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Comparative Genomics of Vancomycin-Resistant Staphylococcus aureus Strains and Their Positions within the Clade Most Commonly Associated with Methicillin-Resistant S. aureus Hospital-Acquired Infection in the United States

Veronica N. Kos,a,b Christopher A. Desjardins,b Allison Griggs,b Gustavo Cerqueira,b Andries Van Tonder,c Matthew T. G. Holden,c Paul Godfrey,b Kelli L. Palmer,a Kip Bodi,d Emmanuel F. Mongodin,e Jennifer Wortman,b Michael Feldgarden,b Trevor Lawley,c Steven R. Gill,f Brian J. Haas,ab Bruce Birren,b and Michael S. Gilmoreab

Departments of Ophthalmology and of Microbiology and Immunobiology, Harvard Microbial Sciences Initiative, Harvard Medical School, Massachusetts Eye and Ear Infirmary, Boston, Massachusetts, USAa; The Broad Institute, Cambridge, Massachusetts, USAab; The Wellcome Trust Sanger Institute, Hinxton, Cambridge, United Kingdomab;Tufts University School of Medicine, Boston, Massachusetts, USAa; Institute for Genome Sciences, University of Maryland School of Medicine, Baltimore, Maryland, USAa; and School of Medicine and Dentistry, University of Rochester, Rochester, New York, USAa

ABSTRACT  Methicillin-resistant Staphylococcus aureus (MRSA) strains are leading causes of hospital-acquired infections in the United States, and clonal cluster 5 (CC5) is the predominant lineage responsible for these infections. Since 2002, there have been 12 cases of vancomycin-resistant S. aureus (VRSA) infection in the United States—all CC5 strains. To understand this genetic background and what distinguishes it from other lineages, we generated and analyzed high-quality draft genome sequences for this species. In contrast to existing hypotheses on what predisposes this lineage to acquire Tn1546 from enterococci in a clonal cluster 5 (CC5) background, the most common hospital lineage causing hospital-acquired MRSA infection. To understand the nature of these strains, we determined and examined the nucleotide sequences of the genomes of all available VRSA. Genome comparison identified candidate features that position strains of this lineage well for acquiring resistance to antibiotics in mixed infection.

IMPORTANCE  Invasive methicillin-resistant Staphylococcus aureus (MRSA) infection now ranks among the leading causes of death in the United States. Vancomycin is a key last-line bactericidal drug for treating these infections. However, since 2002, vancomycin resistance has entered this species. Of the now 12 cases of vancomycin-resistant S. aureus (VRSA), each was believed to represent a new acquisition of the vancomycin-resistant transposon Tn1546 from enterococcal donors. All acquisitions of Tn1546 so far have occurred in MRSA strains of the clonal cluster 5 genetic background, the most common hospital lineage causing hospital-acquired MRSA infection. To understand the nature of these strains, we determined and examined the nucleotide sequences of the genomes of all available VRSA. Genome comparison identified candidate features that position strains of this lineage well for acquiring resistance to antibiotics in mixed infection.

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Address correspondence to Michael S. Gilmore, michael_gilmore@meei.harvard.edu.
Each of the 12 U.S. VRSA strains are believed to have resulted from acquisition of Tn\textsubscript{1546} from enterococci during the course of infection (15). Tn\textsubscript{1546} confers the VanA phenotype. Curiously, most of the Tn\textsubscript{1546} were from an Enterococcus faecalis donor (15–17), as opposed to the more common vancomycin-resistant Enterococcus faecium (18). Nine VRSA strains arose in southeast Michigan, where Tn\textsubscript{1546} transfer was mediated by a broad-host-range conjugative Inc18 plasmid (15–17, 19).

The repeated acquisition of vancomycin resistance by CC5, along with its involvement in the early acquisition of methicillin resistance and resistance to other antibiotics (11), suggests that it is genetically or biologically predisposed to horizontal acquisition of resistance and possibly other genes. Such transfer requires that donors and recipients coexist intimately in a mixed community, and that they achieve a population size that allows them to overcome inefficiencies and obstacles to transfer, genetic element establishment, and resistance expression. Thus, it was of interest to examine VRSA genomes for barriers to entry of foreign DNA as well as for traits that could foster their existence in mixed infection with potential resistance donors. We therefore generated high-quality draft genome sequences of the available 12 CC5 VRSA isolates from the first 11 VRSA cases in the United States and examined them for traits that may have predisposed this lineage to vancomycin resistance acquisition.

