Transfer of Tn5385, a Composite, Multiresistance Chromosomal Element from *Enterococcus faecalis*

LOUIS B. RICE,¹,²* AND LENORE L. CARIAS²

Medical Service, Department of Veterans Affairs Medical Center,¹ and Department of Medicine, Case Western Reserve University School of Medicine,² Cleveland, Ohio

Received 8 August 1997/Accepted 1 December 1997

Tn5385 is a ca. 65-kb element integrated into the chromosomes of clinical *Enterococcus faecalis* strains CH19 and CH116. It confers resistance to erythromycin, gentamicin, mercuric chloride, streptomycin, tetracycline, minocycline, and penicillin via β-lactamase production. Tn5385 is a composite structure containing regions previously found in staphylococcal and enterococcal plasmids. Several transposons and transposon-like elements within Tn5385 have been identified, including conjugative transposon Tn5381, composite transposon Tn5384, and elements indistinguishable from staphylococcal transposons Tn4001 and Tn552. The divergent regions of Tn5385 are linked by a series of insertion sequence (IS) elements (IS256, IS237, and IS1216) of staphylococcal and enterococcal origin. The ends of Tn5385 consist of directly repeated copies of enterococcal IS1216. Within the chromosomes of strains CH19 and CH116, Tn5385 has interrupted an open reading frame with substantial homology to previously described alkyl hydrogen peroxide reductase genes. Segments of this open reading frame in both CH19 and CH116 have been deleted, but the amount of deleted DNA differs for the two insertions. Transfer of Tn5385 from both donors into *E. faecalis* recipients occurs at a low frequency. Two types of transconjugants have been identified. In one type, the target alkyl hydrogen peroxide reductase open reading frame has been deleted, and sequences flanking Tn5385 in the respective donors are carried over to the transconjugants. These data suggest that the mechanism of Tn5385 insertion into the recipient chromosome in these transconjugants was recombination across flanking regions in the donors and homologous sequences in the recipients. The second type of transconjugant appears to have resulted from excision of Tn5385 from the CH19 chromosome by recombination across the terminal IS1216 elements and insertion into the recipient chromosome by recombination across Tn5385 (within Tn5385) and a previously transferred Tn5381 copy in the recipient chromosome. These data confirm that Tn5385 is a composite structure with genetic material from diverse genera and suggest that it is a functional transposon. They also suggest that chromosomal recombination is a mechanism of genetic exchange in enterococci.

Large, chromosomally located conjugative elements have been found with some frequency in gram-positive bacteria (3, 6, 7, 16). The most prevalent of these elements are the conjugative transposons, which generally confer tetracycline-minocycline resistance encoded by *tetM* genes and exhibit broad host ranges (23). Conjugative transposons are most commonly 18 to 20 kb in size, although larger variants conferring additional resistance determinants or genes for nisin production have been described in pneumococci and lactococci, respectively (7, 16). The broad host ranges of conjugative transposons suggest that they are important in the dissemination of resistance determinants to diverse genera.

Classic conjugative transposons may be integrated within larger conjugative elements in streptococci and pneumococci. Tn5257, for example, is a Tn916-like *tetM*-encoding element inserted within a larger transposon that encodes chloramphenicol resistance (Tn5252). Tn5252 bears no structural similarity to conjugative transposons and exhibits site-specific integration into recipient cell genomes on transfer. Integration of Tn5252 into recipient chromosomes is thought to be mediated by transposon-encoded gene products that exhibit structural similarity to site-specific recombinases (14, 27).

We previously reported the chromosomal locations of multiresistance (β-lactamase production, erythromycin resistance, high-level gentamicin resistance, mercuric chloride resistance, streptomycin resistance, and tetracycline resistance) encoding transferable elements in *Enterococcus faecalis* CH19 and CH116. The transferable elements within which these resistance genes are incorporated in CH19 and CH116 are structurally indistinguishable, if not identical, and have been given the common designation Tn5385.

