in South America: comparative review to emergence of strains in North America and worldwide

Staphylococcus aureus (cetas MRSA e CA-MRSA) na América do Sul: revisão comparativa com emergência de cepas na América do Norte e no mundo

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ABSTRACT

Background: In the last few years, 3 different strains of MRSA have emerged: Community-associated Methicillin-resistant S. aureus (CA-MRSA), Hospital-associated (HA-MRSA), and Livestock-associated (LA-MRSA). The most common CA-MRSA strain is USA 300 lineage. In Brasil, this superbacteria is an important public health problem, once they are associated with severe infections (sepsis, shock and osteomyelitis), high mortality rates (including babies) and low response to usual treatments.

Aim: To review attempts to compare CA-MRSA strains in South America and propose an interconnection with patterns of North America and worldwide strains. Methods: Non-systematic review. Findings: Epidemiological and Genotyping definitions were used to compare different strains in different continents. Thus, we could determine ST30+ as the most common lineage in the Brazil and South America, USA 300 lineage as the most common in North America and ST80+ as the most common in Europe. Conclusion: MRSA is a seriously public health problem in Brazil and worldwide. In few years scientist will need a better understand of bacteria-derived factors that participate in enhanced MRSA pathogenesis & host susceptibility. Also, scientists will need to improve tools for an early diagnosis and they will need to enhance preventative/therapeutic modalities. However, new challenges will keep emerging.

Keywords: Staphylococcus aureus. Methicillin-resistant Staphylococcus aureus. Superinfection. Drug resistance.

RESUMO

Introdução: Nos últimos anos, emergiram três diferentes cepas de Staphylococcus aureus resistentes à meticilina (MRSA): Adquirida na Comunidade (CA-MRSA), Adquirida em Hospitais (HA-MRSA) e Adquirida na Pecuária (LA-MRSA). A ceps mais comum de CA-MRSA é a linhagem USA 300. No Brasil, essas superbactérias são um importante problema de saúde pública, uma vez que eles estão associados com infecções graves (sepsis, shock and osteomyelitis), altas taxas de mortalidade (incluindo mortalidade neonatal) e baixa resposta aos tratamentos usuais. Objetivo: Revisar as tentativas de comparação das cepas de CA-MRSA na América do Sul e propor uma interconexão com padrões de ocorrência de cepas na América do Norte e mundialmente.

Métodos: Revisão não sistemática. Resultados: Definições epidemiológicas e de genotipagem foram usadas para comparar diferentes cepas nos diferentes continentes. Determinamos que a linhagem ST30+ foi a mais comum no Brasil e na América do Sul, a USA 300 como a mais comum na América do Norte e a linhagem ST80+ como a mais comum na Europa. Conclusão: A emergência de MRSA é um problema de saúde pública no Brasil e no mundo. Num futuro próximo, os pesquisadores necessitam de um melhor entendimento dos fatores bacterianos envolvidos na patogênesis do MRSA e no aumento da suscetibilidade do hospedeiro. Será necessário otimizar os instrumentos para o diagnóstico precoce, bem como as modalidades de prevenção e os métodos de tratamento. No entanto, novos desafios continuarão emergindo.

INTRODUCTION

*Staphylococcus aureus* is a Gram-positive bacteria and it is a pore-forming toxic molecule producer. The leading cause of human bacterial infections worldwide usually lives as a commensal organism in humans and it affects asymptomatically 1/3 of human population. Nostrils are the most common colonization site, but there are known extra-nasal sites (e.g. throat, axilla, groin, perirectal area). Colonization sites are commonly related with subsequent infections.1-3

Transmission occurs via direct contact with the organism, in other words, skin-to-skin contact with colonised or infected individuals. Fomites and sexual transmission are other possible ways to get infected. The main predisposing risk factors are skin trauma, infection drug use, and poor personal hygiene. According to CDC, there are 5C-risk factors: crowding, contact, contaminated, compromised, and cleanliness.1,2