RESULTS
North American CC5 phylogeny. We determined a core gene sequence-based phylogeny for VRSA, based on 1,822 single-copy orthologs present in all genomes (Fig. 1). Strains do not cluster on site or time of isolation (Fig. 1), supporting their independent development into VRSA. Strain VRS3a, isolated in New York in 2004 (the only PFGE type USA800 strain) is the most divergent. MRSA strain JH1 and its GISA derivative JH9, which arose during antimicrobial therapy (13, 20), are nested deeply within the VRSA. This phylogeny shows that all VRSA strains stem from a monophyletic source, supporting the hypothesis that they harbor a trait or traits that predispose them for vancomycin resistance acquisition or expression.

Tn\textsubscript{1546} haplotype network. To determine whether Tn\textsubscript{1546} evolved along a path different from the rest of the chromosome (as would be expected if Tn\textsubscript{1546} were repeatedly, independently acquired as opposed to having been acquired once and then passed along vertically with the rest of the chromosome), Tn\textsubscript{1546} sequences were compared, and their relationships to each other and to possible donor elements were calculated (17, 19) (Fig. 2). Tn\textsubscript{1546} sequences segregate regionally, as opposed to temporally (e.g., transposons from strains isolated in New York, Pennsylvania, and Delaware [strains VRS2, VRS3a, and VRS11a or VRS11b] share features with each other that are not shared by strains from Michigan [e.g., strains VRS1 and VRS4], consistent with an independent acquisition model (Fig. 2). All Tn\textsubscript{1546} sequences from Michigan VRSA strains form a tight cluster with few single-nucleotide polymorphisms (SNPs). Tn\textsubscript{1546} in E. faecalis VRE5, coisolated with VRS5 (17), possesses a 491-bp duplication missing in VRS5, suggesting that either VRE5 is not the Tn\textsubscript{1546} donor or that the duplication occurred in the donor after transfer.
Tn1546 of candidate enterococcal donor VRE4 is identical to that in VRS4, as well as to transposons in putative enterococcal donor strains VRE6 and VRS6. Tn1546 from the plasmid in putative donor VRE9 differs from that in VRS9 by a single SNP (Fig. 2).

Tn1546 of strains VRS2 and VRS3a were reported to possess insertions of IS1251 between vanS and vanH, as well as copies of IS721 within orf1 that are inverted in the two strains (21, 22). The Tn1546 portions of these composite elements have 4 SNPs in common, distinguishing them from the VRS1 prototype and supporting their origination from a closely related source. Tn1546 of strain VRS11a/b shares a SNP with strains VRS2 and VRS3a, placing it between the New York and Pennsylvania elements, and the Michigan transposons. VRS11a/b possesses an ISFaecalis1 (ISEf1) insertion in the intergenic region between vanX and vanY (Fig. 2). VRS11a and VRS11b also possess a common SNP that is not found in other strains. Finally, even though isolated from a single patient, a third SNP distinguishes VRS11b from VRS11a, probably the result of microevolution within the patient.

Comparison of the Tn1546 haplotype network to the single-copy core gene phylogeny showed that there was little congruence and that the Tn1546 haplotype network was a poor predictor of core gene phylogeny (P < 0.0001 by the Shimodaira-Hasegawa test [23]). Phylogenies based upon typing schemes for the hypervariable repeat regions of clumping factor B (clfB), coagulase (coa), and protein A (spa) and based on SCCmec and agr loci also failed to show a consistent line of VRSA strain descent (see Table SIA and Fig. SIA, SIB, and SIC in the supplemental material). The results of each analysis strongly support the model that each VRSA isolate arose as the result of an independent transposon acquisition.