In this paper, we present the details of the overall structure of Tn5385. We provide evidence that its ends are formed by directly repeated copies of enterococcal insertion sequence IS1216. In addition, we present evidence that integration of Tn5385 into recipient chromosomes occurs by recombination between flanking sequences in the donor and homologous sequences in recipient genomes. A second mechanism of Tn5385 integration into recipient chromosomes, which involves excision of the element by use of the IS1216 ends and integration into the recipient chromosome by recombination across copies of Tn5381, is also described.

**MATERIALS AND METHODS**

**Strains and plasmids.** Relevant bacterial strains, cloning vectors, and recombinant plasmids are listed in Table 1. CH19 and CH116 are clinical *E. faecalis* strains that were isolated from different patients in the same hospital in 1987. There was no specific epidemiological relationship between these two patients, and the two strains exhibit different plasmid profiles, but their *Sm* resistance profiles are identical (18). *E. faecalis* JH2-7 is a plasmid-free recipient strain used in mating experiments (13). *E. faecalis* OG1RF is an OG1 derivative (12). It was constructed by mutating OG1X (Str') to resistance to fusidic acid and rifampin by sequential inoculation of fusidic acid (25 μg/ml) and rifampin (100 μg/ml) plates with ca. 10⁷ CFU of an overnight culture. Single colonies were harvested and purified, and their resistance phenotypes were confirmed by replating on 

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* Corresponding author. Mailing address: Infectious Diseases Service 1110(W), VA Medical Center, 10701 East Blvd., Cleveland, OH 44106. Phone: (216) 791-3800, ext. 4399. Fax: (216) 231-3482. E-mail: lbr@po.cwru.edu.
selective plates containing both rifampin and fusidic acid at the above-mentioned concentrations. CX19 and CV61 are transconjugants resulting from matings between CH19 and HJ2-7. CV123 is a transconjugant resulting from a mating between CH116 and JH2-7.

**Conjugation experiments.** We previously reported that the entire complement of CH19 resistance determinants transferred to enterococcal recipients at a very low frequency (ca. 10⁻⁷ transconjugant/recipient CFU) (18). Conjugation experiments were carried out by mixing 50 μl of each overnight cultures of donor and recipient strains (grown in nonselective brain heart infusion [BHI] broth) in a sterile test tube and then spreading the mixture across a BHI agar plate. Plates were incubated at 37°C overnight. The following day, the confluent growth on the plate was removed with a platinum loop and suspended in 500 μl of sterile saline.

**Hybridization experiments.** Genomic DNA was extracted as described previously (21), with the following modifications. After the lysozyme-RNase-proteinase K step (which was shortened to 2 h), the resulting suspension was mixed with a CTAB (hexadecyltrimethyl ammonium bromide)-NaCl solution and heated to 60°C for 10 min. This mixture was then extracted once with phenol-chloroform-isomyl alcohol (25:24:1) and once with chloroform-isomyl alcohol. DNA was precipitated with 100% isopropanol, washed with 70% ethanol, and resuspended in TE (50 mM Tris, 10 mM EDTA, pH 7.0) buffer. Genomic DNA was digested with restriction enzymes for 1 to 2 h at 37°C in accordance with the specifications of the manufacturer (Promega, Madison, Wis.). Digested DNA was separated on 0.7% to 1% agarose gels overnight. Separated DNA was denatured, neutralized, transferred to nylon filters by using a negative-pressure transfer apparatus (Pharmacia, Uppsala, Sweden), and baked at 80°C for 1 to 2 h. The DNA was then recovered from the filter. Filters were prehybridized and hybridized with digoxigenin-labeled probes overnight at 68°C and washed under conditions of high stringency in accordance with the specifications of the manufacturer (Boehringer Mannheim, Indianapolis, Ind.).