Symptoms varies widely in severity. The most common is skin and soft tissue infections (SSTI) – abscesses and cellulitis – responsible for around 90% of the worldwide cases. Invasive infections can cause bacteraemia, leading to complex cases and syndromes (e.g. Purpura fulminans, Myositis, Necrotizing fasciitis, Osteomyelitis, Necrotizing pneumonia and Endocarditis).1,2

Diagnosis is mainly based in culture from blood, tissue or pus. The exam has a high specificity, thus a culture failure is a strong evidence against *S. aureus* infection. Antimicrobial susceptibility can also be tested, but it does not discriminate differences between strains. It is important to highlight that selection of the appropriate treatment for infection requires antibiotic susceptibility patterns plus a careful assessment of patients’ history.1,2

*S. aureus* was first described in 1881 by Sir Alexander Ogston, when the infection was usually fatal because the lack of antibiotics.1 The beginning of a new age started with the discovery of Penicillin by Alexander Fleming in 1928. Previously untreatable infectious disease as *S. aureus* infection became treatable. Although, scientists and doctor’s happiness did not last for a long time, once antibiotic resistance started to be described.

*Staphylococcus* became resistant to sulfonamide in 1940s. In 1944, it became resistant to Penicillin (PRSA) by direct inactivation of the drug. In 1960s, it finally became resistant to Methicillin (MRSA) by acquisition of an alternative penicillin-binding protein (PBP-2) with lowered affinity to B-lactam.1,2

In the past 60 years, MRSA strains started an age of antibiotic resistance epidemic. Normally associated with nosocomial infections, besides this notion has changed greatly in the past few years. MRSA is among the leading causes of death by one single infectious agent, causing more deaths annually than HIV/AIDS in the USA.1-3

In the last 20 years, MRSA became hyper-virulent. Thus, it became multidrug resistant. The pandemic clones ST5, ST8, ST22, ST30, and ST45 are nowadays one of the most seriously health problems worldwide.1-3

DEFINITION & HISTORY: CA-MRSA, HA-MRSA & LA-MRSA

The discovery of Community-associated Methicillin-resistant *S. aureus* (CA-MRSA) occurred in 1990s, attributed to paediatric cases in the Midwestern United States. However, there are older description in cases associated with IV drug user in Detroit, MI, and Aboriginal populations in Western Australia (WA-1) – both strains were subsequently classified as MLST clonal complex (CC) 1 (Figure 1).4

Figure 1. Definitions: CA-MRSA, HA-MRSA and LA-MRSA.

Source: Prepared by the author.

In 2000, a different lineage emerged CC8 or “USA300”. The latter quickly eclipsed MW2 (USA400), becoming the primary cause of skin and soft tissue infections (SSTI) in the US, and greatly increasing the burden of community MRSA carriage and transmission. During the same period, genetically distinct CA lineages were reported from numerous countries.4

In the last few years CA-MRSA became a health problem worldwide. For instance, in the period of 2000-2003, it was responsible for more than 50% of USA infections while in France it was responsible for just 1-3%.1,3

According to CDC, CA-MRSA is a disease contracted within 48 hours of hospital admission by patients not having recently undergone surgery, haemodialysis, and prolonged hospitalization. Meanwhile, this is an epidemiological definition and it has lots of misclassifications, once CA-MRSA is an increasing cause of health care-acquired infections.1-3

CA-MRSA is traditionally regarded as MRSA strains causing infection (cellulitis or abscesses) in previously healthy young patients without prior healthcare contact, susceptible to most non-β-lactam antimicrobial agents, and carrying Panton Valentine (PVL) genes and SCCmec types IV and V.1-3

