vanG Element Insertions within a Conserved Chromosomal Site Conferring Vancomycin Resistance to *Streptococcus agalactiae* and *Streptococcus anginosus*

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**ABSTRACT** Three vancomycin-resistant streptococcal strains carrying vanG elements (two invasive *Streptococcus agalactiae* isolates [GBS-NY and GBS-NM, both serotype II and multilocus sequence type 22] and one *Streptococcus anginosus* [Sa]) were examined. The 45,585-bp elements found within Sa and GBS-NY were nearly identical (together designated vanG-1) and shared near-identity over an ~15-kb overlap with a previously described vanG element from *Enterococcus faecalis*. Unexpectedly, vanG-1 shared much less homology with the 49,321-bp vanG-2 element from GBS-NM, with widely different levels (50% to 99%) of sequence identity shared among 44 related open reading frames. Immediately adjacent to both vanG-1 and vanG-2 were 44,670-bp and 44,680-bp integrative conjugative element (ICE)-like sequences, designated ICE-r, that were nearly identical in the two group B streptococcal (GBS) strains. The dual vanG and ICE-r elements from both GBS strains were inserted at the same position, between bases 1328 and 1329, within the identical RNA methyltransferase (*rumA*) genes. A GenBank search revealed that although most GBS strains contained insertions within this specific site, only sequence type 22 (ST22) GBS strains contained highly related ICE-r derivatives. The vanG-1 element in Sa was also inserted within this position corresponding to its *rumA* homolog adjacent to an ICE-r derivative. vanG-1 insertions were previously reported within the same relative position in the *E. faecalis rumA* homolog. An ICE-r sequence perfectly conserved with respect to its counterpart in GBS-NY was apparent within the same site of the *rumA* homolog of a *Streptococcus dysgalactiae* subsp. *equisimilis* strain. Additionally, homologous vanG-like elements within the conserved *rumA* target site were evident in *Roseburia intestinalis*.

**IMPORTANCE** These three streptococcal strains represent the first known vancomycin-resistant strains of their species. The collective observations made from these strains reveal a specific hot spot for insertional elements that is conserved between streptococci and different Gram-positive species. The two GBS strains potentially represent a GBS lineage that is predisposed to insertion of vanG elements.
vancomycin resistance in strains of *Enterococcus faecalis* through the synthesis of peptidoglycan precursors with C-terminal D-Ala–D-serine (D-Ala–D-Ser) residues that have low vancomycin affinity while concurrently directing the removal of precursors ending with D-Ala–D-Ala. The *vanG* resistance proteins include the *vanG*-encoded D-Ala–D-Ser ligase, the *vanT*-encoded serine racemase, the *vanXY*-encoded bifunctional D,D-carboxypeptidase/D,D-peptidase, and the *vanY*-encoded D,D-carboxypeptidase which in some elements may be inactive due to a frameshift mutation (5). Detailed transcriptional analysis of a *vanG* operon in *E. faecalis* and assessment of *vanG* product enzymatic activities have been previously described (5). Although *vanG*-like elements with relatively divergent sequences in comparison to *E. faecalis* *vanG* elements are found within *Clostridium* and *Ruminococcus* (9, 10), functional *vanG* elements shown to confer vancomycin resistance have been characterized only from a small number of *E. faecalis* strains recovered in Australia and Canada (4–7). Here we provide a preliminary description of the first known isolates of *Streptococcus agalactiae* and *Streptococcus anginosus* that express vancomycin resistance. These three strains are also the first known members of the genus *Streptococcus* that carry elements of the *vanG* type.

**RESULTS**

**Transferability of *vanG-1* and *vanG-2*.** Many attempts were made to detect transfers of these elements from the three streptococcal strains; however, all were unsuccessful. These included interspecies transfers (GBS-NY and GBS-NM donors to *E. faecalis* recipients; *S. anginosus* urine isolate [Sa] donor to *E. faecalis* and GBS recipients). Intraspecies transfer events employing GBS-NM and a diverse set of erythromycin-resistant GBS strains were also attempted. Since strains GBS-NY and GBS-NM are both serotype II and genotype sequence type 22 (ST22), it appeared possible that strain GBS6 (also serotype II and ST22) would be an ideal recipient strain for *vanG* element transfer events; however, no vancomycin-resistant recipients were detected.

**Insertional inactivation of *vanG*.** Inactivation of the *vanG* gene in GBS-NY was facilitated by transforming the strain to chloramphenicol resistance with plasmid pVE6007-G*’* via a single-crossover event. This conferred a vancomycin-susceptible (MIC, <1.0 μg/ml) phenotype. The vancomycin-sensitive phenotype was reversible at a low frequency, and vancomycin-resistant colonies were detected following overnight growth at 37°C in Todd-Hewitt broth (THB) without drug and plating on TH agar containing 2 μg/ml vancomycin. Insertional inactivation and homologous excision events were verified through PCR employing appropriate primers annealing to *vanG* and pVE6007-G*’* sequences.

**Resistance phenotypes.** GBS-NY, GBS-NM, and *S. anginosus* strain Sa exhibited a vancomycin MIC of 4.0 μg/ml and were tetracycline resistant. Despite the observed relatedness between GBS-NY and GBS-NM (both multilocus sequence type 22, serotype II), only GBS-NY was resistant to erythromycin and clindamycin. As previously described for vanG-positive *E. faecalis* (5), all three streptococcal strains displayed inducible vancomycin resistance in that growth in the presence of 2 μg/ml vancomycin was accelerated by preincubation in the presence of a subinhibitory (0.4 μg/ml) concentration of the antibiotic (Fig. 1). As previously described in a report of an *E. faecalis* study (4), these vanG-positive strains were susceptible to the glycopeptide teicoplanin (data not shown). Growth in the presence of 2 μg/ml vancomycin was detected in GBS-NM within 2 h; however, preincubation with 0.4 μg/ml shortened this time period to 1.5 h. In contrast, detectable growth of both GBS-NY and Sa was observed only under noninducing conditions after 8 and 10 h, respectively, and these time periods were reduced under inducing conditions to 6 and 8 h, respectively. *E* test results revealed that strains GBS-NY and Sa had a vancomycin MIC of 3.5 μg/ml, while GBS-NM had a slightly higher MIC of 4.0 μg/ml. The broth dilution vancomycin MIC for all 3 strains was 4.0 μg/ml.

