Insight into the Diversity of Penicillin-Binding Protein 2x Alleles and Mutations in Viridans Streptococci

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ABSTRACT The identification of commensal streptococci species is an everlasting problem due to their ability to genetically transform. A new challenge in this respect is the recent description of Streptococcus pseudopneumoniae as a new species, which was distinguished from closely related pathogenic S. pneumoniae and commensal S. mitis by a variety of physiological and molecular biological tests. Forty-one atypical S. pneumoniae isolates have been collected at the German National Reference Center for Streptococci (GNRCS). Multilocus sequence typing (MLST) confirmed 35 isolates as the species S. pseudopneumoniae. A comparison with the pbp2x sequences from 120 commensal streptococci isolated from different continents revealed that pbp2x is distinct among penicillin-susceptible S. pseudopneumoniae isolates. Four penicillin-binding protein x (PBPx) alleles of penicillin-sensitive S. mitis account for most of the diverse sequence blocks in resistant S. pneumoniae, S. mitis, and S. infantis and S. oralis sequences were found in S. pneumoniae from Japan. PBP2x genes of the family of mosaic genes related to pbp2x in the S. pneumoniae clone Spain23F-1 were observed in S. oralis and S. infantis as well, confirming its global distribution. Thirty-eight sites were altered within the PBP2x transpeptidase domains of penicillin-resistant strains, excluding another 37 sites present in the reference genes of sensitive strains. Specific mutational patterns were detected depending on the parental sequence blocks, in agreement with distinct mutational pathways during the development of beta-lactam resistance. The majority of the mutations clustered around the active site, whereas others are likely to affect stability or interactions with the C-terminal domain or partner proteins.

KEYWORDS S. pseudopneumoniae, PBP2x, penicillin resistance, PBP2x mutation, viridans streptococci

S. pneumoniae is one of the major human pathogens. It colonizes the upper respiratory tract asymptomatically (1), but can lead to a variety of diseases ranging from sinusitis, otitis media, and pneumonia to meningitis (2). By contrast, related streptococci from the mitis group of viridans streptococci (SMG) are commensal organisms that rarely cause disease. S. pneumoniae is differentiated from other species by a variety of phenotypic and genotypic tests on the basis of the presence of important virulence factors carried by almost all S. pneumoniae strains (for a review, see reference 3). This includes the capsule biosynthesis cluster responsible for the expression of more than 90 serotypes (4). Moreover, S. pneumoniae cells lyse in the presence of sodium deoxycholate (DOC) due to the presence of the L-alanine amidase LytA, an autolysin that is also responsible for stationary-phase autolysis. The gene lytA is located
on an island together with ply encoding a potent cytolysin (5). Furthermore, S. pneumoniae is optochin susceptible. Nevertheless, tests for these phenotypes are not completely reliable for identifying S. pneumoniae due to variations in their specificity and sensitivity (6).

S. pneumoniae can be genetically differentiated from its close relatives S. mitis and S. oralis by comparing DNA sequences of housekeeping genes, such as with multilocus sequence typing (MLST) (7), which has been considered the gold standard for studying the epidemiology of pathogenic bacteria. A greater number of species can be resolved by multilocuss sequence analysis (MLSA) (8). However, several studies revealed the problems associated with identifying the species of SMG due to their competence for genetic transformation. Lateral intra- and interspecies gene transfer results in mosaic genes, a highly variable and relatively large accessory genome, and consequently, to diffuse species borders, resulting in “fuzzy species” (8). This is specifically evident in the lineages comprising S. mitis and S. oralis, thereby challenging the definition of species (6, 9–11).

The recently described S. pseudopneumoniae (12) provides additional complications for identifying SMG species. Although closely related to S. pneumoniae, S. pseudopneumoniae may be differentiated by a variety of phenotypic and molecular assays, consistent with the previous classification as “atypical S. pneumoniae.” Some of the key characteristics of S. pseudopneumoniae are the expression of optochin resistance in the presence of 5% CO2, the lack of a pneumococcal capsule, and bile insolubility (12). An analysis of the first S. pseudopneumoniae genome of strain IS7493 documented the lack of some pneumococcal virulence factors, such as the capsule biosynthesis genes and the choline-binding proteins PspC, PcpA, and PspA (13). However, it contained the ply-lytA islet. Comparative genomic hybridization showed that this islet is common among S. pseudopneumoniae strains (13) and among eight sequence types of SMG that have been identified (14, 15). The sequence of lytA is S. pneumoniae is distinct from that of S. pseudopneumoniae and other members of the viridans streptococci (14, 15) and can also be used to differentiate S. pseudopneumoniae by specifically designed PCR.

