Methicillin-Susceptible, Vancomycin-Resistant *Staphylococcus aureus*, Brazil

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We report characterization of a methicillin-susceptible, vancomycin-resistant bloodstream isolate of *Staphylococcus aureus* recovered from a patient in Brazil. Emergence of vancomycin resistance in methicillin-susceptible *S. aureus* would indicate that this resistance trait might be poised to disseminate more rapidly among *S. aureus* and represents a major public health threat.

Acquisition of high-level vancomycin resistance by *Staphylococcus aureus* represents a major public health risk because this antimicrobial drug continues to be the first-line and most inexpensive therapy to treat methicillin-resistant *S. aureus* (MRSA) despite concerns about its clinical efficacy. Recently, we described vancomycin-resistant MRSA (VR-MRSA) recovered from the bloodstream of a patient in Brazil (1). VR-MRSA belongs to sequence type (ST) 8 and is phylogenetically related to the community-associated (CA) MRSA USA300 genetic lineage that has rapidly disseminated in the United States and the northern region of South America (USA300-Latin American variant [USA300-LV]) (1,2). The vanA gene cluster in VR-MRSA was carried by a transferable staphylococcal plasmid (pBRZ01). We characterized a clinical isolate of vancomycin-resistant, methicillin-susceptible *S. aureus* (VR-MSSA) and document the in vivo transfer of the vanA gene cluster to 2 unrelated *S. aureus* strains causing bacteremia within the same patient.

**The Study**

On August 28, 2012, a blood culture from a patient in Brazil was reported positive for 2 isolates of MSSA while the patient was receiving daptomycin therapy (online Technical Appendix, http://wwwnc.cdc.gov/EID/article/21/10/14-1914-Techapp1.pdf). One MSSA isolate was susceptible to all antimicrobial drugs tested (VS-MSSA). The second isolate (VR-MSSA) had a vancomycin MIC of 256 μg/mL and was also resistant to gentamicin (Table 1). Both isolates were susceptible to daptomycin (MIC 0.5 μg/mL). Thirteen days earlier, 2 MRSA isolates, 1 of which was resistant to vancomycin (VR-MRSA), were recovered from the blood of the same patient (online Technical Appendix) (1). The daptomycin MICs for both MRSA strains were also 0.5 μg/mL.

Bacterial strains used in this study (Table 1) were grown in brain–heart infusion broth and agar. Plasmid pBRZ01 was transferred by using filter mating (3) and VR-MSSA and VR-MRSA as donors and VS-MSSA, VS-MRSA, and RN4220RF as recipients (Table 1). Transconjugants were selected on brain heart infusion medium containing vancomycin (32 μg/mL) and fusidic acid (25 μg/mL). Colonies from each mating experiment were subjected to digestion with SmaI and pulsed-field gel electrophoresis to investigate genetic relatedness (1). Plasmids carrying the vanA gene cluster were detected by using S1 nuclease digestion followed by hybridization with a vanA probe (4).

Whole-genome sequencing of VR-MSSA, VS-MSSA, and 2 representatives of the Chilean/Cordobes clone (M1, M91) was performed by using MiSeq PacBio RS II (Illumina, San Diego, CA, USA) to close the VR-MSSA genome (5) (online Technical Appendix). Phylogenetic analysis was performed by using the maximum-likelihood framework within RAxML v7.4.2 (6). For cell wall analysis, extraction and separation of peptidoglycan precursors was performed as described (7).

The PFGE patterns of both isolates (VR-MSSA and VS-MSSA) were indistinguishable, and in vitro growth rates were similar (Figure 1, panel A). S1 nuclease analyses indicated that VR-MSSA harbored a plasmid of $\approx$55
kb, which yielded a positive result when hybridized with a vanA probe (Figure 1, panels B, C) and was similar in size to the previously described vanA-containing plasmid pBRZ01 identified in the same patient (1). pBRZ01 of VR-MSSA was readily transferred to S. aureus RN4220 RF (efficiency = 3 × 10^8/donor). In vitro conjugative transfer of pBRZ01 between MRSA and MSSA strains recovered from the patient’s bloodstream was also readily achieved with efficiencies ranging from 4.3 × 10^7/donor to 2.5 × 10^6/donor. Acquisition of the pBRZ01 by corresponding strains resulted in resistance to vancomycin and gentamicin (Table 1).