**Growth rate and β-lactam susceptibility comparisons of GBS-NY with a *vanG*-inactivated derivative.** To determine if there is an easily detectable growth disadvantage conferred by the *vanG*-1 element, growth rates of GBS-NY were compared to those of its derivative, the pVE6007-G’ insertion strain, and to the plasmid-excised revertant of the latter strain in nonselective Todd-Hewitt broth supplemented with 1% yeast extract (THYB). We also retested the antimicrobial profiles for these three strains, primarily to determine if *vanG* conferred hypersensitivity to β-lactam antibiotics as previously described for vancomycin-resistant *Staphylococcus aureus* laboratory mutants (11). Growth and MIC comparison experiments were performed in parallel in the presence and absence of a subinhibitory vancomycin concentration (0.04 μg/ml). In brief, no significant differences in growth rates or in MICs of any antibiotics were observed. These included susceptibilities to the β-lactam antibiotics ampicillin (range of 0.12 to 0.25 μg/ml), cefazolin (0.25 μg/ml), cefoxitin (range of 4 to 8 μg/ml), penicillin (range of 0.06 to 0.12 μg/ml), and cefotaxime (range of 0.06 to 0.12 μg/ml); those values, while not indicative of basal sensitivities, are not unusual in our population-based surveillance (data not shown). The lowest MICs possible that were detectable in these assays were ≤0.004 μg/ml (ampicillin, penicillin), ≤0.004 μg/ml (cefotaxime), ≤0.016 μg/ml (cefoxitin), and ≤0.015 μg/ml (cefoxolin).

**Features of *vanG* operons from *vanG-1* and *vanG-2*.** The 7,927-bp 8-gene *vanG* operon (Fig. 2) from GBS-NY and Sa (orf29 to orf36) shared complete sequence identity (see Table S1 in the supplemental material) with the previously described operon from *Enterococcus faecalis* strain G1-0247 (7), consisting of a previously mapped constitutively transcribed sigma70-type promoter (5) 29 bp upstream of a 3-gene (vanU, vanR, and vanS) transcriptional regulatory region and a 5-gene (vanY, vanW, vanG, vanXY, and vanT) resistance region previously found to be under the control of a vancomycin-inducible promoter (5). The sequence also shared near-identity with the corresponding sequence from *E. faecalis* BM4518; however, BM4518 contained a T frameshift insertion within *vanY* (5).

GBS-NY and GBS-NM contained *vanG* elements (designated *vanG-1* and *vanG-2*, respectively) that differed considerably in size (45,585 bp versus 49,319 bp) and in sequence (Fig. 2; see also Table S1 in the supplemental material). The 7,667-bp *vanG* operon within *vanG-2* differed substantially from the corresponding 7,927-bp *vanG-1* sequence from GBS-NY, especially within the regulatory gene region (see Table S1 [compare orf29 to orf31 of *vanG-1* with orf29 to orf31 of *vanG-2*]). From *vanG-2*, the deduced first 10 VanUR amino acids shared identity to the predicted *vanG-1* VanU N terminus, followed by 10 unrelated amino acids fused to a 236-residue sequence with 61% identity to the corresponding 236 residues of the GBS-NY/Sa VanK protein (66.8% DNA sequence identity shown in Table S1). The putative...
sigma70-directed promoter sequence, 34 bp upstream of vanUR, shared 72.4% identity with the corresponding vanG-1 promoter sequence, with a −10 region perfectly matching the consensus prokaryotic sigma70-directed promoter (Fig. 2). Immediately upstream of the GBS-NM vanY gene was a sequence with homology (20 of 29 matching positions) to the putative vancomycin-inducible promoter in GBS-NY and Sa. These results indicate that it is likely that vanG operon products from these streptococcal strains are transcriptionally regulated in the same manner as previously described in E. faecalis (5), where a constitutive promoter lies upstream of the vanU, vanR, and vanS (vanUR and vanS in GBS-NM) transcriptional regulatory genes and a vancomycin-inducible promoter upstream of vanY further directs transcription of the resistance enzyme genes.

The 5 resistance genes from GBS-NM shared progressively increased homology with their GBS-NY/Sa counterparts toward the 3’ end, with the two downstream genes vanXY and vanT sharing 95.7% and 99.5% sequence identity, respectively. The vanG genes shared 91% identity, with both containing codons at positions (D243, F252, and R324) postulated to be molecular determinants of terminal D-serine selectivity (12).

**Additional features of vanG-1 and vanG-2.** Different sections of vanG-1 and vanG-2 displayed markedly differing GC/AT ratios, suggesting at least two different origins for components of these elements. Bases 1 to 33,205 of vanG-1 revealed a G+C content of 47.8%, while the vanG operon itself (bases 33,382 to 41,308 in Fig. 2) revealed a markedly lower G+C content of 37.2%. Similarly, bases 41,308 to 45,585 had a G+C composition of 39.9%. A similar scenario was observed for vanG-2. While the vanG operon (bases 37,378 to 45,044) had a G+C content of 35.4% and bases 45,044 to 49,321 revealed a G+C content of 40.2%, the left section (bases 1 to 37,338) had a G+C content of 45.2%.

A detailed analysis of the orfs within vanG-1 and vanG-2 is beyond the scope of this report; however, we do highlight certain features here. The complete 45,585-bp sequences from GBS-NY and Sa differed in only 2 positions, specifically, a missense substitution within orf8 (position 7913) with no predicted function and a synonymous substitution within orf38 (position 41,951 [Fig. 2]). Although all three strains were tetracycline resistant and GBS-NY was resistant to erythromycin and clindamycin, neither vanG element contained genes encoding resistance to antibiotics other than vancomycin.
There were 47 orfs in vanG-1, ranging from 183 bp to 8,076 bp in length (Fig. 2; see also Table S1 in the supplemental material). Forty-four of the 47 orfs, including the vanG operon genes and all orfs that were >360 bp in length, were in the same orientation. Matches with >90% sequence identity to any of these orfs were not found within complete or incomplete S. agalactiae or S. anginosus genomes in NCBI databases. The respective vanG operons in vanG-1 and vanG-2 are indicated as bases 33,382 to 41,308 and bases 37,378 to 45,044, respectively, each starting at the first base of the putative constitutive promoter and ending at the vanT stop codons. The N-terminal 10 residues deduced from orf29 and orf30 (vanUR) from vanG-2 are identical to vanU of vanG-1; however, the major portion of orf29 and orf30 shares homology with vanG-1 orf30 (vanR). Arrows pointing right to left indicate orfs in the orientation opposite that of the other orfs in the diagram.