Diversity in plasmid content and Tn1546 location. Tn1546 resides on plasmids in all VRSA strains, not on the chromosome (even though the chromosome is much larger and would seem to be a more probable insertion target). Each VRSA strain harbors plasmids of enterococcal and staphylococcal origin, and in some cases, cointegrates of the two (Fig. 3), highlighting their history of cooccurrence in mixed infections with enterococci. Plasmids involved in Tn1546 acquisition vary. Strains VRS1 (24), VRS8, VRS9, and VRS11a/b possess large portions of the staphylococcal plasmid pSK41. pSK41 has been shown to enhance gene transfer from E. faecalis strains harboring pheromone-responsive plasmid pAD1 because of pheromone cross talk (25) and the pheromone encoded by a pSK41 gene is detected in the supernatant of strain VRS1 (15). The transposition of Tn1546 onto pSK41, with concomitant loss of the enterococcal donor plasmid, was previously reported to have occurred in strain VRS1 (24). In strains VRS8 and VRS9, Tn1546 also transposed to pSK41, but at unique sites. Strains VRS2, VRS3a, VRS4, VRS5, VRS6, VRS7, and VRS10 possess small fragments with sequence identity to pSK41, but these identities do not include pSK41 traH, the gene encoding the enterococcal plasmid pAD1 pheromone precursor.
Strains VRS2 and VRS3a acquired Tn1546 on large (>100-kb) enterococcal plasmids, which were retained (21, 22). Identical sequences surround Tn1546 in both strains, indicating a closely related source. These megaplasmids likely represent cointegrates of enterococcal plasmids pS177, p7573cB, and possibly pLG2 (Fig. 3). Mosaic plasmid structures are common in enterococci (26). Interestingly, the plasmid content of strain VRS7 is similar to that of strain VRS2, but with Tn1546 remaining on the broad-host-range Inc18 donor plasmid (Fig. 3). In addition to VRS7, VRS4 and VRS5 also retain Tn1546 on the Inc18 enterococcal donor plasmid (17). The sequence of the Inc18 plasmid in VRS10 was completed and found to possess an additional 2.97-kb transposon insertion not present in the others. In VRS4, VRS5, VRS7, and now VRS10, the location of Tn1546 in the Inc18 donor plasmid is identical, highlighting the role of this common element in the Michigan outbreak (16). In VRS6, Tn1546 was known to have transposed onto an S. aureus resident plasmid, with loss of the donor plasmid (16). In this strain, insertion occurred in a novel plasmid of apparent staphylococcal origin.

Strains VRS11a and VRS11b originated from a single patient in Delaware, and each strain possesses a cointegrate of enterococcal and staphylococcal plasmids (see Fig. S2 in the supplemental material) as confirmed by PCR. The cointegrate is a fusion of S. aureus plasmid pLUH02 (including genes for beta-lactamase, enterotoxin, cadmium resistance, and replication), and an enterococcal pCF10-like plasmid carrying phenomere-responsive genes. Tn1546 resides on this fusion plasmid in a location that is identical in both strains (Fig. S2).

No functional lesions identifiable in restriction-modification systems of most VRSA strains. The Sau1 restriction system (27) and a type III-like restriction system (28) represent known barriers to foreign DNA entry. The Sau1 restriction system consists of an endonuclease, HsdR, encoded by a gene distant from two pairs of specificity and modification subunits, which occur on genomic islands sensu2 and sensuβ (27). In VRSA, most Sau1 restriction systems are intact (Fig. 4A). Strain VRS3a possesses a mutation in the hsdR Sau1 endonuclease gene that results in a large truncation of the HsdR subunit (Fig. S2B in the supplemental material). Several polymorphisms in the Sαα sauCC5 hsdM1 copy of the HsdM modification subunit were observed (Fig. 4A), most representing minor amino acid changes. However, in the Sαα copy of HsdM in strain VRS9, a frameshift occurs, caused by an adenine duplication at nucleotide position 1213, which truncates the primary translation product (Fig. 4A). In the Sαα sauCC5 hsdS1 encoded specificity subunit of VRS11a/b, a nonsense mutation occurs in the second codon, likely causing a large truncation (Fig. 4A). In contrast to polymorphic copies in Sαae, only one polymorphism was noted in the Sαβ copies of HsdS and HsdM subunits (Fig. 4A). That polymorphism corresponds to an addition of 3 amino acids to the carboxy terminus of HsdS in strain VRS2, which seems unlikely to alter function. Lack of mutations in the Sαβ genes for HsdM and HsdS, as well as in most copies of counterparts in the Sαβ island, indicate that Sau1 is likely functional in all VRSA strains, except VRS3a.

