Chromosomally and Extrachromosomally Mediated High-Level Gentamicin Resistance in Streptococcus agalactiae

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Streptococcus agalactiae (group B Streptococcus [GBS]) is a leading cause of sepsis in neonates. The rate of invasive GBS disease in nonpregnant adults also continues to climb. Aminoglycosides alone have little or no effect on GBS, but synergistic killing with penicillin has been shown in vitro. High-level gentamicin resistance (HLGR) in GBS isolates, however, leads to the loss of a synergistic effect. Our study performed a multicenter study to determine the frequency of HLGR GBS isolates and to elucidate the molecular mechanisms leading to gentamicin resistance. From eight centers in four countries, 1,128 invasive and colonizing GBS isolates were pooled and investigated for the presence of HLGR. We identified two strains that displayed HLGR (BSU1203 and BSU452), both of which carried the aacA-aphD gene, typically conferring HLGR. However, only one strain (BSU1203) also carried the previously described chromosomal gentamicin resistance transposon designated Tn3706. For the other strain (BSU452), plasmid purification and subsequent DNA sequencing resulted in the detection of plasmid pIP501 carrying a remnant of a Tn3 family transposon. Its ability to confer HLGR was proven by transfer into an Enterococcus faecalis isolate. Conversely, loss of HLGR was demonstrated by curing both GBS BSU452 and the transformed E. faecalis strain from the plasmid. This is the first report showing plasmid-mediated HLGR in GBS. Thus, in our clinical GBS isolates, HLGR is mediated both chromosomally and extrachromosomally.

Streptococcus agalactiae, alternatively designated group B Streptococcus (GBS), is a leading cause of morbidity and mortality in neonates and pregnant women. Recommendations for diagnosing maternal GBS colonization and administering intrapartum antimicrobial prophylaxis have lead to a significant decrease in these infections (1). The rate of invasive GBS disease in nonpregnant adults, however, continues to climb (2). Elderly persons and those with underlying diseases—two expanding segments of the population—are at increased risk (3). Treatment concepts for invasive GBS infections in nonpregnant adults have not yet been established. Clinical isolates of GBS are susceptible to penicillin, the antimicrobial agent of choice for treating invasive diseases. Several publications advocate the addition of an aminoglycoside to penicillin or ampicillin for infective endocarditis (4) and periprosthetic joint infections (5), although aminoglycosides have ototoxic and nephrotoxic side effects, in particular in the elderly. Aminoglycosides alone have little or no effect on GBS, but synergistic killing with penicillin has been shown in vitro (6). In case of the presence of high-level gentamicin resistance (HLGR) in a bacterial isolate, there is a lack of a synergistic effect.

While HLGR in Enterococcus spp. is frequently found (7), to the best of our knowledge, only two HLGR GBS strains have been previously reported (8, 9). Most diagnostic laboratories do not test routinely for HLGR in GBS. Thus, little is known about the frequency of HLGR GBS, the mechanisms of acquiring HLGR, and the potential to spread genetic elements associated with HLGR. The aim of this study was to estimate the frequency of HLGR GBS isolates (i) in systematically and continuously collected GBS isolates from colonized pregnant and nonpregnant women and (ii) in GBS isolates pooled in a collection that stems from various selected patient populations. Upon detection of HLGR isolates, we elaborated the molecular mechanism conferring this resistance.

MATERIALS AND METHODS

GBS isolates. The study included 1,128 GBS isolates. Of these, 464 (41%) were pooled from various GBS collections (Table 1). These isolates stem from various centers and were previously investigated in other contexts (10–15). The other 664 (59%) GBS isolates were prospectively collected and screened for the presence of HLGR. The origin of GBS isolation, the association with diseases or colonization, and the sampling period are presented in Table 1.