The best overall matches to the entire vanG-1 and vanG-2 elements from the NCBI Nucleotide collection (nr/nt) database were from the genomes of Roseburia intestinalis strains XB6B4 and M50/1, respectively (see Fig. S1 and S2 in the supplemental material). Strain XB6B4 shared 34 orfs with vanG-1, with levels of sequence identity ranging from 60.5% to 99.4% (see Table S1 in the supplemental material). Differences from the consensus bacterial sigma70 promoter are indicated in lowercase characters. The single –10 region difference in the constitutive sigma70 class promoter in vanG-2 is underlined. In the same fashion, the putative vancomycin-inducible promoter sequences (5) are indicated upstream of resistance genes vanY, vanW, vanG, vanXY, and vanT. Arrows pointing right to left indicate orfs in the orientation opposite that of the other orfs in the diagram.
mology, with 35 orfs sharing 80% to 99.8% DNA sequence identity (see Fig. S2 and Table S1). As with the comparison of vanG-1 with the corresponding sequence from R. intestinalis XB6B4, the most obvious difference between the vanG-1 element and the corresponding sequence from R. intestinalis strain M50/1 was the lack of vanG operon counterparts, and there was no recognizable vanY-like orf in strain M50/1.

ICE-r and vanG elements in GBS are inserted within rumA codon 443. On the basis of comparisons to serotype II, ST22 control strain GBS6, the two heterologous vanG elements were inserted at the same relative positions after base 1328, within codon 443, of their identical putative RNA methyltransferase (rumA) genes (Fig. 3A, C, and D). Immediately downstream of both of the GBS vanG elements and immediately fused to the corresponding identical GBS6 rumA5 = rumA3 segments (see panels B to D) were on one side only of the ICE-r derivative shown in panel E). The 8-bp and 3-bp sequences that are shared between vanG and ICE-r element termini are indicated in green. (C and D) The tandem vanG and ICE-r insertions in strains GBS-NY and GBS-NM with the vanG element imperfect inverted repeats (ILR2 and IRR2) and ICE-r inverted repeats (ILR1 and IRR2). Base substitutions in the left repeat (LR) and right repeat (RR) relative to vanG-1 are shown in red (D). (E) The vanG-1 insertion within the S. anginosus rumA gene is followed by an ICE-r derivative that shares high intermittent homology with ICE-r from GBS and S. equisimilis ATCC 12394 (see panels B, C, D, and F). Although the rumA 5’ segment was identical to its counterpart in S. anginosus F0211 (see panel A), a corresponding ICE-r IRR1 and rumA 3’ segment was absent. Base changes in the unmatched ICE-r derivative inverted left repeat relative to ICE-r (panels B to D) are shown in red. (F) ICE-r element nearly perfectly conserved with GBS-NY inserted within S. dysgalactiae subsp. equisimilis rumA gene. (G) Diagram of previously published information (5, 7) describing vanG insertions in two different E. faecalis strains. The slanted vertical lines for the E. faecalis vanG elements indicate that the lengths are presently unknown. (H) The vanG-like insertion element in R. intestinalis XB6B3, which has high homology with vanG-1 (see Fig. S1 in the supplemental material).
tative codon 549 serine recombinase gene situated at the right end of the element that shared 52.9% sequence identity over a 537-residue overlap of the \( \text{vanG-1} \) and \( \text{vanG-2} \) 547 codon orf44 (putative serine recombinase gene) situated at the right ends (data not shown) (note that the orf44s from the two \( \text{vanG} \) elements share sequence identity [see Table S1 in the supplemental material]). Although the inverted imperfect repeats of the \( \text{vanG-1} \) and \( \text{vanG-2} \) elements shared little similarity with those of the ICE-\( \text{r} \) elements (Fig. 3C and D), the first 8 bases and last 3 bases of each \( \text{vanG} \) element in this region were identical with those of the other Gram-positive elements shown (Fig. 3 [bases in green]). Other than a 10-base deletion within a noncoding sequence in GBS-NM, the two GBS 44,670-bp and 44,680-bp ICE-\( \text{r} \) elements shared complete sequence identity. In all three GBS strains, the ICE-\( \text{r} \) element was followed by the identical 11 \( \text{rumA} \) 3’ codon segment situated 7,525 bp upstream of the \( \text{cylX} \) gene.

The \( \text{S. anginosus} \text{vanG-1} \) and ICE-\( \text{r} \)-like element is inserted within codon 443 of its \( \text{rumA} \) homolog. We also found that the \( \text{vanG-1} \) element within \( \text{S. anginosus} \) strain Sa was inserted within a \( \text{rumA} \) gene completely identical over its 5’ 1,328-bp sequence with the corresponding sequence from \( \text{S. anginosus} \) (e.g., strain F0211) in the NCBI database (Fig. 3A and E). Immediately following the \( \text{vanG-1} \) direct right repeat (GA), an ICE-\( \text{r} \) derivative was evident with approximately 80% sequence identity to the GBS \( \text{ICE-r} \) ; however, this homology was intermittent along approximately 57 kb of the \( \text{Sa} \) ICE-\( \text{r} \) derivative. An inverted right repeat region (IRR1) similar to that found in GBS-NY and GBS-NM was not evident in strain Sa. The full-length \( \text{S. anginosus} \) F0211 \( \text{rumA} \) gene was the typical 1,362 bases found in other streptococcal species; however, we were unable to deduce a corresponding \( \text{rumA} \) 3’ 11-codon sequence from strain Sa (Fig. 3A and E).