On the other hand, Hospital-associated Methicillin-resistant
**S. aureus** (HA-MRSA) is related to MRSA strains causing infections (bacteraemia) in patients with prolonged hospital stay, care in ICU’s, prolonged antibiotic treatment, surgical infections, close contact with MRSA-positive individuals. They are usually related with risk groups of population or risk factors: history of colonization/infection with CA, close contact (family) with a person colonized/infected with CA, membership of an indigenous community; being member of special communities (participation in contact sports, injection drug use, living in correctional facilities or shelters, military personnel, and men who have sex with men). HA-MRSA is less virulent than CA-MRSA.\(^1,2,5\)

Livestock-associated Methicillin-resistant *S. aureus* (LA-MRSA) or CC398 (most common strain) is related to MRSA in animals. It is widely disseminated among pigs in European countries with high-density pig farming (The Netherlands, Denmark and Germany), but it also affects veal calves, horse and dogs. The main concern about those strains is a possible adaptation to human, as already related in some Chinese Hospitals. Thus, animals will be an endless reservoir of infection.\(^1,2,6\)

**VIRULENCE**

*S. aureus* genome structure has 3 principal components: backbone of core genes, dispersed core of variable genes, and mobile genetic elements. The backbone of core genes is founded in all strains and they are highly conserved. Core variable genes (CV) are dispersed through the backbone composing a high variable group. MGE are constantly transferred and they are especially related to acquisition of PBP-2, an alternative penicillin-binding protein encoded by mecA and harboured in a genetic element: Staphylococcal cassette chromosome (SCCmec) that encodes resistance to methicillin. Depending on the particular SCCmec type, these mobile islands can confer multidrug-resistance.\(^1,7,10\)

MRSA defence is based in 2 mains strategies: Immune evasion and Virulence factors. The main components related to Immune evasion are: cytotoxins, immunomodulatory proteins, protease and factors that prevent immune cell recognition and killing (protein A/ capsule/ catalase). On the other hand, virulence factors are divided in 3 main groups: surface proteins, enzymes and secreted toxins.\(^1,7,10\)

**Virulence factors**

1. **Panton-Valentine Leucocidin (PVL)**

Leucocidins are molecules transferred via Mobile Genetic Elements that attack and lysed neutrophils (main cell involved in pathogenesis of infection associated with *S. aureus*). Leucocidin remain as the forefront of pathogenesis research initiatives.\(^1,9,10\)

Panton-Valentine leucocidin is a bicomponent pore-forming toxin located in prophage phiSA2pvl and encoded in genes lukS-PV and lukF-PV, responsible for induce necrosis and apoptosis in leukocytes by binding to host membranes and forming B-barrel pores that span the phospholipid bilayer.\(^1,9,10\)

2. **Enterotoxins**

High resistant (heat stable and resistant to enzymatic degradation) and variable (more than 20 types) toxins. Enterotoxins activate T cell population by passing the traditional pathway of MHC-TCR generating an excessive inflammatory response: leading to toxic shock, multi-organ failure and death. Thus, they are commonly related to immune-mediated diseases: Kawasaki disease, atopic dermatitis, and chronic rhinosinusitis.\(^1,8,10\)

3. **Spa**

Polymorphism in Protein A coding sequence.\(^3\)

4. **Multilocus sequence type (MLST/ ST) or CC**

Multi-Locus Sequence Typing (MLST) involves the sequencing of fragments from seven “housekeeping” genes (arcC, aroE, glpF, gmk, pta, tip and yqiL) yielding unique sequence types (STs). STs sharing identity at the majority of these loci are grouped into Clonal Complexes (CCs) encompassing related lineages of MRSA. Scientist discovered most MRSA disease worldwide was caused by five major CCs: CC5, CC8, CC22, CC30 and CC45.\(^5\)

5. **Staphylococcal cassette chromosome (SCCmec)**

There are 8 SCCmec types (I-VIII). The most common is SCCmecIV. SCCmec types could be divided in 2 main groups: SCCmec I-III and SCCmecIV-V. The first group is related to HA-MRSA: large length, little movement and contain additional drug resistance to several classes of antibiotics. The second group is related to CA-MRSA: small length, spread easily, more mobile and drug resistance only to beta-lactam.\(^10\)