S. pseudopneumoniae has been isolated mainly from patients with respiratory diseases (16–19). Many of these isolates showed resistance to antibiotics, including a reduced susceptibility to penicillin. The penicillin-binding proteins (PBPs) of S. pseudopneumoniae IS7493 are more similar to PBPs of resistant S. pneumoniae than to the highly conserved PBPs of sensitive strains (13). Since this strain has a decreased susceptibility to beta-lactams, its PBPs might not be representative for this species. The spread of penicillin resistance among populations of S. pneumoniae has been observed for many decades (20), fostered by genetic transformations mediating interspecies gene transfer among closely related streptococci of the mitis group. This results in highly variable mosaic genes encoding the main players in the development of penicillin resistance, the PBPs 2x, 2b, and 1a (for a review, see reference 21). By contrast, PBPs of penicillin-sensitive S. pneumoniae are strictly conserved. Interestingly, a variety of different alleles circulate within the population of S. mitis, and some have been shown to be highly related to mosaic blocks in PBPs from penicillin-resistant S. pneumoniae (PRSP). S. mitis has been identified as the main donor of PBP sequences in PRSP (22–25).

A large number of atypical S. pneumoniae suspected to be S. pseudopneumoniae have been collected over the years at the German National Reference Center for Streptococci (GNRCS) in Germany. One goal of this study was to clarify their species identification by multilocus sequence typing (MLST). Moreover, the mosaic structures of the PBP2x genes were compared with those from a large number of sensitive and resistant commensal streptococci to reveal their relationships to pbp2x in S. pneumoniae and S. mitis, the closest relatives. This gene was chosen as alterations in PBP2x are essential for penicillin resistance, and thus represent a driving force for the evolution of this phenotype. Clusters of related pbp2x sequences were analyzed to reveal group-specific alterations associated with beta-lactam resistance.
RESULTS

Differentiation of *S. pseudopneumoniae* by MLST. Within the large collection of invasive *S. pneumoniae* strains held at the German National Reference Center for Streptococci (GNRCS), 41 atypical isolates were identified that did not react with specific pneumococcal anti-polysaccharide antibodies or showed rough colony morphologies. These isolates were not consistently bile soluble, and optochin resistance was frequently expressed in the presence of CO₂. Moreover, most of them were DOC insoluble and optochin resistant (see Table S1A in the supplemental material). A restriction analysis of PCR-amplified *lytA* with *Ppu21I* and 16SrRNA (see Materials and Methods) resulted in fragments atypical for *S. pneumoniae* in most cases (15). These data suggested that the strains represent *S. pseudopneumoniae*.

Strains that differed from *S. pneumoniae* according to the criteria listed above were further subjected to multilocus sequence typing (MLST) (7) to reveal their phylogenetic relationships to *S. pneumoniae* and *S. mitis*. At least five of the seven genes defined by MLST were amplified from 41 strains (see Table S1B). Genes identified by MLST of *S. pseudopneumoniae* IS7493, whose genome sequence is available (13), were included in further analyses. In addition, we investigated the 38 whole-genome shotgun contigs of *S. pseudopneumoniae* listed in GenBank. Before extracting the seven genes from MLST, we used MLSA to compare *map* and *pyk*, which differentiate *S. pseudopneumoniae* from *S. mitis* (8). Moreover, we compared the *D,D*-carboxypeptidase PBP3 gene, which is also remarkably species specific (10) (see Fig. S1). Only nine out of the 38 *S. pseudopneumoniae* genomes showed 99 to 100% identities for the MLSA loci and >94% identity for the PBP3 gene (only strain 276-03 showed a low identity for *pyk* [93%]) and were considered to be *S. pseudopneumoniae*. Nineteen genomes were defined as *S. mitis* and five each were defined as *S. infantis* and *S. oralis* (see Table S1C).

For the final MLST analysis, all of the *S. pseudopneumoniae* and *S. mitis* were included and compared with MLST data from 29 *S. pneumoniae* and 36 *S. mitis* strains from different parts of the world (22); *S. oralis* Uo5 (26) was used as a reference for this species. Concatenated sequences of six genes were analyzed by the MEGA6 program, including those from 23 atypical strains of this study. The gene *ddl* was omitted as it is frequently highly diverse in penicillin-resistant isolates. The reason is that it maps closely to *pbp2b*, which was acquired by horizontal gene transfer during the evolution of penicillin resistance (27).