Previously reported \( \text{vanG} \) insertions within the \( \text{E. faecalis rumA} \) gene. The same relative insertion sites of \( \text{vanG} \) elements within a more heterologous \( \text{rumA} \) gene were apparent for \( \text{E. faecalis} \) strains G10247 and BM4518, with the separated 5’ and 3’ segments exactly matching those of the uninterrupted wild-type \( \text{E. faecalis rumA} \) gene (5, 7; Fig. 3A and G). These two partially sequenced \( \text{vanG} \) elements (GenBank accession numbers DQ212987 and DQ212986 and accession numbers AY271781 and AY271782) from \( \text{E. faecalis} \) shared near-identity in their overlaps with \( \text{vanG-1} \) from GBS-NY and Sa.

\( \text{vanG-like} \) and ICE-\( \text{r} \)/ICE-\( \text{r} \)-like insertions within the conserved \( \text{rumA} \) location from additional Gram-positive species. \( \text{vanG-like} \) elements were observed in \( \text{R. intestinalis} \) genomes (see Fig. S1 and S2 in the supplemental material), and an ICE-\( \text{r} \)-element nearly identical to that found within ST22 GBS was observed in \( \text{S. dysgalactiae subspp. equisimilis} \). In all 5 species examples shown, these elements are inserted after the same dinucleotide target within the 3’ proximal 11th codon (GAG) (Fig. 3).

It was interesting that, other than the \( \text{S. agalactiae ST22} \) strains, the only nearly perfectly conserved ICE-\( \text{r} \) (44,672-bp to 44,680-bp identity with GBS-NY ICE-\( \text{r} \)) was found inserted within codon 443 of the \( \text{rumA} \) gene in \( \text{S. equisimilis} \) strain ATCC 12394 (Fig. 3F). Large regions of identity to ICE-\( \text{r} \) (84% to 91% coverage, with 96% to 99% sequence identity) were also seen within \( \text{rumA} \) codon 443 of \( \text{S. equisimilis} \) strains RE378 and AC-2713, within total insertion sizes of about 60.5 kb and 98.4 kb, respectively (data not shown). A highly conserved element also observed in \( \text{Streptococcus pyogenes} \) is the ICESp2905 element (89% identity, 60% coverage [GenBank accession number FR691055]) inserted within its chromosomal \( \text{rumA} \) 3’ site (13). Within \( \text{Streptococcus equi} \) subspp. \( \text{equi} \) strain 4047, a homologous ICE-\( \text{r} \)-like derivative was also observed to be inserted within codon 443 of its \( \text{rumA} \) gene (GenBank accession number FM204883), with approximately 90% identity with ICE-\( \text{r} \) over a 50% overlap.

Frequent insertions within GBS \( \text{rumA} \) and conservation of ICE-\( \text{r} \)-within ST22. The two sections of the interrupted \( \text{rumA} \) gene of GBS-NY, GBS-NM, and GBS6 (5’ bases 1 to 1328 and 3’ bases 1329 to 1362) exactly matched full-length 1,362-bp GBS \( \text{rumA} \) genes from 31 strains in the wgs (whole-genome shotgun contig) NCBI database (e.g., strain MC623). In the majority (>230) of GBS genomes, there were insertions present within this exact site. All 10 GBS whole-genome sequences within the nt/nr NCBI database contained insertions within this \( \text{rumA} \) site (after base 1328 within codon 443). In the majority of all GBS strains, and in all 31 genomes in the NCBI database with an intact \( \text{rumA} \), the 3’ 11-codon segment of \( \text{rumA} \) was situated approximately 7,525 bp upstream of the \( \text{cyl operon} \) (data not shown) found in all hemolytic GBS strains (14, 15) (depicted in Fig. 3A to D).

Of the total of 279 \( \text{S. agalactiae} \) whole-genome sequences within the NCBI databases, only 12 ST22 isolates and one single-locus variant (SLV) of ST22 contained large sections of sequence identity to the GBS-NY/GBS-NM ICE-\( \text{r} \)-insertions (Fig. 4), and in each isolate these sections were contained between bases 1328 and 1329 of the \( \text{rumA} \) gene (data not shown). None of the ST22 isolates contained a single ICE-\( \text{r} \) with 99% to 100% sequence identity to the ICE-\( \text{r} \) in GBS-NY, while 3 ST22 isolates (including GBS6) contained larger ICE-\( \text{r} \)-derivatives containing 2 to 3 segments that together encompassed the GBS-NY ICE-\( \text{r} \) with complete sequence identity. The complete GBS6 57,583-bp ICE-\( \text{r} \), also inserted between bases 1328 and 1329 of its \( \text{rumA} \) gene, contained an internal nonhomologous 12,900-bp segment (bases 23233 to 36132) that shared extensive homology (~95% identity over 5,700 bp) with mobilization genes from an \( \text{Anaerococcus prevoti} \) plasmid (GenBank accession number CP001709). In one ST22 strain, there were 3 interrupted sequences of near-identity to the GBS-NY ICE-\( \text{r} \) encompassed within an approximately 80-kb fragment found within the \( \text{rumA} \) target site. One SLV (MC633 in Fig. 4) contained an ICE-\( \text{r} \) derivative within \( \text{rumA} \) with 7 sections that together exactly encompassed perfect matches to 87% of the GBS-NY ICE-\( \text{r} \) (data not shown). In all three of the ST22 GBS study strains, and in all 10 ST22 genomes present in the NCBI wgs database, the GA at the start of ICE-\( \text{r} \) was followed by a left inverted repeat sequence that had a corresponding imperfect right inverted repeat at the ICE-\( \text{r} \) right end and a matching “far right” GA repeat (Fig. 3B, C, and D). Lower levels of identity (83% to 94% identity) were evident in additional strains, primarily in SLVs of ST22 (data not shown).

Observations relative to ICE-\( \text{r} \) divergence within the conserved \( \text{rumA} \) target site in GBS are consistent with the phylogenetic clustering of ST22 core genome sequences (Fig. 4) and with the general congruence of the core phylogeny with multilocus sequence typing (MLST). The data are consistent with the notion that ICE-\( \text{r} \) was initially introduced into ST22 GBS and subsequently diverged. It is also interesting that the two GBS-NY and GBS-NM \( \text{vanG-positive} \) strains were resolved together from the remaining ST22 strains in Fig. 4A, even though they contain clearly divergent \( \text{vanG} \) elements.