In most cases, DNA probes were derived from cloned fragments and were labeled either by a random primer method in accordance with the protocol supplied by the manufacturer (Boehringer Mannheim) or by PCR amplification of the cloned inserts, using the forward and reverse pUC18 primers and labeling mix as recommended by the manufacturer (Boehringer Mannheim). Probes for the joint region of circularized forms of conjugal transposon Tn5381 were amplified directly from enterococcal genomic DNA as previously described (21). The PCR products were labeled with digoxigenin by a random primer method (Boehringer Mannheim Biochemicals). Since Tn5381 has a single EcoRI site, this probe will hybridize to two fragments of genomic DNA for every Tn5381 insertion, Clonconjugants were identified by colony hybridization technique as previously described (4). These colonies were purified and subcloned as necessary for further sequencing.

**PCR amplification.** Several regions were amplified with primers derived from sequences within or flanking Tn5385. Amplification of genomic DNA was performed on a Perkin-Elmer Cetus 9800 thermal cycler, using Taq DNA polymerase, in accordance with standard protocols as recommended by the manufacturer (Perkin-Elmer Cetus, Rockville, Md.) by electroporation (Bio-Rad, Hercules, Calif.). Transformed preparations were inoculated onto plates containing antimicrobial agents selective for the cloning vectors, and colonies with the appropriate insert were identified by colony hybridization techniques as previously described (4). These colonies were purified and subcloned as necessary for further sequencing.

**TABLE 1. Strains and plasmids used in this study**

<table>
<thead>
<tr>
<th>Strain or plasmid</th>
<th>Resistance phenotype</th>
<th>Description (reference or source)</th>
</tr>
</thead>
<tbody>
<tr>
<td>E. faecalis strains</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CH19</td>
<td>Bla⁺ Em' Gm' Mer' Sm' Tc'</td>
<td>Clinical isolate (18)</td>
</tr>
<tr>
<td>CH116</td>
<td>Bla⁺ Em' Gm' Mer' Sm' Tc'</td>
<td>Clinical isolate (18)</td>
</tr>
<tr>
<td>JH2-7</td>
<td>Fus⁺ Rif'</td>
<td>Recipient strain (13)</td>
</tr>
<tr>
<td>OGI1XR</td>
<td>Fus⁺ Rif'</td>
<td>Recipient strain (12)</td>
</tr>
<tr>
<td>UV202</td>
<td>Fus⁺ Rif'</td>
<td>Recombinant deficient recipient strain (28)</td>
</tr>
<tr>
<td>CX19</td>
<td>Bla⁺ Em' Gm' Mer' Sm' Tc' Fus⁺ Rif'</td>
<td>Transconjugant resulting from mating between CH19 and HJ2-7 (18)</td>
</tr>
<tr>
<td>CV34</td>
<td>Bla⁺ Em' Gm' Mer' Sm' Tc' Fus⁺ Rif'</td>
<td>Transconjugant resulting from mating between CH116 and JH2-7 (this study)</td>
</tr>
<tr>
<td>CV123</td>
<td>Bla⁺ Em' Gm' Mer' Sm' Tc' Fus⁺ Rif'</td>
<td>Transconjugant resulting from mating between CH19 and OGI1XR (this study)</td>
</tr>
<tr>
<td>CV61</td>
<td>Bla⁺ Em' Gm' Mer' Sm' Tc' Fus⁺ Rif'</td>
<td>Transconjugant resulting from mating between CH19 and HJ2-7 (this study)</td>
</tr>
<tr>
<td>Plasmids</td>
<td></td>
<td></td>
</tr>
<tr>
<td>pUC18</td>
<td>Amp'</td>
<td>Cloning vector (Bethesda Research Laboratories)</td>
</tr>
<tr>
<td>pACYC184</td>
<td>Amp' Kmr'</td>
<td>Cloning vector (22)</td>
</tr>
<tr>
<td>pCRII</td>
<td>Amp' Km'</td>
<td>Cloning vector for PCR products (Invitrogen)</td>
</tr>
<tr>
<td>pDL412</td>
<td>Sm'</td>
<td>aad' gene probe (15)</td>
</tr>
<tr>
<td>pCWR81</td>
<td>Amp'</td>
<td>398-bp internal Alu-ClaI fragment of IS2126 cloned into pUC18 (20)</td>
</tr>
<tr>
<td>pCWR281</td>
<td>Amp'</td>
<td>4.3-kb EcoRI-ClaI fragment of CX19, containing the region between the end of Tn5381 and the internal IS2126 within Tn5385, cloned into pUC18 (this study)</td>
</tr>
<tr>
<td>pCWR303</td>
<td>Amp'</td>
<td>650-bp HindIII-ClaI fragment of CV34, containing the left Tn5385-chromosome junction, cloned into pUC18 (this study)</td>
</tr>
<tr>
<td>pCWR308</td>
<td>Amp' Sm'</td>
<td>2.6-kb Clai fragment from CV34, containing the intact streptomycin resistance determinant, cloned into pUC18 (this study)</td>
</tr>
<tr>
<td>pCWR318</td>
<td>Amp'</td>
<td>1.7-kb Clai fragment from CV123, containing the left Tn5385-chromosome junction, cloned into pUC18 (this study)</td>
</tr>
<tr>
<td>pCWR334</td>
<td>Cm' Te'</td>
<td>7-kb Clai fragment of JH2-7, containing the target for Tn5385 insertion, cloned into pACYC184 (this study)</td>
</tr>
<tr>
<td>pCWR347</td>
<td>Cm' Te'</td>
<td>3.1-kb HindIII fragment of CV34, containing the right Tn5385-chromosome junction, cloned into pACYC184 (this study)</td>
</tr>
<tr>
<td>pCWR349</td>
<td>Amp'</td>
<td>Insert from pCWR347 cloned into pUC18 (this study)</td>
</tr>
<tr>
<td>pCWR351</td>
<td>Amp'</td>
<td>1-kb HindIII subfragment of pCWR343, containing the target for Tn5385 insertion, cloned into pUC18 (this study)</td>
</tr>
<tr>
<td>pCWR389</td>
<td>Amp'</td>
<td>5-kb HindIII fragment of CV61, containing the junction of the ends of Tn5385 across a single copy of IS2126, cloned into pUC18 (this study)</td>
</tr>
</tbody>
</table>