Genome sequencing (online Technical Appendix) showed that VR-MSSA and VS-MSSA belong to clonal complex (CC) 5 (sequence type ST5) and harbor staphylococcal protein A (Spa) type t002. VS-MSSA and VR-MRSA have the characteristic CC5 genetic traits described by Kos et al. (8). The genome of VR-MSSA has a 2,906,602-bp chromosome and 3 extrachromosomal elements, including a plasmid of 55,713 bp identical to the previously described vanA-carrying pBRZ01 (1), which also harbors aac(6′)-aph(2′′), which confers gentamicin resistance.

Comparison of the core genomes of VR-MSSA and VS-MSSA showed only 20 single-nucleotide polymorphism differences, which suggested a close genetic relationship and probably representing the same organism that acquired pBRZ01. Phylogenetic analysis (Figure 2) confirmed that VR-MSSA is not a derivative of VR-MRSA (1).

Table 1. Staphylococcus aureus strains used in analysis of methicillin and vancomycin resistance, Brazil*

<table>
<thead>
<tr>
<th>Strain</th>
<th>Strain characteristics</th>
<th>MIC, µg/mL</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>VS-MRSA</td>
<td>Isolated from the bloodstream of a patient in Brazil</td>
<td>Vancomycin 0.5</td>
<td>(1)</td>
</tr>
<tr>
<td>VR-MRSA</td>
<td>Isolated from the blood of the same patient above and carrying vanA-containing pBRZ01</td>
<td>Gentamicin 0.5</td>
<td>(1)</td>
</tr>
<tr>
<td>VS-MSSA</td>
<td>Isolated from the blood of the same patient 13 d after isolation of VR-MRSA</td>
<td>Vancomycin &gt;256</td>
<td>32</td>
</tr>
<tr>
<td>VR-MSSA</td>
<td>Isolated from the same blood culture as VS-MSSA</td>
<td>Gentamicin 1</td>
<td>0.75</td>
</tr>
<tr>
<td>RN4220-RF</td>
<td>Laboratory strain of S.aureus used as recipient for mating experiments; fusidic acid and rifampin-resistant</td>
<td>Vancomycin 256</td>
<td>48</td>
</tr>
<tr>
<td>Transconjugant 1†</td>
<td>Transconjugant obtained from a mating experiment using VR-MSSA as donor and VS-MRSA as recipient</td>
<td>Gentamicin &gt;256</td>
<td>48</td>
</tr>
<tr>
<td>Transconjugant 2†</td>
<td>Transconjugant obtained from a mating experiment using VR-MRSA as donor and VS-MSSA as recipient</td>
<td>Gentamicin &gt;256</td>
<td>64</td>
</tr>
<tr>
<td>Transconjugant 3†</td>
<td>Transconjugant obtained from a mating experiment using VR-MSSA as donor and RN4220-RF as recipient</td>
<td>Gentamicin &gt;256</td>
<td>64</td>
</tr>
</tbody>
</table>

*VS-MRSA, vancomycin-susceptible, methicillin-resistant S. aureus; VS-MSSA, vancomycin-susceptible, methicillin-susceptible S. aureus; VR-MRSA, vancomycin-resistant, methicillin-susceptible S. aureus.
†A fusidic acid–resistant derivative was generated for mating experiments. All mating experiments were performed on brain heart infusion agar in the presence of vancomycin (32 µg/mL) and fusidic acid (25 µg/mL) to select for transconjugants.