Definition and identification of HLGR in GBS. No definition of HLGR in GBS has been previously published. According to the recommended screening tests for the detection of HLGR in Enterococcus spp., the resistant isolates have an MIC of ≥500 mg/liter (16). In addition, HLGR isolates with an MIC >500 mg/liter have been reported (17, 18). Therefore, HLGR in GBS was defined when the gentamicin MIC determined by Etest was ≥512 mg/liter. All MIC determinations were confirmed with ≥3 measurements.

Two different methods for the identification of HLGR in GBS were compared: (i) in GBS isolated from colonized pregnant women and (ii) in GBS isolates pooled in a collection that stems from various selected patient populations. Upon detection of HLGR isolates, we elaborated the molecular mechanism conferring this resistance.
applied. Five hundred sixty-one isolates (49.7%) were plated on Mueller-Hinton agar supplemented with 256 mg/liter of gentamicin. Subsequently, the MIC of growing GBS colonies was determined by Etest. For 567 (50.3%) isolates, the MIC was primarily determined by Etest without a prior HLGR screening test.

**Bacterial strains.** The strains used in this study are presented in Table 2. The plasmid-free recipient used in the mating experiments was *Enterococcus faecalis* (BSU386), a clinical blood culture isolate without HLGR.

**Genetic basis of HLGR.** Standard recombinant DNA techniques were used for nucleic acid preparation and analysis. Plasmid DNA was isolated and purified using the QIAprep Spin Miniprep kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. PCR was performed with *Taq* polymerase according to the manufacturer’s protocol (Roche Diagnostics, Mannheim, Germany), with 35 cycles of amplification steps consisting of 1 min at 94°C, 1 min at 55°C, and 1 min at 72°C. PCR products were sequenced on an ABI PRISM 310 genetic analyzer, using the ABI PRISM BigDye Terminator v1.1 cycle sequencing kit (Applied Biosystems, Weiterstadt, Germany). To identify the HLGR resistance gene, we performed multiplex PCR as described by Vakulenko et al. (19). For the detection of Tn3706-specific nucleotide sequences, we used PCR with primers O1, O2, and O3 as described by Horaud et al. (20). The primers used for this study are presented in Table 3. Detection of open

**TABLE 1** Pooled collection of isolates investigated for the presence of high-level gentamicin resistance

<table>
<thead>
<tr>
<th>Reference</th>
<th>No. of GBS isolates</th>
<th>Disease/case definition</th>
<th>Origin of GBS isolation</th>
<th>Study type</th>
<th>Collection period</th>
<th>Geographic origin(s)</th>
<th>HLGR</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>75</td>
<td>No disease/colonization</td>
<td>Vaginal and rectal swabs from pregnant and nonpregnant women</td>
<td>Cross-sectional study</td>
<td>2001–2003</td>
<td>Aachen and Munich, Germany</td>
<td>0</td>
</tr>
<tr>
<td>12</td>
<td>60</td>
<td>EOD&lt;sup&gt;b&lt;/sup&gt; with invasive neonatal GBS infections</td>
<td>Isolation of GBS from blood or CSF&lt;sup&gt;b&lt;/sup&gt; and other sterile body fluids</td>
<td>Part of the prospective active surveillance study</td>
<td>2001–2003</td>
<td>Freiburg, Germany</td>
<td>0</td>
</tr>
<tr>
<td>13</td>
<td>50&lt;sup&gt;c&lt;/sup&gt;</td>
<td>Suspicion of EOD without proven invasive GBS disease</td>
<td>GBS isolates from nonsterile sites</td>
<td>Part of the prospective active surveillance study</td>
<td>2001–2003</td>
<td>Freiburg, Germany</td>
<td>0</td>
</tr>
<tr>
<td>14</td>
<td>30</td>
<td>Patients with cystic fibrosis</td>
<td>Respiratory samples</td>
<td>Collection of isolates&lt;sup&gt;d&lt;/sup&gt;</td>
<td>2002–2008</td>
<td>Münster, Germany</td>
<td>1</td>
</tr>
<tr>
<td>11</td>
<td>150</td>
<td>No disease/colonization</td>
<td>Rectovaginal specimens from pregnant and nonpregnant women</td>
<td>Part of the national surveillance study</td>
<td>2005–2009</td>
<td>Lisbon, Portugal</td>
<td>0</td>
</tr>
<tr>
<td>97</td>
<td>No disease/colonization</td>
<td>Vaginal swabs from pregnant and nonpregnant women</td>
<td>—&lt;sup&gt;e&lt;/sup&gt;</td>
<td>2009</td>
<td>Ulm, Germany</td>
<td>0</td>
<td></td>
</tr>
<tr>
<td>15</td>
<td>99</td>
<td>No disease/colonization</td>
<td>Vaginal swabs from pregnant and nonpregnant women</td>
<td>Cross-sectional study</td>
<td>2010</td>
<td>Ismailia, Egypt</td>
<td>0</td>
</tr>
<tr>
<td>36</td>
<td>364</td>
<td>No disease/colonization</td>
<td>Vaginal swabs from pregnant women</td>
<td>Cross-sectional study</td>
<td>2009–2010</td>
<td>Bern, Switzerland</td>
<td>1</td>
</tr>
<tr>
<td>203</td>
<td>Invasive GBS infections</td>
<td>Isolation of GBS from blood, CSF and other sterile body fluids</td>
<td>—&lt;sup&gt;f&lt;/sup&gt;</td>
<td>1998–2013</td>
<td>Bern, Switzerland</td>
<td>0</td>
<td></td>
</tr>
</tbody>
</table>