* Bla⁺, β-lactamase positive.
DNA sequence analysis. In most cases, DNA sequencing was performed from cloned DNA on double-stranded templates with an A.L.F. automated sequencing kit and fluorescein- or Cy5 indodicarbocyanine dye-labeled forward and reverse primers. In some cases, sequencing was performed directly from PCR amplification products, using a nested primer. PCR products were purified with QIA-quick PCR purification columns (Qiagen, Inc., Chatsworth, Calif.). Cycle sequencing of these products was performed with a GeneAmp PCR System 9600 thermal cycler (Perkin-Elmer Cetus), using the Cy5 autocycle sequencing kit (Pharmacia LKB) in accordance with the manufacturer’s specifications. Primers used in these experiments are listed in Table 2. Sequences were determined with the ALFExpress automated sequencer (Pharmacia LKB). Sequences were compared with sequences entered into the GenBank, EMBL, DDBJ, and PDB databases by using blastn and blastx local alignment search tools (1) and then further analyzed by using the MacDNAsis version 2.0 (Hitachi, Ltd.) and DNASTar (DNAStar, Madison, Wis.) sequence analysis programs.

GenBank accession number. The sequence of the target open reading frame (ORF) within the chromosome of JH2-7 has been entered into the GenBank database. The accession number is AF016233.

### RESULTS

**Structure of Tn5385.** The structure of Tn5385 has been deduced by using a combination of hybridization, cloning, sequencing, and PCR amplification techniques and is shown in Fig. 1. The region between the internal and right-end IS1216 elements has been described in several previous publications (4, 17, 19, 20). The internal structure of Tn5381 has also been described previously (21). Analysis of clones pCWR281 and pCWR308 (Fig. 1) allowed us to determine the relative positions of the internal IS1216 element and the right end of Tn5381 (ca. 3 kb apart) and to show that the ClaI site used to clone the aadeE streptomycin resistance gene was located 75 bp from the left terminus of Tn5381. The distance between the aadeE gene and the left-end IS1216 element is approximately 4