The phylogenetic tree clearly differentiated between *S. pneumoniae*, *S. mitis*, and the sole *S. oralis* strain Uo5 (Fig. 1). In addition, one large cluster of *S. pseudopneumoniae* strains was observed that included 21 of the atypical isolates, strain IS7394, and all nine strains identified as *S. pseudopneumoniae* from GenBank. Four atypical isolates were located on two branches between *S. mitis* and *S. pneumoniae* (PS3719, CAP245, PS2113, and PS886). All 19 strains from GenBank identified as *S. mitis* as described above clustered within the *S. mitis* group, as did the four atypical strains RKI1158, RK884, RRR756, and MSR2_Pn407. RRR1136 was found among *S. pneumoniae* (Fig. 1). This strain was shown by a later analysis to express a rare serotype 31, which was not identified in the serotype analysis, and thus had been classified initially as a nontypeable strain. The MLST genes either matched known alleles of *S. pneumoniae* or differed from known alleles in up to 4.6%; only RRR468 contained one allele (*recP*) that is common among *S. oralis*.

The seven MLST loci were also analyzed individually (see Fig. S2 in the supplemental material). The phylogenetic tree generated with *aroE* sequences showed a tight cluster of 40 strains well separated from other branches, and also, the tree generated with *recP* showed one cluster of 30 *S. pseudopneumoniae* strains that included two *S. mitis* (Fig. S2). Smaller clusters of 8 to 26 strains that consisted only of *S. pseudopneumoniae* were obtained with the other six alleles, and one *S. mitis* was found within the *spi* cluster of 12 *S. pseudopneumoniae* strains (Fig. S2). Clustering of *S. pseudopneumoniae* alleles within *S. pneumoniae* was also observed.
MLST tree of strains from this study and reference *S. pneumoniae* and *S. mitis* strain. A neighbor-joining tree was constructed using the concatenated sequences of six MLST loci, excluding *ddl* from 30 atypical *S. pneumoniae* strains from this study, combined with reference *S. mitis* and *S. pneumoniae* from Chi et al. (22). In addition, MLST genes were extracted from the genomes of *S. pseudopneumoniae* IS7394, 9 *S. pseudopneumoniae*, and 19 *S. mitis* listed in GenBank under whole-genome shotgun contigs of *S. pseudopneumoniae* (see Table S1B in the supplemental material). *S. oralis* Uo5 was included as a reference for this species. Bootstrap values (percentages) are based on 1,000 replications. The bar refers to genetic divergence as calculated by the MEGA software.
Forty-two strains (32 atypical strains and the 10 sequenced *S. pseudopneumoniae* strains) contained at least four of the six MLST alleles in *S. pseudopneumoniae*/ *S. pneumoniae* clusters. Strains PS3719 and RRR468 contained two MLST alleles, and strains PS886 and PW2113 contained only one (aroE and/or recP) positioned within *S. pseudopneumoniae*/ *S. pneumoniae* clusters (Table S1B), but they still clustered with *S. pseudopneumoniae* in the MLST analysis that included six alleles (Fig. 1). Further analyses are required to confirm the species of these strains. Finally, the MLST data identified RK11158, RK884, RRR756, MSR2_Pn407, and RRR720 as *S. mitis*, as were the 19 strains from GenBank (Table S1A and S1B).

**Phylogenetic analysis of PBP2x.** One purpose of this part of the study was to see how divergent PBP2x genes are in *S. pseudopneumoniae*. Moreover, the diversity of *pbp2x* in *S. mitis* was analyzed to further our understanding on the distribution of such sequences among *S. pseudopneumoniae* and other SMG and to identify mutations relevant for developing resistance. Sequence information covered all three PBP2x domains to gain more insight into gene transfer events. Unique penicillin-sensitive *pbp2x* alleles for SMG (MIC for penicillin G, <0.05; for cefotaxime, <0.003), not containing any mutations known to be relevant for penicillin resistance, were used as reference sequences for defining mosaic blocks. These include *pbp2x* in *S. pneumoniae* R6 and *S. mitis* M3, and in NCTC10712, which contains large unique sequence blocks predominantly within the transpeptidase domain and flanked by *S. mitis* M3-related sequences as, described recently (23), as well as two unique *pbp2x* alleles in *S. mitis* 658 and SV01 from the Kaiserslautern strain collection. Finally, *pbp2x* in the PRSP clone Spain^23F^-1 was used as a reference for resistant mosaic genes, as it is widespread among penicillin-resistant SMG (22); it contains sequences related to *pbp2x*^M3^-mainly in the transpeptidase domain.