A nonsense mutation in the type III-like restriction system has been reported to occur in Japanese CC5 strains (28). The impact of this specific mutation on function is unclear, although the type III-like restriction system clearly poses a barrier to DNA uptake (28). Only strain VRS10 possesses a polymorphism in this locus (MQ5_01626; a point change that creates a nonsense mutation in the 12th codon) that is likely to be of functional consequence (Fig. 4A). Strain VRS9 possesses a polymorphism that leads to a conservative N89D amino acid substitution. Otherwise, the type III-like restriction systems of all other VRSA strains appear to be intact.

Frameshift in dprA in all VRSA strains except VRS3a. One coding difference related to DNA metabolism that stood out in comparison of CC5 to non-CC5 genomes was an adenine duplication at positions 333 and 334 in dprA. This duplication introduces a frameshift (Fig. 4B), potentially eliminating or altering DprA function. This change was found in all North American CC5 strains except strain VRS3a. DprA influences DNA transformation efficiency in Bacillus subtilis (29). In addition to identifying potential loss-of-function mutations, such as the dprA frameshift, we also searched for other potentially functionally altering nonsynonymous-codon-changing SNPs unique to North American CC5 strains. We identified 45 SNPs common to all except the most divergent strain, VRS3a (see Table S2 in the supplemental material).

VRSA strains possess polymorphisms in the agr locus. Similar to recent observations for phage 80/81 isolates of S. aureus that were prevalent in hospitals in the early 1990s (30), several VRSA strains (VRS1, VRS2, and VRS8 [see Fig. S1C in the supplemental material]) possess lesions in a key global regulator of virulence, agr (31). The agr locus is known to attenuate during infection (32), and this parallels reduced vancomycin susceptibility (33, 34).

CC5 gene content differences include lack of a bacteriocin operon and presence of genes for diverse superantigens. We next examined North American CC5 strains, including VRSA, for differences in gene content that could promote coexistence in polymicrobial infection (see Table S3A and S3B in the supplemental material). Most of the variable gene content unique to the North American CC5 strains occurs within the vSaβ island (35, 36) (Table S3A; Fig. 5). Of potentially high importance, an operon (bsa) encoding a lantibiotic bacteriocin (active against other Gram-positive bacterial species [37, 38]), is absent in all members of the CC5 lineage. Instead, this island contains a unique cluster of enterotoxin genes (Fig. 5). Interestingly, this pattern—the absence of the bacteriocin operon and the presence of nearly identical complement of superantigens—also occurs in the vSaβ pathogenicity islands in the EMRSA-15 and MRS252/EMRSA-16 lineages that are prevalent in hospitals in the United Kingdom (39), even though the genetic backgrounds are highly dissimilar (Fig. 5). This suggests active selection for this version of vSaβ in the hospital environment. The main difference in vSaβ islands in strains from the United States and United Kingdom is a polymorphism that breaks the seu superantigen gene into seu1 and seu2 pseudogenes in the CC5 strains. In contrast to a previous hypothesis (40), our results indicate that the complete seu gene is in the ancestral state, and a deletion at the base of the CC5 clade created the pseudogenes. The North American CC5 vSaβ island also includes leukocidin genes, which encode a toxin that prevents phagocytic clearance (41, 42). CC5 strains lack the phage carrying the Panton-Valentine leukocidin (PVL) toxin (43) gene. Synteny analysis independently confirmed each of the above differences and identified other changes in gene position of unknown consequence (see Fig. S3 in the supplemental material).