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**TABLE 2. Custom primers used in amplification and sequencing experiments**

<table>
<thead>
<tr>
<th>Primer (Fig. 1 designation)</th>
<th>Sequence (5’→3’)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>346for-80 (A)</td>
<td>TCTGTACCTGTCCTCCTCTTT</td>
<td>Alkyl hydrogen peroxide reductase primer</td>
</tr>
<tr>
<td>IS1216-608Cy5 (B)</td>
<td>TCTCTTGCGGTTTCGCTTG</td>
<td>Nested primer for sequencing from within IS1216</td>
</tr>
<tr>
<td>IS1216For (C)</td>
<td>AATTATAATGCTCTTCTTACGGA</td>
<td>PCR primer for amplifying IS1216-containing fragments</td>
</tr>
<tr>
<td>IS1216-110Cy5 (D)</td>
<td>CCCACGCTACAATAACACA</td>
<td>Nested primer for sequencing from within IS1216</td>
</tr>
<tr>
<td>IS1216Rev (E)</td>
<td>CCCGAGGTACATCTCTGTTT</td>
<td>PCR primer for amplifying IS1216-containing fragments</td>
</tr>
<tr>
<td>CHATAT1216 (F)</td>
<td>TGAAGAAAACAAAAGGAGATG</td>
<td>Primer downstream of aadeE gene; used to read outward from left end of Tn5385</td>
</tr>
<tr>
<td>Strep DOWN (G)</td>
<td>TGAACACCCCTAAACACATACTTT</td>
<td>Downstream of aadeE, toward left end of Tn5385</td>
</tr>
<tr>
<td>Strep UP (H)</td>
<td>TGAACACCCCTAAACACATACTTT</td>
<td>Upstream of aadeE ORF, toward Tn5381</td>
</tr>
<tr>
<td>JointR (I)</td>
<td>CGTGAAGATATCTCTTACAGT</td>
<td>Primer reading outward from end of Tn5381</td>
</tr>
<tr>
<td>JointL (J)</td>
<td>GGATAATACTGCTATCAAGG</td>
<td>Primer reading outward from end of Tn5381</td>
</tr>
<tr>
<td>86-301 (K)</td>
<td>AAATAAAAGACCGAAATACAGG</td>
<td>Primer downstream of β-lactamase gene; used to read outward from right end of Tn5385</td>
</tr>
<tr>
<td>351 forward (L)</td>
<td>GGACAATTTAAGACCTTGAG</td>
<td>Alkyl hydrogen peroxide reductase primer</td>
</tr>
</tbody>
</table>

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**FIG. 1. Schematic portrayal of the structure of Tn5385.** Symbols for the different insertion sequences are shown at the top of the figure. Inserts from relevant plasmids are indicated above the figure. Relevant restriction sites are also indicated above the transposon (Cl, ClaI; Ec, EcoRI). The positions of the different resistance genes are noted above the transposon (Bla, β-lactamase gene; Em, ermA erythromycin resistance gene; Gm, aac6’-aph2’ high-level gentamicin resistance gene; Mer, merKAB mercuric chloride resistance genes; Sm, aadeE streptomycin resistance gene; and Tc, tetM tetracycline-minocycline resistance gene). Different transposons and transposon-like elements are indicated by boxes below the transposon. Arrows below the transposon represent primers used for amplification and sequencing in these experiments (see Table 2 for details of primers).
Partial sequencing of the region immediately internal to the left-end IS1216 revealed 127 bases with 97% sequence identity to an internal region of ORF zeta from Streptococcus pyogenes plasmid pSM19035 (5). This ORF, which is interrupted by IS1216, has no defined function. Sequence analysis of another 400-bp region between the aadE gene and IS1216 revealed no homology with sequences in the database. These data allowed us to construct the detailed map of Tn5385 shown in Fig. 1. Details of primers listed are in Table 2, and their positions are shown in Fig. 1. The total distance between the terminal IS1216 elements of Tn5385 is approximately 65 kb.