Total 1,128 2

<sup>a</sup> GBS isolates investigated and published in a context other than the presence of HLGR. The number of GBS isolates investigated for HLGR may vary from the number in the source publication for technical reasons.

<sup>b</sup> EOD, early-onset disease.

<sup>c</sup> For this study, only GBS isolates with serotype III were available.

<sup>d</sup> GBS isolation occurred during a routine visit or during a visit due to an exacerbation of clinical symptoms.

<sup>e</sup> Prospective collection during routine diagnostic microbiology laboratory analysis. GBS isolates were investigated for this study.

<sup>f</sup> Collection of invasive GBS isolates (all age groups) during routine diagnostic microbiology laboratory analysis. GBS isolates were investigated for this study.

<sup>g</sup> CSF, cerebrospinal fluid.

**TABLE 2** Bacterial strains and their corresponding genetic elements conferring HLGR

<table>
<thead>
<tr>
<th>Species</th>
<th>Strain</th>
<th>Description</th>
<th>MIC of gentamicin (mg/liter)</th>
<th>aac(6')-Ie-aph(2')-Ia gene</th>
<th>Transposon</th>
</tr>
</thead>
<tbody>
<tr>
<td>GBS</td>
<td>BSU1203</td>
<td>Wild-type strain</td>
<td>≥1,024</td>
<td>Yes</td>
<td>Tn3706</td>
</tr>
<tr>
<td>GBS</td>
<td>BSU452</td>
<td>Wild-type strain</td>
<td>512</td>
<td>Yes</td>
<td>Tn3-like</td>
</tr>
<tr>
<td>E. faecalis</td>
<td>BSU386</td>
<td>Wild-type strain</td>
<td>12</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>E. faecalis</td>
<td>BSU580</td>
<td>BSU386 + pIP501&lt;sub&gt;BSU452&lt;/sub&gt;</td>
<td>≥1,024</td>
<td>Yes</td>
<td>Tn3-like</td>
</tr>
<tr>
<td>E. faecalis</td>
<td>BSU720</td>
<td>BSU580 cured</td>
<td>12</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>GBS</td>
<td>BSU729</td>
<td>BSU452 cured</td>
<td>24</td>
<td>No</td>
<td>No</td>
</tr>
</tbody>
</table>
reading frames (ORFs) was carried out by using ORF Finder (http://www.ncbi.nlm.nih.gov/gorf/gorf.html). Sequence comparison was performed by using the BLAST system (http://blast.ncbi.nlm.nih.gov/Blast.cgi).