Figure 1. Molecular typing of Staphylococcus aureus strains, Brazil. A) SmaI digestion of total DNA, followed by pulsed-field gel electrophoresis. Lane M, lambda ladder (molecular masses are indicated in kilobases on the left); lane 1, vancomycin-susceptible, methicillin-resistant S. aureus (VS-MRSA) isolated from the blood of a Brazilian patient (1); lane 2, vancomycin-resistant MRSA (VR-MRSA) isolated from the same patient and blood culture (1); lane 3, transconjugant 1 obtained from a mating experiment using vancomycin-resistant MSSA (VR-MSSA) as donor and RS-MRSA as recipient; lane 4, vancomycin-susceptible MSSA (VS-MSSA) isolated from the blood of the same patient 13 days after isolation of VR-MRSA; lane 5, VR-MSSA isolated at the same time as VS-MSSA; lane 6, transconjugant 2 obtained from a mating experiment using VR-MRSA as donor and VS-MSSA as recipient; lane 7, S. aureus RN4220 RF, lane 8, transconjugant 3 obtained using VR-MSSA as donor and RN4220 RF as recipient. B) S1 digestion of total DNA using the same strains shown in panel A. C) Hybridization with vanA probe using the same strains shown in panel A. Arrow indicates a positive signal for the vanA gene.
DISPATCHES

(isolated days before from the same patient) and emphasized the relationship of this strain to other vancomycin-resistant *S. aureus* and MRSA isolates with intermediate susceptibility to vancomycin (VISA).

We analyzed the pool of cytoplasmic peptidoglycan precursors of VR-MSSA grown in the absence or presence of 50 μg/mL of vancomycin for induction of the *vanA* cluster (Table 2). Tandem mass spectrometry analysis identified 3 nucleotide precursors ending in d-alanyld-alanine (UDP-MurNAc-pentapeptide), d-alanyl-d-lactate (UDP-MurNAc-pentadepsipeptide), and d-Ala (UDP-MurNAc-tetrapeptide). Upon induction with vancomycin, UDP-MurNAc-pentapeptide was not detected, and UDP-MurNAc-pentadepsipeptide accounted for most of the precursors (Table 2). These results indicate that the *van*-encoded enzymes required for incorporation of d-Lac into the precursors were fully functional in VR-MSSA. Our results also show that the *vanA* cluster was inducible by vancomycin in the *S. aureus* host because only a small proportion of the precursors (4%) ended in d-Lac in the absence of the drug.

Analyses of cell wall muropeptides from VR-MSSA showed 2 modifications of the L-Alaγ-α-D-Glu2-L-Lys3-d-Ala4-d-Ala5 stem peptide that are highly conserved in *S. aureus* strains, namely the amidation of the α-carboxyl of D-Glu2 to form D-iGln2 and the addition of a pentaglycine side chain on the ε-amino group of L-Lys3 by the Fem amino-acyltransferases (9). Induction of the *vanA* gene cluster

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**Figure 2.** Phylogenetic analyses of *Staphylococcus aureus* strains, Brazil. Whole-genome phylogenetic tree (dataset = 325,732 single-nucleotide polymorphisms, gamma-based log likelihood − 1909607.06950) of the *S. aureus* species showing position of vancomycin-resistant, methicillin-susceptible *S. aureus* (VR-MSSA) and vancomycin-susceptible MSSA (VS-MSSA) isolates sequenced for this study. Vancomycin-resistant *S. aureus* (VRSA) strains are shown in red. Numbers on branches are bootstrap values based on 1,000 resampling iterations. All branches without numbers had bootstrap values of 100%. Branch lengths are proportional to number of nucleotide substitutions per site (scale bars). Inset labeled CC5 is expanded to emphasize the polyphyly of VRSA strains. *Genomes sequenced for this study. M1 and M91 are members of the Chilean/Cordobes clone that is widespread in Latin America (online Technical Appendix, http://wwwnc.cdc.gov/EID/article/21/10/14-1914-Techapp1.pdf). CC, clonal complex.
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Table 2. Relative abundance of peptidoglycan precursors in Staphylococcus aureus strains, Brazil*