In addition to *pbp2x* of *S. pseudopneumoniae*, we analyzed 38 genes from GenBank listed in Table S1C in *S. mitis* and *S. oralis* from our collection, including nonhuman isolates (10). Recently, eight clusters of *pbp2x* in SMG have been described on the basis of sequences covering the transpeptidase domain (25). We included *pbp2x* sequences representative of three of these clusters that did not match any of the reference *pbp2x* sequences described above, as well as singletons from this study. Furthermore, we searched GenBank for *S. pneumoniae* *pbp2x* that contained blocks of the reference sequences.

A total of 120 *pbp2x* sequences comprising 104 distinct alleles were phylogenetically analyzed using the MEGA6 program. Five clusters were defined that contained one reference sequence each (Fig. 2): *S. pneumoniae* R6 (cluster X-1) and *S. mitis* strains M3 (X-2a), 658 (X-3a), NCTC10712 (X-4), and SV01 (X-5). Sequences of *pbp2x* related to the mosaic *pbp2x* of the PRSP clone Spain^23F^-1 were found in cluster X-2b close to X-2a and consisted exclusively of genes in penicillin-resistant strains, as were those in cluster X-3b, which was close to X-3a. Fourteen *pbp2x* alleles were located on branches outside these clusters. Most *pbp2x* alleles in *S. oralis* formed one group, which included strain ATCC 35037, as did most of the *pbp2x* alleles of *S. infantis*; *pbp2x* alleles in penicillin-resistant strains of these species were found outside clusters or were part of cluster X-2b. In summary, several highly divergent *pbp2x* alleles are apparent in the three species *S. mitis*, *S. infantis*, and *S. oralis*.

**Mosaic structures of *pbp2x*.** The mosaic structures of the *pbp2x* alleles of *S. pseudopneumoniae* in comparison to those of *S. mitis* and *S. pneumoniae* are shown in Fig. 3. *S. oralis* and *S. infantis* *pbp2x* are shown in Fig. 4; only those of *S. infantis* strain JR and *S. oralis* strain ATCC 35037 are included in Fig. 3, as closely related sequence blocks were detected in *S. pneumoniae*. The overall view shows several highly divergent *pbp2x* alleles not only in *S. mitis,* but in *S. infantis* and *S. oralis* as well. Within one cluster, there were mainly sequences of the reference *pbp2x* in addition to sequence blocks of an unknown origin (gray in Fig. 3 and 4). Small sequence blocks of the reference *pbp2x* in *S. mitis* occurred in several clusters. The 14 *pbp2x* alleles which were scattered
FIG 2 Comparative analysis of PBP2x genes. An evolutionary tree was generated from 95 PBP2x sequences using the MEGA software. Clusters X-1 to X-5 were defined according to the reference sequences of *pbp2x* in *S. pneumoniae* R6 (X-1) and those of *S. mitis* strains M3 (X-2b), 658 (X-3a and 3b), NCTC10712 (X-4), and SV01 (X-5). *S. oralis* and *S. infantis* *pbp2x* clusters are also indicated. Cluster X-2b refers to the family of mosaic genes related to *pbp2x* in the *S. pneumoniae* clone Spain23F-1. The species are indicated by different colors. Bootstrap values (percentages) are based on 1,000 replications. The bar refers to genetic divergence as calculated by the MEGA software.
outside the clusters had unique mosaic structures and contained sequences from at least two different sources.

In the four groups related to the *S. mitis* reference *pbp2x*, multiple recombinations were observed that were frequently located within the central transpeptidase/penicillin binding domain (PBD), whereas the 5' and 3'-sequences encoding the N- and C-terminal domains were more uniform. It is also remarkable that in many cases, one distinct sequence block covered the spacer region between the transpeptidase and the C-terminal domain, mimicking the structural organization of the protein. Although most mosaic blocks covered large regions of up to 450 codons, very small regions of divergence of <10 codons were noticeable. *S. pneumoniae* sequences were frequently observed in *pbp2x* in *S. mitis*.