Shared features of \( \text{vanG} \) and ICE-\( \text{r} \) elements. It was beyond the scope of this work to investigate the nature of the \( \text{vanG} \) and
ICE-r element gene content in depth; however, there are some obvious conserved features that are consistent with their shared target-site specificity. Both elements contain an array of genes homologous to previously identified genes involved in conjugal transposition. In particular, both elements contain a related 547-549 codon predicted serine recombinase/integrase gene that shares 52.8% identity in amino acid sequence (data not shown). Both putative recombinase genes encompass the right end of their element and lack a stop codon. The two elements also share homology at their left ends (Fig. 5). The vanG and vanG-like left ends are comprised of a predicted 115-codon segment, lacking a translational start, that is fused in frame to the chromosomal 5’ rumA gene segment. These predicted 115-amino-acid codon sequences share obvious homology with the corresponding 101 codons of ICE-r derivatives of the four streptococcal study isolates (Fig. 5A). Also depicted here is the related R. intestinalis strain M50/1 ICE-r derivative that lies immediately downstream of its vanG-like element. Although vanG and ICE-r share this common rumA gene target, and share the above-described limited homologies, their respective terminal imperfect inverted repeats are dissimilar (Fig. 3).

As a result of vanG-vanG-related and ICE-r insertions within the different Gram-positive rumA genes (Fig. 5), different translational fusions were generated. For strains GBS6 (and other ST22 GBS strains) and S. equisimilis, the rumA 5’ region is translationally fused to 101 codons encompassing the ICE-r left end (Fig. 3B and F and 5A). This was also true for at least one other streptococcal species in that the corresponding rumA 5’ region in the S. equi subsp. equi 447 strain was translated in this manner, where the 101 N-terminal rumA–ICE-r fusion derivative residues shared 89.1% sequence identity with their counterpart ICE-r elements from ST22 GBS and S. equisimilis (data not shown).

Within strains containing vanG or vanG-like elements (GBS-NY, GBS-NM, S. anginosus strain Sa, E. faecalis strains G1-0427 and BM4518, R. intestinalis XB6B3), the rumA 5’ region is translationally fused to 115 codons encompassing the left end of the respective vanG elements (Fig. 3C, D, E, G, and H). In addition, the rumA 3’ 11 codon segment in all examples shown in Fig. 3, with the exception of S. anginosus Sa, is translationally fused to 547-549 codon putative serine recombinase genes that encompass the right ends of the vanG-vanG-like and ICE-r elements (data not shown). It was interesting, but of unknown significance, to observe the sequence similarity between the vanG-vanG-related and ICE-r portions of the two different translational fusions to the rumA 5’ domains (Fig. 5A), and the obvious relatedness between the 11 residue RumA termini from various Gram-positive species and the first deduced 11 residues from the various GBS insertion elements was also interesting (Fig. 5A and B). Although the rumA 11th 3’ proximal codon is a conserved insertion target for vanG-vanG-related and ICE-r/ICE-r-related elements in the different species shown, the reason for this is not evident on the basis of the weak interspecies rumA sequence homology (Fig. 6).

Numerous GBS genomes from the NCBI database displayed a diverse array of related, often short sequences translationally fused at the 3’ rumA target site (a partial listing of shorter deduced fusions is shown in Fig. 5C), and these were all associated with the ends of a wide variety of insertional elements. Only the coding sequence for the GBS wild-type rumA 3’ end (ECVALLQRSKG)
FIG 5  (A) Deduced amino acid sequence homology between the indicated left ends of \( \text{van} \text{G} \)-\( \text{van} \text{G} \)-like elements (red arrows), ICE-\( r \) elements (green arrows), and \( \text{rum} \text{A} \) segments (black arrows) of different translational fusions (diagrammed at the top of the panel). The residues that are completely conserved among the 7-element left ends are blue. Residues conserved in 4 to 5 left elements are red. Residues conserved among 3 elements are green. (B) The RumA C-terminal 11 residues from different species that are often translationally fused to different serine recombinase genes (\( \text{srg} \)) are directly aligned, with those conserved among the various elements shown in red. (C) Deduced related ICE-like element 5\(^\prime\) residues in the \( \text{S. agalactiae} \) NCBI database found to be fused with the wild-type chromosomal \( \text{rum} \text{A} \) 5\(^\prime\) end (below the horizontal line). Residues conserved with \( \text{S. agalactiae} \) wild-type \( \text{rum} \text{A} \) 11 codons indicated at the top (uninterrupted by an insertion element) are shown in red. Isolate identifiers or GenBank accession numbers are provided. The approximate lengths of fusion elements inserted within \( \text{rum} \text{A} \) are indicated in parentheses, based on the distance of the wild-type \( \text{rum} \text{A} \) 11 codons from \( \text{rum} \text{A} \) base 1328 and the \( \text{cylX} \) gene, consistently situated about 7,525 to 7,650 bp downstream (strains COH1, 260/V, NEM316, GB201008-001, A909, 09mas01888, and GB00003), or based on the distance of the wild-type \( \text{rum} \text{A} \) 11 codons from \( \text{cylX} \) (strains ILRI005 and ILRI112). Other strains with no lengths shown were from contigs that did not include the entire insertion element.

FIG 6  Comparisons of 4 insertion element target regions from 4 Gram-positive species with the corresponding regions in \( \text{S. agalactiae} \). Percent DNA sequence identities are shown at right. Amino acid sequence identities (entire deduced RumA proteins) are shown below each alignment.
was consistently situated ~7.5 kb upstream of the cylX operon (either as an uninterrupted rumA gene or as a rumA gene interrupted by an insertion element). For many genomes, the deduced wild-type 5′ rumA 1,328-bp sequence was bisected from its 3′ 11 codons by the presence of insertion elements of various sizes (sizes of approximately 16 to 71 kb as depicted in Fig. 5C; see the fusions ending in ECVALLVKA and ECIALIQRVKS). In other genomes, the wild-type rumA 3′ sequence was not readily apparent; however, insertion sizes after rumA base 1328 were estimated on the basis of the location of the genomic cylX gene (strains ILRI005 and ILRI112). It is interesting that the left ends of these various elements and the apparent wild-type rumA gene 3′ 11 codons from various species share appreciable deduced amino acid sequence homology (Fig. 5), potentially consistent with an extensive history of insertion and excision events centering at the GBS rumA target.