<table>
<thead>
<tr>
<th>Precursor</th>
<th>Monoisotopic mass</th>
<th>Abundance (%) in corresponding strains</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Observed</td>
<td>Calculated</td>
</tr>
<tr>
<td>UDP-MurNaAc-tetrapeptide</td>
<td>1,078.35</td>
<td>1,078.35</td>
</tr>
<tr>
<td>UDP-MurNaC-pentapeptide</td>
<td>1,149.37</td>
<td>1,149.39</td>
</tr>
<tr>
<td>UDP-MurNaC-pentadepsipeptide</td>
<td>1,150.37</td>
<td>1,150.37</td>
</tr>
</tbody>
</table>

*aBacteria were grown on brain heart infusion (BHI) broth (not induced) or BHI supplemented with vancomycin (50 μg/mL). VS-MSSA, vancomycin-susceptible, methicillin-susceptible S. aureus; VR-MSSA, vancomycin-resistant, methicillin-susceptible S. aureus; ND, not detected.

led to 2 major modifications. First, stem peptides ended in d-Ala4, indicating that the peptidyl- d-Ala4- d-Ala2 target of vancomycin, and d-Ala4-d-Lac termini, were fully eliminated. Second, the pentaglycine side chain was frequently missing (online Technical Appendix), indicating that replacement of d-Ala by d-Lac at the extremity of peptidoglycan precursors might have impaired the ability of Fem transferases to add Gly on L-Lys.

Conclusions

In this study, we demonstrated that the vanA-containing pBRZ01 plasmid previously described in MRSA was acquired by an invasive MSSA isolate within the same patient. Our findings also suggest that a vanA-containing plasmid (pBRZ01) was horizontally acquired at least twice during a short period by distinct S. aureus lineages within the same host (MRSA belonging to ST8 and an ST5 MSSA). VR-MSSA belongs to the ST5 lineage of CC5, a major hospital-associated lineage (10). The prevalent hospital-associated lineages circulating in Brazil are ST5 (New York/Japan and Pediatric clones), ST239 (Brazilian clone) and ST1 (USA400 clone) (11), and recent epidemiologic data showed replacement of the endemic Brazilian (ST239) clone by ST5 strains (11–13). Moreover, VR-MSSA is related to ST5 vancomycin-resistant S. aureus strains recovered in the United States (8) and to VISA isolates, including Mu50 and the hetero-VISA strain Mu3, initially recovered in Japan (14). It remains unclear why CC5 strains appear more likely to exhibit vancomycin resistance.

Our biochemical analysis indicates that the vanA gene cluster is fully functional in VR-MSSA, which leads to vancomycin-inducible production of d-Lac ending precursors and elimination of d-Ala- d-Ala containing peptidoglycan, as found in the enterococci (15). Our results also revealed a defect in side chain synthesis, although this did not prevent the synthesis of a functional and highly cross-linked peptidoglycan in VR-MSSA.

In summary, we report the in vivo acquisition of high-level vancomycin resistance in a bloodstream MSSA isolate. Of note, vanA-containing pBRZ01 was maintained even after the selective pressure of vancomycin had been removed, raising serious concerns about the possibility of further spread of resistance to this agent. However, no other MSSA strains containing this plasmid have been isolated so far in Brazil.

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References

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http://dx.doi.org/10.1046/j.1365-2958.1998.01114.x

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Assessment of Arbovirus Surveillance 13 Years after Introduction of West Nile Virus, United States
Results from the National Legionella Outbreak Detection Program, the Netherlands, 2002–2012
Seroprevalence for Hepatitis E and Other Viral Hepatitides in Diverse Populations, Malawi
Swine Influenza A(H3N2) Virus Infection in an Immunocompromised Man, Italy, 2014
Severe Pediatric Adenovirus 7 Disease in Singapore Linked to Recent Outbreaks across Asia
Hemagglutinin Receptor Binding of a Human Isolate of Influenza A(H10N8) Virus
Schmallenberg Virus Reoccurrence, Germany 2014
Detection of Circovirus in Foxes with Meningoencephalitis, United Kindom, 2009–2013
Readability of Ebola Information on Websites of Public Health Agencies, United States, United Kingdom, Canada, Australia, and Europe