Lipoproteins unique to CC5. S. aureus strains are known to harbor clusters of lipoprotein genes typically at four locations in the chromosome—within a vSaα element and at three other sites

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FIG 4 Restriction systems and dprA in VRSA. (A) (1) Alignment of inferred primary translation products of genes encoding the SauI restriction nuclease subunit (HsdR) compared to the CCS strain originating in Japan, *S. aureus* N315. Polymorphisms are highlighted. Periods in blocks of sequence denote stretches that are identical and not shown. Numbers above the sequences represent the amino acid position in the prototype, from which distances represented by dots can be discerned. (2) Alignment of inferred primary translation products of genes encoding the modification (HsdM) and specificity (HsdS) subunits in the /H9263/Sa/H9251/i island. The locations of restriction-modification subunits relative to other key genes are indicated. (3) Alignment of inferred primary translation products of genes encoding the specificity (HsdS) subunits in the /H9263/Sa/H9252/i island. No polymorphisms occur in the /H9263/Sa/H9252/i HsdM-encoding gene (not shown). (4) Alignment of inferred primary translation products of genes encoding the type III-like restriction system compared to those of N315 and the CC8 *S. aureus* NCTC8325 prototype sequence. (B) Alignment of inferred primary translation products of dprA genes of VRSA. VRS3a is the only isolate that encodes a dprA product identical to that encoded by the Japanese CCS isolate N315. All other VRSA strains possess a truncating frameshift mutation predicted to generate a premature termination, and possibly overlapping reinitiation product, as generically designated in the figure as VRSAa and VRSAb.
CC5 genomes possess a significantly larger set of lipoprotein genes than non-CC5 genomes (P < 0.05 by the Mann-Whitney U test; see Fig. S4 in the supplemental material), suggesting that selection favors their occurrence in these hospital isolates. We observed strong congruence between lipoprotein- and whole-genome-based phylogenies, showing that lipoprotein variation largely parallels chromosome divergence patterns. Interestingly, BLAST identified Listeria grayii as the only non-staphylococcal species possessing highly related homologs of S. aureus lipoproteins (<1e-24). Gram-positive bacteria process lipoprotein signal peptides into septa- or octapeptide pheromones (45), which have been hypothesized to contribute acquisition of vancomycin resistance by staphylococci (46). Potential pheromone sequences encoded by the North American CC5 S. aureus genomes were identified (Fig. S4).

DISCUSSION

MRSA emerged in the early 1960s and remained largely restricted to the hospital environment (43). Not until resistance occurred in other lineages 30 years later did it rapidly spread in the community (43). CC5 is often the first lineage in which new antibiotic resistance genes appear (10, 11, 47). Our work provides quantitative support for an earlier proposition, based largely on PFGE (15–17), that each occurrence of Tn1546-conferred vancomycin resistance in S. aureus represents an independent acquisition, rather than patient-to-patient spread. Modeling evolutionary distances on a time scale (Fig. 6) shows that the last common ancestor of all VRSA strains occurred at about the time of methicillin introduction, about 1960, 40 years earlier than vancomycin resistance was found to have entered the species. The early 1960s also is the approximate time when the Japanese CC5 isolates diverged. The modeling analysis employed was based on a calculated mutation rate of 3.46 × 10^-6 per site per year and employed conservative, relaxed-clock assumptions. This rate is in excellent agreement with the 3.3 × 10^-6 mutation rate calculated independently by others in a rigorous application of second-generation sequencing in a study of the global spread of the MRSA lineage ST239 that is prominent in Asia (48). Only in the case of strains VRS11a and VRS11b, which were isolated from the same patient, does the uncertainty with respect to the time of strain divergence extend beyond the date of isolation of that VRSA strain. The 1960 time point represents the time when the most divergent VRSA strain, VRS3b (PFGE type USA800), branched from the rest, most being USA100 PFGE type (Fig. 5). Within the USA100 group, extensive diversification occurred next in about 1978.

In contrast to expectations, VRSA restriction barriers appear to be largely intact. Strain VRS3a possesses a defect in the Sau1endonuclease that is likely to be of functional consequence. However, most other polymorphisms in the Sau1 system were limited to the Sa-encoded copy of a modification gene, with the Sa-encoded copy fully intact. Another strain, VRS10, possesses change in the type III-like restriction, shortening the predicted primary translation product from 953 amino acids to 856 amino acids by removal of the amino terminus. The functional consequence of this truncation is currently unknown. It may be important that all VRSA strains, except for the phylogenetic outlier strain VRS3a, possess a nonsense mutation early in the dprA gene.