**DISCUSSION**

In this report, we have described the first vancomycin-resistant strains within two different streptococcal species, *S. agalactiae* and *S. anginosus*. To our knowledge, there have been no previous validated reports of vancomycin resistance within these two species. Among more than 16,000 invasive GBS strains from Active Bacterial Core surveillance tested during 1996 to 2013, strain GBS-NM was the only isolate detected with a vancomycin MIC of 2 μg/ml or greater (unpublished data). Nonetheless, the detection of two independent vancomycin-nonsusceptible invasive GBS strains within such a limited time frame (2011 to 2012) suggests the possible emergence of vancomycin resistance within *S. agalactiae*. The implications of emergence of GBS strains with the observed level of resistance are not entirely clear. The adult patient infected with the GBS-NM strain recovered on vancomycin monotherapy (3). Vancomycin is among the agents recommended for perinatal GBS infection prevention within a very small subset of women and is rarely used (16); however, it is important that vancomycin is a common therapy for adults with mixed wound infections, which sometimes do include GBS. No growth defects were apparent in comparing the two vancomycin-resistant GBS isolates and the highly related invasive GBS68 isolate grown under laboratory conditions (data not shown) or in comparing GBS-NY and its vanG-inactivated insertion derivative. In addition, unlike the previously described β-lactam antibiotic hypersensitivity exhibited by laboratory-selected vancomycin-resistant *Staphylococcus aureus* mutants (11), the vanG-inactivated derivative of GBS-NY exhibited no increased β-lactam susceptibility. Although not conclusive, these observations are consistent with minimal or no fitness costs imposed upon streptococcal strains containing vanG elements.

Based upon homology to known vanG elements, the donors to these streptococcal strains of the vanG elements may be *E. faecalis* strains. From the surprising near-identity of the vanG-1 elements (45,583 identical bases of 45,585) of the distantly related species *S. agalactiae* (strain GBS-NY) and *S. anginosus* (strain Sa), it is logical to suspect that events of transfer into these strains from a common source occurred very recently. The mechanism of vanG element transfer remains puzzling. The presence of imperfect inverted repeats of 28 to 30 bp on the ends of these elements is consistent with previously described conjugative elements; however, we were unable to demonstrate transfer events under laboratory conditions. In addition, conjugative transposition involves the generation of a circular intermediate. Similar to previous examination of vanG-positive *E. faecalis* strains, we were also unable to detect a circular form (5) (data not shown). If restricting our observations to *S. agalactiae* at the intraspecies level, the conserved location of insertion would appear potentially consistent with homologous recombination as a possible mechanism of vanG insertion. The vanG-1 element, however, has obviously crossed multiple species barriers, since near-perfect copies of it exist in at least two different species, with the vanG operon and other segments of this element nearly perfectly conserved with previously described *E. faecalis* counterparts (5, 7). It also appears likely that vanG and ICE-r insertion events share the same mechanism, and in the vancomycin-resistant strains, ICE-r insertion was likely to have preceded vanG element insertion.

The rumA target specificity of vanG, ICE-r, and insertion elements within multiple species (5, 7), where various heterologous chromosomal rumA genes are interrupted through an insertion event which simultaneously reconstitutes in-frame translational fusions to rumA and the element-terminal serine recombinase gene, is particularly intriguing. We have not attempted here to describe the scope of what must be a very diverse array of insertion elements that are inserted within the rumA target. The structural basis of this cross-species conserved insertion site for the highly related vanG-vanG-related and ICE-r elements described here is not apparent, especially since the level of DNA sequence identity between the different rumA homologs is often quite low (Fig. 6). Potentially, the identical 8-bp and 3-bp termini shared between the GBS vanG and ICE-r elements are involved in the mysterious target site specificity of these elements.

Further analysis and comparisons of these study strains and other streptococcal genomes in the vicinity of the conserved vanG insertion site may lead to further insights. Analysis of the rumA 3′ region within *S. agalactiae* revealed a great deal of sequence variation, associated by a great diversity of introduced genetic elements that appeared to center upon this locus (Fig. 5C). The available sequence information from the various streptococcal species discussed in this paper and elsewhere (13, 17, 18), from *Roseburia intestinalis* and *E. faecalis*, and from other Gram-positive species not mentioned in this report is also consistent in the implication of this 2-base rumA target as a very active site for resistance-conferring insertion events among a broad range of species. The high occupancy of a diverse array of insertion elements within this specific target suggests a strategy for routine localized screening of these insertions in streptococcal and other species for the introduction of new resistances. Although not described in this report, the ICE-r like element within *S. anginosus* strain Sa contained genes predicted to confer resistance to tetracycline and aminopenicillins.

Potentially, both vanG and ICE-r insertion events were mediated through serine recombinases, since both contained predicted genes on their 3′ end that share significant homology with serine recombinases, and both contain 2-bp direct repeats (GA) flanking each insertion which are representative of the target site employed by the serine recombinase family. We are unable at present to test the requirement of these or any other orfs for transfer events, since we are presently unable to facilitate vanG transfer events between different strains.

Analysis of the difference in G+C content between the left and right sections of both vanG-1 and vanG-2, where the vanG operon and downstream genes display a much lower value, suggests dif-
fferent species origins for at least these sections of these elements. The role, if any, of *R. intestinalis* in the evolution of *vanG*-1 and *vanG*-2 elements is unclear. Despite the strong homology between *vanG*-1 and *vanG*-2 with corresponding *vanG*-like elements in *R. intestinalis*, it is unknown if these elements represent prior intermediates in the evolution of *vanG* elements or if they arose from functional *vanG* elements through the loss of resistance genes. The presence of these highly homologous *vanG* or *vanG*-like elements within these diverse species (*E. faecalis, R. intestina\-lis, S. agalactiae, and S. anginosus*) suggests that a very broad host range for *vanG*-conferred vancomycin resistance is conceivable.