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Technical Appendix

Case-Patient Summary

The patient was a 35-year-old man with mycosis fungoides, cocaine addiction, diabetes mellitus, and a history of repetitive skin and soft tissue infections. He was first hospitalized and treated for leg cellulitis in November 2011 and readmitted for recurrent skin and soft tissue infections and worsening concurrent conditions in June 2012. During his hospitalization, repetitive febrile episodes developed, and he had blood cultures positive for different *Staphylococcus aureus* isolates. The clinical course of the patient, *Staphylococcus aureus* isolates, and antimicrobial drugs provided are summarized in Technical Appendix Figure 1. Further details can be found in a prior publication by Rossi et al. (1).

Genome Sequencing

MiSeq assembly was performed by using ABySS (2), and PacBio assembly was performed by using the HGAP2 v2.1 de novo assembly pipeline (Pacific Biosciences, Menlo Park, CA, USA). Comparison of single-nucleotide polymorphisms (SNPs) between genomes used in this study was performed by using the short read alignment to the *S. aureus* genome for strain N315 as a reference and the Burrows-Wheeler Alignment tool (http://bio-bwa.sourceforge.net). SNP calls were detected by using samtools (http://samtools.sourceforge.net), and SNPs were identified as high quality if they were unambiguous and had a q score \( \geq 20 \). For preassembled genomes available from public databases, we used whole-genome alignment with reference to the N315 genome by using the show-snps utility of NUCmer (http://mummer.sourceforge.net). We created phylogenetic datasets by combining results of both SNP calling techniques above. We excluded potentially repeated regions from the reference genome that had \( >80\% \) nucleotide similarity over 100 bp on the basis of BLAST.
(http://blast.ncbi.nlm.nih.gov/Blast.cgi) of the genome against itself. All locations in the genome annotated as mobile genetic elements were also excluded.

**Phylogenetic Methods**

Maximum-likelihood phylogenies were constructed by using the POSIX-threads version of RAxML v8.0.19 (3). For SNP data, we used an ascertainment bias correction and a general time-reversible substitution model accounting for among-site rate heterogeneity by using the gamma distribution and 4 rate categories (ASC_GTRGAMMA model) for 100 individual searches with maximum parsimony random-addition starting trees. Node support was evaluated with 1,000 nonparametric bootstrap pseudoreplicates and filtering the optimal maximum-likelihood tree through the bootstrap trees so that node support values shown indicate the percentage proportion of bootstrap trees that contained a given internode branch.

**Peptidoglycan Precursor and Cell Wall Analyses**

Extraction of peptidoglycan precursors was performed as described (4). Separation of precursors by reversed-phase, high-performance liquid chromatography was conducted by using a C18 column (Nucleosil 4.6 × 250 mm; Macherey-Nagel, Hoerdt, France). Peaks were collected and precursors were identified by mass spectrometry (Qstar Pulsar I; Applied Biosystems, Courtaboeuf, France) (4). The peptide moiety of the precursors was sequenced by tandem mass spectrometry (4). Relative abundance of precursors was estimated by the percentage of the integrate peak area at 262 nm. Peptidoglycan was prepared as described (5), and covalently attached proteins were removed from peptidoglycan by digestion with pronase and trypsin. Muropeptides were obtained by digestion with lysozyme and mutanolysin. The ether bond internal to N-acetylmuramic acid was cleaved with 3% ammonia, and the resulting lactoyl peptides were separated by reversed-phase, high-performance liquid chromatography for sequencing by tandem mass spectrometry (Qstar Pulsar I).