**Directly repeated copies of IS1216 form the ends of Tn5385.** In previously published experiments, we identified IS1216 downstream of the β-lactamase gene within Tn5385 (20). An internal fragment of this IS element was cloned into pUC18 for use as a probe (pCWR91). We detected five copies of IS1216 within the genomes of donor strains CH19 and CH116 (Fig. 2) (the two additional copies observed in CH116 represent a recent [after March 1997] change in the IS1216 profile of this strain). Two transconjugants in which all resistance determinants were transferred were selected for further study. The first, CV34, was the product of a mating between CH116 and JH2-7. The second, CV123, was the product of a mating between CH19 and OG1XRF. JH2-7 has two copies of IS1216 present within its genome (Fig. 2), neither of which is involved in the transfer event. OG1XRF is devoid of IS1216 copies. Hybridization of genomic DNA from CV34 and CV123 after digestion with either HindIII or HincII (neither of which cuts sites within IS1216) revealed the transfer of three copies of IS1216 in association with the resistance determinants of Tn5385 (Fig. 2). No change in restriction fragment size was observed when the IS1216-hybridizing fragments from donors and transconjugants were compared, suggesting that the three copies of IS1216 were all internal to the transferred region. Pulsed-field gel electrophoresis of SmaI-digested DNA from donors and transconjugants revealed two similarly sized IS1216-hybridizing fragments in CH19, CH116, CV34, and CV123 (Fig. 3). The sizes of these fragments added up to ca. 100 kb, which supported the hypothesis that the transferred region was larger than the 65 kb between the directly repeated IS1216 elements. However, IS1216 hybridization of NotI-digested DNA from OG1XRF and CV123 revealed only one hybridizing fragment (data not shown). The OG1XRF fragment into which insertion occurred increased in size from ca. 280 kb to ca. 350 kb. This size was consistent with the transferred DNA being Tn5385. The assumption that the DNA acquired by the transconjugant consisted of at least 100 kb (based on SmaI digestion of donor and transconjugant DNA) was therefore incorrect. The alternative hypothesis suggested by these findings was that Tn5385 had transferred into a very specific site within the JH2-7 chromosome in which flanking SmaI sites were conserved in the donor and the recipient.

To investigate this possibility, we cloned the left IS1216-chromosome junction from CV123, a subfragment devoid of IS1216 sequences was then used as a probe of genomic digests of OG1XRF, CV123, JH2-7, and CV34. This probe was used to identify the target site for insertion in JH2-7, which was found to be within a 7-kb ClaI fragment. Sequence analysis of the target site revealed an ORF with significant homology to previously described alkyl hydrogen peroxide reductase genes from several different species (2, 10). This target fragment was then used to confirm that the terminal IS1216 elements of Tn5385 represented the ends of the transferred DNA. The junction sequences on both sides of Tn5385 were determined for CH19, CH116, CV34, and CV123, and the target region for OG1XRF was also determined. In all cases, Tn5385 was inserted within the putative alkyl hydrogen peroxide reductase ORF. A comparison of the junction sequences from the different strains is shown in Fig. 4. The right-end Tn5385-chromosome junctions in CH19 and CH116 were identical. The left junctions of the two strains differed. Both left junctions demonstrated deletions of the target ORF. However, the deletion adjacent to Tn5385 (CH116) was 151 bp larger than that adjacent to Tn5385 (CH19). When sequences of junctions from transconjugants CV34 and CV123 were compared with those of the donors, it was noted that the identical deletions present in the donors were carried over to the transconjugants. In addition, base changes characteristic of flanking sequences in the donor (as opposed to the target sequence in the recipient) were also present in the transconjugants. Moreover, the target ORF, intact after PCR amplification in both JH2-7 and...
OG1XRF, was lost in the transconjugants (Table 3), suggesting that insertion of Tn5385 into the recipient genome occurred by a process that deleted the original target, a finding most consistent with homologous recombination being the mechanism of insertion. In support of this hypothesis, we have never observed transfer of Tn5385 into E. faecalis UV2012, a recombination-deficient strain, despite several attempts.