Transfer and mobilization of the genetic element conferring HLGR. To investigate the potential transfer of resistance, we transformed *E. faecalis* (BSU386) with plasmid DNA, as described previously (21). Curing the transformed *E. faecalis* strain and GBS BSU452 with HLGR was achieved as follows. The strains were exposed in an overnight culture to serial 2-fold dilutions of ciprofloxacin in Todd-Hewitt broth plus 0.5% yeast extract and Luria-Bertani medium. Bacterial cultures containing the highest subinhibitory concentrations of ciprofloxacin (i.e., 0.625 mg/liter for GBS BSU452 and 10 mg/liter for *E. faecalis* BSU580) were plated on antibiotic-free tryptic soy agar plates and grown overnight at 37°C. Single colonies were then tested for loss of resistance to gentamicin by subculture onto Mueller-Hinton agar plates containing 256 mg/liter of gentamicin. The MIC for gentamicin was then determined by Etest.

Nucleotide sequence accession number. Nucleotide sequences were submitted to GenBank under accession number KP698941.

RESULTS

GBS isolates with HLGR. Among 1,128 GBS isolates, 2 (0.17%) strains with HLGR were identified. One strain (BSU1203; MIC, >1,024 mg/liter) was obtained from a 35-year-old Swiss woman during prenatal screening. The second HLGR GBS strain (BSU452; MIC, 512 mg/liter) was isolated in a respiratory specimen from a 26-year-old man with cystic fibrosis.

PCR detection of genes conferring HLGR on GBS. To investigate the resistance determinants in HLGR GBS strains, we performed PCRs that were specific for the *aac(6′)-Ie-aph(2′)-Ia* gene and the flanking IS256 element (19). The *aac(6′)-Ie-aph(2′)-Ia* gene was readily detected in both strains, yielding the expected product of 348 bp (19).

Detection of a previously known transposon and a novel element. A PCR with a primer set annealing on the structural gentamicin resistance gene and the IS256 sequence, located downstream of this gene, showed the expected 369-bp product for strain GBS BSU1203 (19), indicating the presence of the previously described chromosomal transposon Tn3706. For further characterization of the resistance determinant, PCRs that were specific for insertion sequence elements of Tn3706 were performed as published previously (20). These PCRs were positive with GBS strain BSU1203 and matched those previously described for HLGR GBS strain B128 (20), confirming the presence of Tn3706. In GBS strain BSU452, the *aac(6′)-Ie-aph(2′)-Ia* gene was found, but none of the PCRs specific for the transposon structures of Tn5281, Tn4001, or Tn3706 (20) yielded a product, suggesting the presence of a novel HLGR determinant in this strain.

Characterization of a novel mobile genetic element conferring HLGR on GBS. To identify the genetic structure of GBS strain BSU452 carrying the *aac(6′)-Ie-aph(2′)-Ia* gene, we performed an inverse PCR on a plasmid preparation of BSU452. Primers annealing to the gentamicin resistance gene (Table 3) and directed toward DNA regions upstream and downstream of *aac(6′)-Ie-aph(2′)-Ia* yielded a PCR product about 2 kb in length, which was completely sequenced. Nucleotide comparison with the GenBank database revealed a 100% identity of nucleotides 1 to 253 and 721 to 2029 with plasmid pTEF1 of *E. faecalis* strain V583 (GenBank accession number AE016833.1). Several ORFs were identified on the 2-kb PCR product, and comparison of the deduced amino acids with the GenBank database revealed a DNA resolvase fragment and one copy of an insertion sequence element with high homology to IS1216. This structure displayed high similarities to a Tn3 family transposon.