**References**


Technical Appendix Table 1. Genome statistics for Staphylococcus aureus, Brazil*

<table>
<thead>
<tr>
<th>Strain</th>
<th>Coverage</th>
<th>No. contigs</th>
<th>Mean subread length, bp</th>
<th>Read length N50/assembly N50</th>
<th>NCBI Bioproject no.</th>
</tr>
</thead>
<tbody>
<tr>
<td>VR-MSSA (HP022)</td>
<td>800×</td>
<td>1,437</td>
<td>NA</td>
<td>NA/183054 bp</td>
<td>PRJNA262896</td>
</tr>
<tr>
<td>VS-MSSA (HP023)</td>
<td>575×</td>
<td>1,438</td>
<td>NA</td>
<td>NA/91,499 bp</td>
<td>PRJNA262282</td>
</tr>
<tr>
<td>M1 (HP012)</td>
<td>250×</td>
<td>1,813</td>
<td>NA</td>
<td>NA/8,727 bp</td>
<td>PRJNA262670</td>
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<tr>
<td>M91 (HP013)</td>
<td>85×</td>
<td>1,808</td>
<td>NA</td>
<td>NA/46,912 bp</td>
<td>PRJNA262672</td>
</tr>
<tr>
<td>VR-MSSA (PacBio)</td>
<td>81.1×</td>
<td>9†</td>
<td>4,955</td>
<td>6,305 bp/ 2.04 Mbp</td>
<td>PRJNA262896</td>
</tr>
</tbody>
</table>

*NCBI, National Center for Biotechnology Information; VR-MSSA vancomycin-resistant, methicillin-susceptible S. aureus; NA, not applicable; VS-MSSA, vancomycin-resistant, methicillin-susceptible S. aureus.
†Manual polishing and additional assembly resulted in 4 contigs (1 closed circular chromosome and 3 extrachromosomal elements).

Technical Appendix Table 2. Mass of muropeptide from vancomycin-susceptible and vancomycin-resistant, methicillin-susceptible Staphylococcus aureus, Brazil*

<table>
<thead>
<tr>
<th>Strain (growth condition)</th>
<th>R substituent of muropeptide</th>
<th>Monomer, n = 0</th>
<th>Dimer, n = 1</th>
<th>Trimer, n = 1</th>
<th>Tetramer, n = 2</th>
<th>Pentamer, n = 3</th>
<th>Hexamer, n = 5</th>
</tr>
</thead>
<tbody>
<tr>
<td>VS-MSSA</td>
<td>d-Ala – d-Ala Gly5</td>
<td>Calculated</td>
<td>844.40</td>
<td>1,599.76</td>
<td>2,355.12</td>
<td>3,110.47</td>
<td>3,865.83</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Observed</td>
<td>844.42</td>
<td>1,599.79</td>
<td>2,355.18</td>
<td>3,110.55</td>
<td>3,865.92</td>
</tr>
<tr>
<td>VR-MSSA (induced)†</td>
<td>d-Ala</td>
<td>Calculated</td>
<td>488.26</td>
<td>1,243.62</td>
<td>1,998.97</td>
<td>2,754.33</td>
<td>3,509.68</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Observed</td>
<td>488.25</td>
<td>1,243.6</td>
<td>1,998.97</td>
<td>2,754.34</td>
<td>3,509.71</td>
</tr>
<tr>
<td></td>
<td>Gly5</td>
<td>Calculated</td>
<td>773.38</td>
<td>1,528.72</td>
<td>2,284.08</td>
<td>3,039.44</td>
<td>3,794.79</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Observed</td>
<td>773.38</td>
<td>1,528.73</td>
<td>2,284.08</td>
<td>3,039.46</td>
<td>3,794.83</td>
</tr>
</tbody>
</table>