**Evidence for a second insertion mechanism.** We noted a systematic and exhaustive fashion. A graphic description of the proposed mechanisms of the two transfer events is illustrated in Fig. 5.

**TABLE 3. Amplification products of PCR with selected primer pairs**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Amplification with primer paira:</th>
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<tbody>
<tr>
<td></td>
<td>A-F</td>
<td>K-L</td>
</tr>
<tr>
<td>CH19</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>CV123</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>CV61</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CH116</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>CV34</td>
<td>+</td>
<td>+</td>
</tr>
</tbody>
</table>

a See Fig. 1 and Table 2 for identities of primers.

**DISCUSSION**

We previously reported analyses of regions of Tn5385 that suggested it had evolved as a composite of staphylococcal and enterococcal plasmids and transposons (4, 17). The data reported in this paper further extend these observations. In sum, Tn5385 is a ca. 65-kb composite structure that includes segments characteristic of diverse species and genera, including *Staphylococcus aureus* (IS257, mercury resistance operon, Tn552-like β-lactamase transposon, and relaxase-mobilization region of small staphylococcal plasmids), broad-host-range en-
enterococci (Tn5381, Tn4001, broad-host-range replication genes, and Tn917), enterococci (IS1216), and S. pyogenes (partial ORF zeta from S. pyogenes plasmid pSM19035). It appears that Tn5385 originated as a plasmid, one which became more complex as it cointegrated with other plasmids. The ability to acquire genetic material from such diverse genera was probably conferred by broad-host-range plasmid conjugation genes that have subsequently been deleted (4). It is intriguing that roughly 38 kb (58%) of Tn5385 is composed of antimicrobial resistance determinants or of structures conferring mobility to such determinants. The concentration of this collection in a small region is reminiscent of integrons in gram-negative bacilli. Although no integron-like mechanism is obvious from the structure of Tn5385, it is tempting to speculate that this collection of determinants represents a gram-positive equivalent of an integron, mediated by the activity of a variety of IS elements. This report represents the first example of such a complex, chromosomally based element in enterococci.

The structure of Tn5385 as described in this paper suggests a mechanism for insertion of the putative composite plasmid into the bacterial chromosome: cointegration mediated by a copy of IS1216 within the presumed plasmid. Supportive of this scenario is the fact that portions of the same ORF are found flanking the terminal, directly repeated IS1216 elements. IS1216 is a member of the large IS1 family of ISs that characteristically generate 8-bp duplications of the target sequence on insertion; these were not observed in the two Tn5385 insertions detailed in this paper (8). The absence of target duplications flanking Tn5385 is explainable by the occurrence of secondary transposition events resulting in deletions of different segments of the target ORF in CH19 and CH116. In most other respects (e.g., SmalI digest pulsed-field gel electrophoresis patterns and IS6770 hybridization patterns), CH19 and CH116 are indistinguishable. The observed differences in sequence immediately adjacent to the Tn5385 insertion sites in the two strains are consistent with the occurrence of a single insertion event followed by divergence in the quantity of deleted adjacent DNA associated with subsequent rearrangements. Tn5385 appears to require the replication functions of either the chromosome or another plasmid, since the presumed replication origin (for broad-host-range plasmids) has been deleted (4).

Although Tn5385 does not meet the classic definition of a transposon based on currently available information, it exhibits many characteristics of known transposons. First, it is flanked by directly repeated copies of an IS element (the characteristic conformation of composite transposons) known to be insertionally active in enterococci (11). Previous reports have implicated IS1216 in the transposition of vancomycin resistance determinants in E. faecium (11). Second, at least one of the terminal IS1216 copies (the one on the right end) appears to have been involved in the original insertion event, since the insertion sites in both CH19 and CH116 are identical. It is unclear whether the left-end IS1216 elements in CH19 and CH116 are the copies involved in the original insertion, since subsequent rearrangements appear to have occurred. Finally, the chromosomal insertion of Tn5385 in CV61 is consistent with circularization of Tn5385 followed by entry into the recipient chromosome by recombination across copies of conjugative transposons. Evidence of circularization, while not definitive, is highly suggestive that Tn5385 is in fact a transposable element.