Transfer and mobilization of the resistance determinant in association with HLGR. The HLGR genes and the flanking DNA sequences in GBS BSU1203 matched those previously identified in GBS B128, and because thorough molecular analyses on the acquisition of HLGR have been published for that strain (8, 20, 22), further investigations focused on GBS BSU452. Tn3 family transposons are typically located on plasmids. To investigate if this is the case in strain GBS BSU452 and to characterize the potential of spreading HLGR to other isolates, we transformed the gentamicin-susceptible *E. faecalis* strain BSU386 with the plasmid preparation obtained from GBS BSU452. Positive clones (i.e., designated *E. faecalis* BSU580, which carries the mobile element of BSU452) were obtained upon plating the transformed strain onto the HLGR screening agar, as described above. A subsequent gentamicin evaluation revealed an increase in MIC from 12 mg/liter to ≥1,024 mg/liter (Table 2). To ensure that the increased MIC was due to the uptake of the plasmid DNA, plasmid preparations were subjected to gel electrophoresis (Fig. 1), which showed the presence of large plasmids in the HLGR strains BSU452 and BSU580. Further confirmation of the successful transfer was achieved by PCR showing the presence of the *aac(6′)-Ie-aph(2′)-Ia* gene and a lack of any flanking IS256 sequences in *E. faecalis* strain BSU580. To confirm that the newly detected mobile genetic element is indeed located on a plasmid, we attempted to cure GBS BSU452 and the transformed *E. faecalis* BSU580 from the plasmid. This was successfully achieved by growing the strains in subinhibitory concentrations of ciprofloxacin, as described by Eliopoulos et al. (23). Under these conditions, clones of both GBS BSU452 and the transformed *E. faecalis* strain BSU580 lost their elevated resistance to gentamicin. The MICs decreased in GBS strain BSU729 (i.e., strain BSU452 after plasmid curing) from 1,024 mg/liter to 24 mg/liter and in *E. faecalis* BSU720 (i.e., strain BSU580 after plasmid curing) from ≥1,024 mg/liter to 12 mg/liter (Table 2). In addition, the lack of a plasmid in the cured strains could be demonstrated by gel electrophoresis (Fig. 1). Plasmid loss in the presence of subinhibitory ciprofloxacin occurred at a frequency of about 0.02% (1 in 4,500 colonies in GBS and 1 in 6,000 colonies of *E. faecalis*). One of the most commonly found plasmids in GBS and enterococci is

### Table 3 Primers used for PCR and DNA sequencing

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5′-3′)</th>
<th>Target gene</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inverse primer</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HLGR1</td>
<td>CTCATCCTTCCACGCTCTG</td>
<td><em>aac(6′)-Ie-aph(2′)-Ia</em></td>
</tr>
<tr>
<td>HLGR2</td>
<td>GCCAGAACATGAATACGAGG</td>
<td><em>aac(6′)-Ie-aph(2′)-Ia</em></td>
</tr>
<tr>
<td>369 Vakulenko</td>
<td>CAGGAAATTTCTGGAACAGTGA</td>
<td><em>aac(6′)-Ie-aph(2′)-Ia</em></td>
</tr>
<tr>
<td>348 Vakulenko</td>
<td>CAGGCCCTTGGGAAAGTGAGA</td>
<td><em>aac(6′)-Ie-aph(2′)-Ia</em></td>
</tr>
<tr>
<td>Primer O1</td>
<td>GACCATCAGTGAATTGA</td>
<td></td>
</tr>
<tr>
<td>Primer O2</td>
<td>CTTTTTACAGAATTTCAATAAGC</td>
<td></td>
</tr>
<tr>
<td>Primer O3</td>
<td>GTATAG CAAATGCAATACTC</td>
<td></td>
</tr>
<tr>
<td>pIP501-for</td>
<td>TGGCTCAATCATTCAAACGC</td>
<td></td>
</tr>
<tr>
<td>pIP501-rev</td>
<td>CTTGAAAGGATATAGGCCCCTT</td>
<td></td>
</tr>
</tbody>
</table>
most Enterococcus spp. with HLGR, the transposon harboring theaac(6')-aph(2')gene is found on a plasmid (28). Truncated forms ofTn4001 are typically located on plasmid DNA (29). Intact
Tn4001 transposons can also be located on chromosomal DNA. In
the previously described HLGR GBS strain B128, theaacA-aphD
gene was found on a Tn4001 derivative (designated Tn3706), lo-
cated on chromosomal DNA (8). In one of our strains (BSU1203),
the finding of transposon Tn3706 conferring HLGR is in agree-
ment with the previously published findings about HLGR GBS
strain B128 (8, 20, 22). Horaud et al. (20) described that its trans-
position from E. faecalis occurred on GBS plasmid pIP501. How-
ever, after conjugative transfer between GBS strains, the hybrid
repli
cons pIP501::Tn3706 were found to be structurally unstable. This
observation indicated that streptococcal pIP501-like plas-
"mids do not constitute appropriate delivery vectors for the dis-
semination of Tn3706 among GBS, and therefore, HLGR is found
relatively rarely among GBS isolates (20). Although these argu-
ments speak against a high potential for spread, the persistence
of HLGR in GBS strain B128 and BSU1203 indicates that Tn3706 can
be stably integrated into the chromosome.