* VS-MSSA, vancomycin-susceptible, methicillin-susceptible S. aureus; VR-MSSA vancomycin-resistant, methicillin-susceptible S. aureus.
†Induction was performed with 10 µg/mL of vancomycin.
**Technical Appendix Figure 1.** Clinical course timeline of the patient, Brazil. Drugs used are indicated by colored rectangles: β-lactams in blue (cephalexin, cefepime, and piperacillin/tazobactam [Pip/Tazo]), clindamycin in green, glycopeptides in pink (vancomycin and teicoplanin), and daptomycin in yellow. The number in each rectangle corresponds to the number of days of treatment with the drug. Drugs are shown in the order in which they were added to therapy. The final days of hospitalization are not included. SSTI, skin and soft-tissue infection; VS-MRSA, vancomycin-susceptible, methicillin-resistant *Staphylococcus aureus*; VR-MRSA, vancomycin-resistant, methicillin-resistant *S. aureus*; VS-MSSA, vancomycin-susceptible, methicillin-susceptible *S. aureus*; VR-MSSA vancomycin-resistant, methicillin-susceptible *S. aureus*; VREF, vancomycin-resistant *Enterococcus faecalis*. 
**Technical Appendix Figure 2.** Diversity in the structure of muropeptides from *Staphylococcus aureus*, Brazil. Diversity of muropeptides is generated by variations at the C-terminus ($R_1 = \text{OH or } \text{D-Ala}^4 - \text{D-Ala}^5$), at the N terminus ($R_2 = \text{H or } \text{D-Gly}^5$) and by the extent of oligomerization (from $N = 0$ for monomers to $N = 6$ for heptamers).

**Technical Appendix Figure 3.** Muropeptides from vancomycin-susceptible, methicillin-susceptible *Staphylococcus aureus*, Brazil. A) Main monomers. The side-chain is assembled by aminoacyl transferases of the Fem family that sequentially add the first (FmHB), second, and third (FemA), and fourth and fifth (FemB) Gly residues. B) Dimer generated by $\text{D,D-} \text{transpeptidation.}$ The $\text{D,D-} \text{transpeptidases cleave the } \text{D-Ala}^4 - \text{D-Ala}^5 \text{ peptide bond of the acyl donor and link the carbonyl of } \text{D-Ala}^4 \text{ to amino group located at the extremity of the side chain of the acyl acceptor.}
Technical Appendix Figure 4. Muropeptides from vancomycin-resistant, methicillin-susceptible Staphylococcus aureus grown in the presence of 10 µg/mL vancomycin, Brazil. A) Main monomers. The C-terminal D-Lac is cleaved by D,D-carboxypeptidase and is not found in mature
peptidoglycan. Most (62%) of the muropeptide monomers did not contain any side-chain ($R_2 = H$ instead of Gly$_5$) because of impaired activity of FmhB with D-Lac ending precursors. B) Dimer generated by D,D-transpeptidation. All cross-links contain Gly$_5$ because unsubstituted stem peptides ($R_2 = H$) are not used as acyl acceptors by D,D-transpeptidases.

Tree

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begin taxa;
dimensions ntax = 50;
taxlabels
VRS3a
HP022
HP023
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SaMSHR1132
Sa08BA02176
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SaTCH60
SaMRSA252
SaJKD6159
SaLGA251
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SaRF122
SaMW2
SaMSSA476
Sa1181997
```
tree tree_1 = [&R] (((VRS3a:8.244799231807832E-4,(((HP022:3.96736443382444E-6,HP023:2.250523383219172E-5):3.411632724734676E-4,(SaED98:9.703221642786964E-4,(SaMSHR1132:0.5632346423186245,(((Sa08BA02176:5.10585699270055E-4,SaST398:5.38525420539065E-4):0.02318758006007828,(SaTCH60:0.0018606904107206922,SaMRSA252:0.0023361910795642177):0.0228694576846738):0.0208677517984276,(SaJKD6159:0.043048096207328516,(SaLGA251:0.012751949141110169,SaED133:0.014657491012749579):0.007702308459562556,SaRF122:0.020706149045281765):0.005715274275517768:0.005352806350899746):0.014270643684360873):0.004215184179548835,(((SaMW2:8.148223984244278E-4,SaMSSA476:9.399261875994422E-4):0.011723988543628189,Sa1181997:0.012031146247175885):0.003472808070423307,(((SaT0131:0.0011290280300672703,SaJKD6008:4.616415737154414E-4):2.4208588870377014E-4,SaTW20:5.876500448633564E-4):;end;

begin trees;

VRS6
VRS8
VRS9
SaMu50
SaMu3
VRS1
VRS2
SaST228
HP013
HP012
;

end;

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end;
Matrix

Large file, available from the authors.