FIG 5 Variation in the Saβ island of VRSA strains compared to other lineages. The gray shading for the schematic representations of the Saβ islands shown to the right of the figure corresponds to the position within the phylogeny showing the relationship of VRSA genomes to other completely sequenced S. aureus genomes shown to the left of the figure. The asterisk in the top right-hand corner of the schematic representation in panel b indicates that S. aureus COL contains an IS1811 transposase upstream of bsaA2. The pair of asterisks in the top right-hand corner of the schematic representation in panel d indicate that the far right endpoint for RF122 is not shown because of phage insertion.
DprA (also known as Smf [49]), is highly conserved and contributes to efficient DNA transformation in naturally competent bacteria (29, 49–54). Transformation efficiency of plasmids in a *B. subtilis* *dprA* mutant is decreased 60-fold (29). Experiments with *Escherichia coli dprA* mutants do not show an obvious role in transformation or conjugation (55). Its function in *S. aureus* remains to be explored.

The most variable feature of the VRSA genome is plasmid content. In all cases, Tn1546 resides on a plasmid, even though it clearly transposed upon entry into some strains, and because of size, the chromosome would seem to be the most probable target for transposon insertion. The basis for the insertion site preference for plasmids over the *S. aureus* chromosome, and also for an apparent incompatibility between the enterococcal Inc18 plasmid that played a major role in the Michigan outbreak and an endogenous *S. aureus* pSK41 plasmid present in several recipients, is unknown. VRSA genomes are replete with plasmids of enterococcal origin, highlighting their cooccurrence in polymicrobial infections and possibly in other ecologies. The multiplicity of plasmid structures conveying Tn1546, including *S. aureus*/enterococcal cointegrate plasmids, increases the odds of future transfers, possibly into staphylococcal lineages or species where a lower fitness cost is incurred.

The genomic island vSaβ is a distinguishing feature of CC5. Of potential ecological importance, the bsa operon usually encoded within this island is absent in VRSA and other CC5 strains. Frequent application of antibiotics in the hospital environment may select for strains with an enhanced ability to comingle with potential resistance donors of other species. Interestingly, this trait is also lacking in otherwise highly divergent strains that are prevalent in hospitals in the United Kingdom. The fact that United Kingdom and U.S. hospital strains with widely different chromosomal backgrounds share very similar vSaβ islands suggests that there may be active selection for this configuration in the hospital environment. Multidrug-resistant enterococci are much more likely to lack clustered regularly interspaced short palindromic repeat (CRISPR) defenses of the genome than commensal strains (56), indicating that the widespread use of antibiotics has selected for hospital-adapted bacteria that have enhanced abilities to exist in mixed communities and exchange resistance determinants. Loss of bacteriocin production as well as immunity may also explain why, for 30 years, CC5 MRSA strains were not able to establish methicillin resistance in the community at a high level. They may have been inhibited by the functional bacteriocin loci of *S. aureus* strains of other sequence types (such as CC8 and CC1) already present in the community niche. This may also be limiting the spread of vancomycin resistance from CC5 strains to other clades.

Staphylococcal enterotoxins, T cell mitogen superantigens (57) that dysregulate the host response by stimulating CD8+ regulatory T cells (Treg) at low concentrations (58) and other mechanisms, are particularly abundant in the CC5 vSaβ element, as well as that of the

**FIG 6** Consensus tree for CC5 under relaxed clock conditions. North American CC5 strains are shown in black, while other strains are shown in gray. Blue bars indicate range of 95% highest posterior density interval (95% confidence interval, 1.76 × 10^-6 to 5.22 × 10^-6). The time of penicillin and methicillin introduction, as well as the emergence of resistances are shown for reference (PRSA, penicillin-resistant *S. aureus*). The arrow indicates the estimated time of divergence of the most distantly related VRSA genome, that of VRS3a (USA800 branch) from the remaining VRSA (mainly USA100) branch of the CC5 clade, showing that strain divergence occurred far earlier than vancomycin resistance acquisition, supporting the model of independent Tn1546 acquisition.
United Kingdom clones that are prevalent in hospitals. Enterotoxins, together with lipopolysaccharides (59) and leukocidin (41, 42) may facilitate overgrowth of mixed populations of bacteria at a site of infection, contributing to the creation of conditions favorable for resistance transfer. Notably, most VRSA strains occurred in mixed infections of plantar ulcers of diabetic patients (2), infections that are known to be highly polymicrobial (60, 61).