6. **Toxins: alfa-toxin & Phenol-Soluble Toxins (PSM)**

a) **Alfa-toxin**

Pore-forming toxin and well-stabilised major virulence determinant. It is not lytic to human neutrophils, but lyses other immune cells; promotes an influx of neutrophils to infected lungs and enhance neutrophil adhesion to endothelial cell.\(^1,9,10\)

b) **Phenol-soluble modulins (PSM)**

Amphipathic/alpha-helical peptides belonging to surfactant-like class, which responsible for neutrophil lysis and for immune response avoiding.\(^1,9,10\)

7. **Mobile-genetic element (MGE)**

Newly acquired sequence of genes that can move around in genome and transferred between strains by horizontal gene transfer leading to selective advantage. MGE are determinants of colonisation/transmissibility rather than virulence. Examples: bacteriophages encoding toxins, pathogenicity
and composite islands (SaPIs), plasmids and transposons, SCCmec, SCC (non-mec), genomic islands. They are more prevalent among the USA lineage. Therefore, MGE are one of the possible factors responsible for the success of adaptation of those clones across the globe.\textsuperscript{1,9,10}

8. Arginine-catabolic mobile element (ACME)

Genetic element acquired by CA-MRSA isolates unique to USA300. Physically linkage of ACME with SCCmec IV is mirrored by an epidemiological linkage. ACME aids primarily in USA300 colonization through the Arc mediated ammonification of the acidic skin environment, though this has never been experimentally verified.\textsuperscript{3}

**GENOTYPE**

Once epidemiological definitions and classifications are not well accurate, genetic typing methods are the most plausible form of classification. However, there is no single, stable genetic marker for CA-MRSA strains. PVL genes were proposed as markers, but there are variations (positive and negative strains).\textsuperscript{11,12}

The most used genotyping methods are:\textsuperscript{12}

1. Spa sequence typing: sequence polymorphism in the variable X region of the spa gene for *S. aureus* surface protein A.

2. Multilocus sequence typing (MLST): sequencing of fragments from seven “housekeeping genes” (arc, aroE, glpF, gmK, pta, tpi yqiL) yielding unique sequence types (ST). ST sharing identity at the majority of these grouped into Clonal Complexes (CC) encompassing related lineages of MRSA. For example, the 5 major types clonal complexes are CC5, CC8, CC22, CC30 and CC45.

3. SCCmec typing (PFGE): MGEs – highly discriminatory approach that can identify genomic rearrangements (insertions/deletions) and it classifies strains into PFGE-types USA100-1200.

4. Macrorestriction pattern analysis: analysis of restriction polymorphisms of the whole chromosome.


**GLOBAL VIEW**

There are around 20 distinct genetic lineages of MRSA around the world. The 5 global predominant are ST1-IV (WA-1, USA400); ST8-IV (USA 300), ST30-IV (South West Pacific clone), St 59-V (Taiwan clone), ST80-IV (European clone). There are 2 strains classified as pandemic: ST8-IV (USA300 lineage); ST30-IV (South West Pacific clone SWP) (Figure 2).\textsuperscript{4,6,13}

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**Figure 2.** Global distribution of MRSA strains.

Source: Prepared by the author.
The most prevalent area of infection by MRSA is the EUA. The most prevalent strain is “USA300” lineage. A reasonable explanation to this condition are the different social panoramas between EUA and Europe. The USA infection panorama is homogeneous with a predominant lineage, while in Europe there is a heterogeneous panorama once none lineage predominates. However, ST80-IV is the most common lineage in Europe.4,6,13 The prevalence of MRSA worldwide could be divided in 3 main zones: Americas and Asia (highest prevalence), East Europe (intermediate prevalence) and West Europe (low prevalence).6,13