There has been debate in the published literature about whether conjugative transposition proceeds by a mechanism

![Diagram of Tn5385 insertion into enterococci](http://www.asm.org/jb)
resembling a cell fusion event. Torres et al. originally reported the transfer of unlinked chromosomal loci between *Bacillus subtilis* strains in the presence of conjugal transposon Tn925 in the donor chromosome (26). Showsh and Andrews failed to detect the occurrence of retrotransfer of nonconjugal plasmids in association with conjugal transposition of Tn916 or Tn925, concluding that a cell fusion-like event was unlikely (24). The data presented in this paper are consistent with the occurrence of a cell fusion-like event. The insertion of Tn5385 in recipient chromosomes, resulting in transconjugants CV34 and CV123, bears all of the marks of recombination across regions of chromosomal homology, which would be consistent with cell fusion. The facts that IS2126-hybridizing SmalI fragments are identical in size in donors and transconjugants and that they add up to ca. 100 kb suggest that the area of crossover is substantial and considerably greater than the limits of Tn5385. The carryover of the specific deletions to the transconjugants argues strongly against site-specific insertion of circularized Tn5385 as the mechanism of entry, although circularization into a much larger element than is defined by the terminal IS2126 elements, one that includes the flanking regions of homology and possesses transfer genes, remains a possibility that would not require invoking cell fusion.

Insertion of Tn5385 into the enterococcal chromosome has allowed us to identify a homolog of alkyl hydrogen peroxide reductase genes found in several other genera (2, 10). Mutant strains of *Salmonella typhimurium* that lack the alkyl hydrogen peroxide reductase gene have been shown to be extremely sensitive to killing by organic hydroperoxides (25). A similar gene in *Staphylococcus aureus* has been shown to be expressed at increased levels following osmotic stress (2). Preliminary experiments comparing the abilities of recipient and transconjugant strains to grow in the presence of increasing concentrations of hydrogen peroxide revealed no difference in growth rates (data not shown). The role played by this gene in the survival and growth of enterococci remains unclear.

We do not at the present time know what promotes the conjugation event resulting in Tn5385 transfer. A single plasmid is present in CH116 but does not transfer in association with Tn5385 (data not shown). CH116 has no detectable plasmids. It remains possible that transfer is mediated by Tn3571, the conjugal transposon within Tn5385. Tn5381 transfers alone at a much higher frequency than it does when within Tn5385 (ca. 10^{-6} and 10^{-7}/recipient CFU, respectively) (21). In addition, conjugal transfer of Tn5381 is increased after preincubation of donor strains with tetracycline, apparently because of an increased rate of excision of the element (21). Preincubation with tetracycline does not lead to an increase in the transfer of Tn5385 from either CH119 or CH116 (data not shown), arguing against the possibility that Tn3571 excision stimulates transfer of Tn5385. If, however, Tn5381 transfers by creating cell fusion events, the random transfer of Tn5385 by chromosomal recombination would be expected to occur at a low frequency. Experiments to investigate the role of Tn5381 in the transfer of Tn5385 are planned.

The data presented in this paper emphasize the important role played by insertion elements in the evolution of antimicrobial resistance in gram-positive genera. A circularized form of Tn5385, if it can exist, would be a highly versatile integration element, since it would possess several active IS elements which would stimulate cointegration with other replicons (IS256, three copies; IS2126, two copies; and IS257, two copies). In addition, each of these frequently repeated elements (including Tn5381) can cointegrate with other replicons by homologous recombination, an event which apparently occurred across copies of Tn5381 to result in the transconjugant CV61. The ability of different mobile elements to cooperate in this fashion is a powerful tool for the dissemination of antimicrobial resistance determinants among gram-positive genera.

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**REFERENCES**