Previous attempts to demonstrate intraspecies *vanG* element transfer either have been unsuccessful (7) or have exhibited very low transfer frequencies (5). In past successful demonstrations of intraspecies transfer in *E. faecalis*, the conserved integration site was noted for all selected recipient strains (5). The 3 different *vanG*-positive streptococcal strains clearly represent three independent acquisitions of these elements within streptococci, since two different species are represented. *E. faecalis* is likely to be the source of the *vanG*-1 element in *S. agalactiae* and *S. anginosus*, since nearly identical elements have been previously documented in *E. faecalis* (5, 7) and all three species are frequently recovered from gut flora. The divergence between the two *vanG* elements found within the two serotype II, ST22 GBS strains is also indicative of independent origins; however, the identity of the donor species of the *vanG*-2 in GBS-NM, which is not highly conserved throughout its length with other *vanG* elements, is open to speculation. In view of the overall high sequence similarity of *vanG*-2 to a *vanG*-like element in *R. intestina\-lis* strain M50/1 (see Fig. S2 in the supplemental material), as well as of the near-identity shared by the *vanXY* gene, the *vanT* gene, neighboring genes, and the putative orf44-encoded serine recombinase of both streptococcal *vanG* elements, it is likely that multiple gut species have contributed to the evolution of *vanG*-2.

Our first analysis of whole-genome sequences from these three strains has not yet shed light upon the origins of these elements and the circumstances allowing for their transfer into streptococci. The coincidence of the two different elements residing within two highly related invasive GBS strains sharing the same serotype and multilocus sequence type suggests that this lineage might have specific features conducive to receiving *vanG* and related elements. It is possible that prior acquisition of the ICE-r element within the ST22 lineage made subsequent acquisition of *vanG* elements possible through providing cellular machinery necessary for the process.

**MATERIALS AND METHODS**

**Strains.** *S. anginosus* strain 2182 (Sa) was recovered from a catheterized urine specimen on 3 June 2012 from a quadriplegic in a long-term-care facility. The patient, a young woman, was involved in a motor vehicle accident in October 2011 and was treated with vancomycin for most of the hospitalization. She was later treated preoperatively with vancomycin for insertion of a ventriculoperitoneal shunt during May 2012. In early June 2012, she presented with fever and vomiting. The patient was again treated with vancomycin, and the shunt was removed. Urine culture yielded *S. anginosus* (MRSA) (unknown dates) and may have been treated with vancomycin then as well. The identification of *S. anginosus* was confirmed using conventional biochemical identification schema, the rapid ID32 Strep identification system (bioMérieux, Inc.), and 16S rRNA gene sequencing as previously described (19). Case reports associated with the two independent invasive *S. agalactiae* isolates (designated GBS-NY and GBS-NM), recovered in two different states (New York and New Mexico) during 2011 and 2012, respectively, have been recently published (3). These two strains, GBS-NY and GBS-NM, were both serotype II, multilocus sequence type 22 (ST22), and vancomycin nonsusceptible. GBS6, an invasive vancomycin-susceptible, serotype II, ST22 control GBS strain, was recovered in New York during 2009 through CDC’s Active Bacterial Core surveillance.

**Resistance phenotypes.** Susceptibility testing of GBS-NY and GBS-NM using the broth microdilution method (20) has been previously described (3), employing an in-house panel that was provided by Trek Diagnostic Systems. Here we additionally performed E tests (bioMérieux, Inc.). Strain Sa was tested using the same methods. All 3 strains were tested for inducible vancomycin resistance by preincubation for 1 h at 37°C in Todd-Hewitt broth containing 0.2% yeast extract and containing a sub-inhibitory concentration (0.4 μg/ml) of vancomycin prior to dilution back to an optical density (OD; absorbance at 600 nm [A600]) of 0.02 in the same medium containing 2 μg/ml vancomycin. The A600 was measured every 60 min over a 13-h period.

**Streptococcus agalactiae** multilocus sequence typing. The 7-locus multilocus sequence type of GBS-NY, GBS-NM, and GBS6 was determined as described at http://pubmlst.org/sagalactiae/.

As previously described (3), initial detection of the *vanG* element in all 3 strains employed PCR with the previously described EG1 and EG2 primers (6).

**Genome sequencing.** Initially, by employing Sanger sequencing and amplification/sequencing primers derived from data corresponding to GenBank accession numbers DQ212987 and DQ212986, ~15-kb sequences comprised of the *vanG* right and left regions from Sa and GBS-NY were obtained (data not shown). Due to the relative divergence of the GBS-NM *vanG*-2 element, the original sequencing strategy employed a previously described unidirectional PCR/sequencing strategy (21), resulting in the sequence of an equivalent ~15-kb region. The left and right chromosomal junctions of the three *vanG* elements were amplified and sequenced using unidirectional PCR primers also derived from data corresponding to GenBank accession numbers DQ212987 and DQ212986, describing the left and right ends of the *vanG* element, respectively (7). Subsequently, the whole genomes from these 3 vancomycin-resistant strains, as well as GBS control strain GBS6, were sequenced employing the complementary PacBio and Illumina platforms. Sequence comparisons of the *vanG* elements, ICE-r regions, and component open reading frames employed the NCBI Blast server (http://blast.ncbi.nlm.nih.gov/blast.cgi) and Protein Homology/analogy Recognition Engine V. 2.0 (http://www.ncbi.nlm.nih.gov/~pyhe2/html/page.cgi?id=1).