In GBS BSU452, we identified a different mobile genetic ele-
ment. The genes surrounding theaac(6')-aph(2')gene did not
display the structures of transposon Tn4001 or any of the closely
related derivatives or its truncated forms. We detected plasmid
pIP501, which is a conjugative plasmid that often carries multi-
resistance genes. It has previously been described for S. agalactiae
in association with HLGR and belongs to the Inc18 group of plas-
"mids (24). Tn3 family transposons are commonly associated with
Inc18 plasmids and often confer antibiotic resistance on Entero-
coccus spp. (30). They are, however, typically associated with gly-
copeptide and macrolide resistance (31) and not HLGR. Investi-
gators have previously reported the presence of an IS1216
transposase on Tn3-like remnants (32), as we found in our GBS
BSU452 strain; however, IS1216 is typically associated with tetra-
cycline resistance in streptococcal species (33). To the best of our
knowledge, the detection of theaacA-aphDgene on a Tn3-like
transposon and the presence of IS1216 in association with HLGR
constitute novel findings. They have been reported neither for
enterococci nor for GBS.

Resistance determinant in GBS BSU452 shows close ho-
"mologies to parts of the enterococcal resistance plasmid pTEF1 of
E. faecalis strain V583 (34), suggesting that it may have been
transferred through horizontal gene transfer. This is, however,
speculative for GBS strain BSU452, since the presence of an
HLGR Tn3-like transposon in GBS has not been previously
described. Nevertheless, horizontal gene transfer of resistance
genes from Enterococcus spp. to other Gram-positive bacteria by
mobile genetic elements is a well-described mechanism in the
spread of antibiotic resistance (30, 31). Horizontal gene transfer
has recently been suggested for the acquisition of vancomycin
resistance genes in GBS (35). GBS strain BSU452 was isolated from
the sputum of a cystic fibrosis patient, but there was no evidence
of enterococcal colonization. Considering that patients with cystic
fibrosis are often treated with antibiotics (including aminoglycosides),
and their microbiome in the respiratory tract is different from that of
untreated healthy patients, it is possible that horizontal gene transfer
to GBS originated from the selected flora. However, this hypoth-
osis cannot be proven in our case and remains speculation.

Though a plasmid-borne HLGR has high potential for further
spread in a GBS population, the likelihood of this happening can-
not be predicted yet. In this study, we demonstrated that pIP501, including the HLGR determinant of GB5452, could easily be transferred to *E. faecalis*. Thus, it is conceivable that transfer to other GBS isolates is also possible, especially in view of the fact that pIP501 is a broad-host-range plasmid, well established in GBS and enterococci.

In conclusion, the overall frequency of HLGR GBS in our large collection of isolates was low. Molecular investigations revealed a transposon located on the chromosome, as previously described for a single isolate (8, 20, 22), and a Tn3 family transposon conferring HLGR in association with pIP501. These findings point toward a new dimension of potential spread of HLGR within GBS.

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We have no conflicts of interest to declare.

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