The decline of prevalence of MRSA worldwide could be divided in 3 main zones: Americas and Asia (low decline), East Europe (intermediate decline) and West Europe (high decline). This data contribute to a social analysis of the infection. Nations with more precarious health systems are less likely to fight and contain MRSA infection, while developed nations where MRSA is combated quickly and effectively.4,6,13

In South America, the 2 predominant HA-MRSA clones are: Brazilian clone (sequence type ST239) – related to Russian clones, which bears the SCCmecIII (MRSA0ST239-III), and the Chilean/ Cordobes clone (MRSA-ST5-I). However, USA300 has disseminated through Latin America as the main cause of CA-MRSA. In Uruguay and Brazil, USA 300 CA-MRSA isolates have been reported to display distinct genetic characteristics (MRSA-ST30-IVc). Interestingly, this strain, recently dubbed “USA300-LV” (Latin America Variant) appears to be a separate lineage which may be distinguished from the North America strain (USA300-0114) by various molecular features including SCCmec subtype (IVc) and absence of ACME. The dissemination of CA-MRSA isolates in South America hospital environments was first reported in Colombia in 2009. In the Andean countries (Colombia, Venezuela, Peru, and Ecuador), the CA-MRSA isolates present similar characteristics to the USA300 clone, including MRSA-ST8-IVc-E, PVL positivity, and multi-susceptibility.10,14

In the USA, the predominant CA-MRSA clone is ST8+ SCCmecIVA - PVL positive or USA300. Meanwhile, a few years ago USA400 was the predominant clone, but after a clonal expansion it became USA300. ST1+ or USA400 is still predominant in northern areas of America (e.g. Alaska and Canada), while USA300 is predominant in USA itself. USA300 lineage cause the majority of complicated SSI infection. After antibiotics discovery, this reality has changed. However, bacteria became resistant and new and more potent drugs were developed.4,13

The fast emergence of USA lineage comes from the combination of 3 main elements: newly acquired virulence genes (gene acquisition), altered expression of common virulence determinants (altered gene regulation), and alterations in protein sequences that increase fitness (sequence polymorphisms – protein sequence divergence). However, it is likely that no single explanation can suffice, and that MRSA represents a continuously emergent phenomenon driven by multi-factorial interactions between the classic triad of host, pathogen, and environment.3

Different from the USA, MRSA infections are less prevalent and more heterogeneous in Europe. In 2003, the first European case was reported in Greece (country where MRSA infections are more prevalent in Europe). The most common clone in Europe is ST80+, but in certain areas there are other clones. For example, ST22-IV or EMRSA-15 in UK; ST239/spa3 (t037)/SCCmecIII, ST239/spa351(t030)/SCCmec III and ST8/spa1 (t008)/SCCmecIV in Russia. The Russian clones are divided according to geographic areas: ST239 is more prevalent in European and Eastern Russia, while ST8 is more prevalent in Siberia Russia. The clones present in Siberia Russia are very similar to Brazilian clones. It could be possible because of migration of strains and transference of genetic material.15

In Asia, the 2 predominant clones are ST59- IV (USA1000) and ST59-V (Taiwan clone). However, Asia is a very diverse region with many different clone: ST22-IV & ST772-V in India; ST59-V in Taiwan, Vietnam and China; ST72+ in Korea; ST121+ and ST834+ in Cambodia; and ST30+ in South Pacifica, Australia and Russia. It also common to Asian strains be PVL negative.2,4,13,16 In Japan, the most prevalent clone is the Japanese ST8 SCCmecIVa (designatedST8 CA/J). The first case was reported in 2003. The Japanese clone is very similar to the USA clone ST8 SCCmecIVa (USA300), but with marked diversity in accessory genes. For example, ST8 CA/J possessed enhanced clytotic peptide genes, but lacked the PVL phage and ACME, unlike USA300. From a genetic point of view, ST8 CA/J is a geographic variant of ST8 CA-MRSA, which is one of the most disseminated lineages. However, ST8 CA/J – PVL negative is the most common clone some PVL positive strains were identified in Japan like: ST6 and ST59.2,4,13,16