**Phylogenetic comparisons of GBS core genomes.** Two different data sets were used to compare the core sequences of the *vanG*-positive GBS strains to those of other related *S. agalactiae*. The first data set was comprised of the three study GBS strains (GBS-NY, GBS-NM, and GBS6), 12 additional ST22 genomes, 4 single-locus variants (SLVs) of ST22, and a single two-locus variant. The second data set contained the 3 GBS study strains, 6 additional ST22 SLVs, eight 2- to 3-locus variants of ST22, and a single 4-locus variant. The full-length genomes of *S. agalactiae* ST22 and ST22-related variants were retrieved from the NCBI ftp site (ftp://ftp.ncbi.nlm.nih.gov/genomes/). For any assemblies that did not contain the GenBank “locus_tag” gene identifiers, *de novo* predictions were performed by Prod-igal V 2.60 (22) with the -c option. PanOCT ver 1.9 (23) was used to find orthologous genes that shared at least 40% sequence identity (determined by an all-against-all blastp search) (24) and 70% coverage across the genomes in each group. PanOCT found 942 orthologous genes in first data set and 943 orthologs in the second. Amino acid sequences from each core genome cluster were concatenated and aligned using FSA version v 1.15.8 (25) [with the -anchored and -nolearn options]. Finally, after removing poorly aligned regions with trimAL 1.2rev59 (26) (with -automated1 option), maximum-likelihood analysis was carried out by the use of
RaxML v 7.3.0 (27) and a PROTGAMMAJITTF protein substitution model. Node support was assessed using 500 bootstrap replicates.

Electrocompetent GBS-NY. GBS-NY was grown overnight in 10 ml Todd-Hewitt broth supplemented with 1% yeast extract and 20-mM glycine (THYB-glycine). The overnight culture was then diluted 1/20 in 100 ml THYB-glycine (5-mL overnight culture transferred to 100 ml THYB-glycine) and incubated at 37°C without shaking under 5% CO₂. The optical density at 600 nm (OD_600) was monitored, and at OD_600 = 0.3, the cells were harvested by centrifugation at 5,000 × g for 20 min at 10°C. The cell pellet was washed with 225 ml prechilled double-distilled water (ddH₂O), followed by a wash with 225 ml prechilled 15% glycerol and 2 successive washes with 35 ml prechilled 15% glycerol. Cells were spun at 6,000 × g for 15 min at 4°C between washes. The pellet was resuspended in 1.5 ml of 15% cold glycerol and divided into Eppendorf tubes as 70-μl aliquots. The aliquots were immediately frozen in dry ice and stored at −80°C.

Insertional inactivation of vanG-1. Campbell-type insertional inactivation of the vanG gene was carried out by transforming GBS-NY electrocompetent cells with a derivative of the plasmid temperature-sensitive pVE6007 plasmid (28), designated pVE6007-G', that contained a 630-bp vanG internal structural gene fragment downstream of the chloramphenicol resistance gene. The pVE6007-derived plasmid was constructed by using an In-Fusion HD cloning kit (Clontech PT5162-1). A linear pVE6007 was derived using PCR with the following pair of primers: pVE6007-Fwd (GATCCACTAGTTCTAGAGCGGCC) and pVE6007-Rev (ACCGTGCACCTGAGGGG). The 630-bp vanG gene internal segment was amplified from GBS-NY with the following pair of primers: VG-pVE6007-Fwd (CCCCCTCGAGGTCGACTTGCCTGTCCCACATCGTT) and VG-pVE6007-Rev (TAGAACTAGTGGATCCGGCATCAATCGTTGCAGGCA). The VG-pVE6007-Fwd and VG-pVE6007-Rev primers contain, respectively, a 5' attachment of 15 bp homologous to the 3' end of the linearized pVE6007 and a 5' attachment of 15 bp homologous to the 5' end of linearized pVE6007. A circular plasmid was obtained by homologous recombination using 100 ng of the vanG PCR product and 100 ng of linearized pVE6007 and 10 μl of an In-Fusion HD cloning reaction mixture as instructed by the manufacturer (Clontech PT5162-1). To propagate the constructed plasmid, chemically competent Escherichia coli Top-10 cells were subsequently transformed with 3 μl of the reaction mixture and plated on LB agar containing 10 μg/ml of chloramphenicol at 30°C. Transformants with the desired construct were screened by PCR and by electrophoresis analysis of extracted plasmids. GBS-NY competent cells were transformed by electroporation (single pulse of 2.5 kV, capacitance at 25 μF, and resistance at 200 Ω) with the constructed plasmid purified from E. coli. The electroporated cells were plated on prewarmed Todd-Hewitt agar supplemented with 0.5% yeast extract (THYB) and containing 10 μg/ml of chloramphenicol after 2.5 h of incubation in 300 μl of THYB-glycine without drug at 30°C. Transformants were identified at the permissive temperature for plasmid replication (30°C). Single-crossover Campbell-type chromosomal insertions were selected by shifting to the nonpermissive temperature (37°C) while maintaining chloramphenicol selection. The insertion mutants were confirmed by PCR employing appropriate primers annealing to vanG and plasmid sequences.

Filter mating. Filter mating experiments were performed as described previously (29). Strains GBS-NY, GBS-NM, and Sa were used as attempted donors. Vancomycin-susceptible, erythromycin-resistant strains of S. agalactiae serotypes I, Ia, II (including ST22 control strain GB56), III, IV, V, VI, VII, VIII, and IX and E. faecalis 200300506S (SS 1297) were used as recipients for GBS-NM and Sa; vancomycin-susceptible and erythromycin-susceptible E. faecalis ATCC 19433 and E. faecalis 2004015741 (SS 1928) were used as recipients for all three strains. Selection for transconjugants of GBS-NY, GBS-NM, and Sa was attempted on Todd-Hewitt agar containing 0.2% yeast extract, 5% sheep blood/Trypticase soy agar, and M enterococcus medium containing 2 μg/ml vancomycin and/or 5 μg/ml erythromycin (Sigma-Aldrich, St. Louis, MO).

Nucleotide sequence accession numbers. The GenBank accession numbers for the four genomes sequenced in this work are CP007570 (GBS-NY; 2,243,708 bp), CP007571 (GBS-NM; 2,214,307 bp), CP007572 (GBS6; 2,231,475 bp), and CP007573 (Sa; 2,036,353 bp).

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at http://mbio.asm.orglookup/suppl?doi=10.1128/mBio.01386-14/-/DCSupplemental.

Figure S1, PPTX file, 0.1 MB.
Figure S2, PPTX file, 0.1 MB.
Table S1, DOCX file, 0.1 MB.

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REFERENCES