The hospital is a unique environment, where colonization and patient-to-patient propagation of a strain may depend less on bacterial virulence traits associated with transmission (62) than transmission vectors in the form of hospital staff and environmental surfaces and antibiotics (63). Strains that are prevalent in hospitals are under continuous antibiotic selection pressure and are exposed to an ever rotating arsenal. CC5 isolates appear to be very well adapted for surviving and evolving in this environment by acquiring resistance to new antibiotics.

MATERIALS AND METHODS

Strains. All available VRSA strains were obtained through the Network for Antimicrobial Resistance in Hospitals (NARSA) (see Table S1B in the supplemental material). Since strain VRS3a is believed to be identical to strain VR3b, the latter was not sequenced. Strains VRS11b and VRS11a had not been characterized, and both were examined because of differences in oxacillin resistance. The VRSA strains were routinely grown on tryptic soy agar containing 10 μg/ml vancomycin.

Sequence analysis. The complete genome of AHBV00000000. The complete genomes of VRS10, AHBT00000000; VRS11a, AHBU00000000; and VRS11b, VRS7, AHBQ00000000; VRS8, AHBR00000000; VRS9, AHBS00000000; VRS4, AHBN00000000; VRS5, AHBO00000000; VRS6, AHBP00000000; VRS10, AHBQ00000000; VRS11a, AHBQ00000000; and VRS11b, AHBQ00000000. The complete genomes of S. aureus strains used for comparison (see Table S1C in the supplemental material) were downloaded from GenBank (http://www.ncbi.nlm.nih.gov) and the Sanger website (http://www.sanger.ac.uk/pathogens).

FOR Illumina sequencing, total DNA was purified from 10-ml overnight cultures using the DNeasy DNA extraction kit (Qiagen). DNA was transferred to the Tufts University DNA Core Facility, Boston, MA, and a modified Illumina protocol (64) was used. Libraries were subjected to multiplexed paired-end sequencing according to the manufacturer’s specifications. Sequencing reads were filtered to exclude reads with a quality score of <25 at any position. The average coverage of the 3-Mb genomes was >110-fold. The genomes were also independently sequenced at the University at Buffalo Next-Generation Sequencing and Expression Analysis Core (Buffalo, NY), by 454 FLX (Roche) to at least 10-fold coverage.

Assemble. Illumina reads were assembled using Velvet version 1.0.18 (65). The 454 and Illumina reads were then combined and assembled using Newbler 2.3 (Roche). Gene annotations were generated using the Prodigal gene caller (66). Draft genomes for strains were submitted to GenBank under the following accession numbers: strain VRS1, AHBK00000000; VRS2, AHBL00000000; VRS3a, AHBM00000000; VRS4, AHBN00000000; VRS5, AHBS00000000; VRS6, AHBP00000000; VRS7, AHBQ00000000; VRS8, AHBR00000000; VRS9, AHBV00000000; VRS10, AHBW00000000; VRS11a, AHBX00000000; and VRS11b, AHBV00000000. The complete genomes of S. aureus were used for comparison (see Table S1C in the supplemental material) were downloaded from GenBank (http://www.ncbi.nlm.nih.gov) and the Sanger website (http://www.sanger.ac.uk/pathogens).

PCR and targeted DNA sequencing. To verify results in some cases or to obtain missing sequence information, PCR was performed using Taq polymerase (New England Biolabs) with the amplification primers listed in Table S4 in the supplemental material. DNA sequencing of individual PCR products was performed by the Massachusetts General Hospital DNA Sequencing Core Facility. Specific nucleotide sequence alignments were routinely performed using mafft (67) and ClustalW (68).

Bioinformatic analysis. See Text S1 in the supplemental material.

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at http://mbio.asm.org/lookup/suppl;doi:10.1128/mBio.00112-12/-/DCSupplemental.

Text S1, DOC file, 0.2 MB.

Figure S4, PDF file, 1.5 MB.

Table S1, DOC file, 0.1 MB.

Table S2, DOCX file, 0.1 MB.

Table S3, DOC file, 0.1 MB.

Table S4, DOC file, 0.1 MB.

Table S5, DOC file, 0.1 MB.

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