Negative point: MRSA classification is not clear whether the incidence of CA-MRSA infections reported in many parts of the world is adequately reflected or influenced by less stringent testing and reporting measure.2,4,13,16

TREATMENT

Until the last century, there was no treatment to S. aureus infection. After antibiotics discovery, this reality has changed a lot. Penicillin was an inexpensive, non-toxic and high effective drug used against Staphylococcus in the first years. However, bacteria became resistant and new and more potent drugs were developed.

a) Antibiotics

Nowadays, SSTI are treated empirically based on antibiotic guidelines. Except abscesses, which, according to literature, are treated with incision and drainage.1

In less severe cases, MRSA infections are conventionally treated with oral antibiotics. Clindamycin oral is the gold pattern treatment. Also, tetracyclines (doxycycline, minocycline) could be used, but doctors should take care of side effects. Possible adjunct drugs are Rifampicin and Fusidic
Acid, but they should be never used alone.  

Linezolid (Oxazolidinone) is an expensive drug reserved for serious infections (comple STTI and MRSA pneumonia) when oral drugs are not an option. Doctors should take care of toxicity: myelosuppression, peripheral neuropathy, optic neuritis and lactic acidosis.  

In severe cases (e.g. bacteraemia), MRSA infections are conventionally treated with intravenous antibiotics. Vancomycin IV is the gold pattern treatment. Doctors should take care of nephrotoxicity with high doses.  

Despite the vast current therapeutic arsenal, multiresistant strains are emerging. Resulting in the need for new and more effective drugs.  

b) Experimental drugs  
These drugs are rapidly effective, but they are related to hypersensibility cases. They are divided in 3 subgroups:  

1. Glycopeptides derivates: telavancin, dolbavancin, ontanvancin;  
2. Anti-MRSA B-lactam: ceftaroline, ceftabiprole;  

c) Vaccine  
There are not any clinically approved vaccine. The researches focus in 2 vaccine mechanisms: enhance opsonophagocytosis or antitoxin. The main reason to vaccine failure is the fact that most researches focus on vaccines based on opsonophagocytosis process. But the bacteria has a neutrophil lysis mechanism after being engulfed which makes phagocytosis ineffective.  

d) Immunotherapy  
Passive immunization with antibodies to sequester main virulence determinants of *S. aureus* may represent a valuable alternative to active immunization in the future, but it requires more research. Example: Monoclonal alpha – toxin.  

**FINAL CONSIDERATIONS**  
MRSA is a seriously health problem across the globe because: lack of knowledge on the genetic control of the bacteria, its high mortality and its multidrug resistance. Thus, in the next years, researchers should focus on better understand the bacteria-derived factors that participate in enhanced MRSA pathogenesis & host susceptibility; improve tools for an early diagnosis; enhance preventative and therapeutic modalities. Otherwise, we will face an increased number of infections, new antibiotic resistance and low positive clinical outcomes.  
Work on developing research for better understanding of the genetic mechanisms involved will bring the possibility of creating vaccines or immunotherapies able to contain the bacteria more effective and specific way.  

However, new challenges remain emerging. For instance: drug resistance to the most potent antibiotics actually used (e.g. vancomycin, daptomycin, linezolid); interactions between HA-MRSA, CA-MRSA, LA-MRSA resulting in possible new and more resistant strains; spread by new reservoirs; pandemic clones and not genotyped strains. Thus, scientist will need to continuously develop new methods to identify markers of hypervirulence, transmissibility and persistence; and new/more effective treatments.  

**ACKNOWLEDGEMENTS**  
I would like to thank Dr Cristiane Cunha for her support during my Microbiology monitoring and in this thesis process. I would like to thank Dr Mal Horsburgh and Dr Viv Dillon for their extraordinary support during this year abroad and in this thesis process. Also, this project would have been impossible without the financial support of the Program Science without Borders-UK (SwB-UK) and the National Research Council (CNPq-BR).  

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