**FIG 3** Mosaic structure of PBP2x variants. Mosaic gene structures were deduced by comparing the reference PBP2x sequences (white arrows on the left side) in *S. pneumoniae* R6 (white sequence blocks), *S. mitis* strains M3 (red), NCTC10712 (yellow), SV01 (light blue), and 658 (green), *S. oralis* ATCC 35037 (dark green), and *S. infantis* JR (orange). Highly similar sequences (≤5% difference) are shown in the same colors; unrelated sequences of unknown origin are in gray. The domain structure and active site boxes of PBP2x are indicated on top; the gray shaded area indicates the central transpeptidase domain. Mutations at sites 338 and 552 are indicated by black arrows. The strains are indicated on the right and are shaded according to species: white, *S. pneumoniae*; light blue, *S. mitis*; gray, *S. pseudopneumoniae*; green, *S. oralis*; orange, *S. infantis*. A pink frame indicates isolates obtained from the GNRCs; four *pbp2x* from Jensen et al. (25) are indicated by an *.
**S. pseudopneumoniae** *pbp2x*. Cluster X-1 consisted almost exclusively of 12 *S. pseudopneumoniae* genes (7 alleles). The four sequences in strains 61-14, 5247, 22725, and 338-14 were identical, as were the *pbp2x* in strains PW2108 and 276-03 (Fig. 3). All of these contained small regions of up to 59 codons in length that differed by 7 to 20% from the R6 *pbp2x*; none of the *S. pseudopneumoniae* alleles were found among *S. pneumoniae* or *S. mitis* in a nucleotide BLAST search.

All of the other *pbp2x* alleles in *S. pseudopneumoniae* were from penicillin-nonsusceptible strains and displayed a variety of mosaic structures. They were found in clusters X-3a/b (RRR475 and RRR472) and X-2b (PW2113, RRR468, and IS7394) or did not cluster (PS3719 and PS15617). Sequences from *S. mitis* NCTC10712, *S. oralis*, or *S. infantis* were not found in these alleles.

**S. mitis** *pbp2x*. Mosaic *S. mitis* PBP2x variants were found in all of the clusters except X-1 (Fig. 2 and 3). In general, the individual makeup of mosaic patterns in *S. mitis* is remarkable, not only in penicillin-resistant isolates, but in penicillin-susceptible isolates as well. There were only a few cases of near sequence identity (>2%): *pbp2x* in *S. mitis* strains 1271, 1272, and DD22 differed from *pbp2x* in *S. mitis* strains 1271, 1272, and DD22 differed from *pbp2x* by ≤15 nucleotides (nt) (0.7%), and *pbp2x* in strain NCTC12261 differed from that of NCTC10712 by 45 nt (2%). Even in cases where the overall mosaic structures were very similar, individual *pbp2x* alleles differed from each other in the lengths of related blocks and the presence of distinct sequences. Examples of this are *pbp2x* in strains 1042, 445, and 1217 (X-4), 578 and 661 (X-3), and MSR2_Pn407 and 888. SV01-related sequences were found in another 14 *S. mitis* strains.
No sequence blocks related to *S. oralis* ATCC 35037 or *S.
infantis* JR were detected. The diversity of 3′ sequences among the 23F-family (X-2b) is of interest, signifying multiple recombination events that aggravate tracking of their evolutionary history.

**S. pneumoniae** *pbp2x*. Sequences from each of the reference *pbp2x* alleles were detected in individual *S. pneumoniae* mosaic PBP2x genes, all of which were from penicillin-resistant isolates (Fig. 3). *S. pneumoniae* sequences related to *pbp2x* in *S. mitis* NCTC10712 (X-4) have been described in serotype 19A *S. pneumoniae* from Hungary (28) and were not included in the present analysis. In most cases, the mosaic structures covered major parts of the PBP2x genes. However, there were two exceptions: *pbp2x* of strain 645.35L98 designated *S. pneumoniae* (29) was completely identical to that of *S. mitis* M3; this is so odd that the species identification is questionable. Moreover, *S. pneumoniae* RRR1136 (group 1) contained only two small sequence blocks of 28 and 124 nt compared with that of *pbp2x*_{23F}, an unusual mosaic structure among resistant *S. pneumoniae* where mosaic blocks are generally much longer. SV18 (X-2b) contained a
large divergent block of unknown origin covering the region encoding the active site S_{337} (see below). PBP2x of *S. pneumoniae* strains 5919 and 14761 were located in group S (X-3), that of strain F1 was in X-4, and four strains contained sequence blocks of *S. mitis* SV01 (X-5) in the 3’-part of the transpeptidase domain (*S. pneumoniae* strain URA4731 from Portugal, F2 and F3 from France, and EU14, member of the clone S.Africa^{19A-7}). *S. pneumoniae* from Japan contained *S. oralis* ATCC 35037 sequences (strains SP00058 and SP00087), and SP00087 (X-2b) contained *S. infantis* JR pbp2x sequences, as well as the regions encoding the N- and C-terminal domains. Taken together, all of the *S. pneumoniae* mosaic PBP2x genes showed signs of multiple gene transfer events involving *S. mitis*, *S. infantis*, and *S. oralis* sequences.

**Altered sites in PBP2x.** PBP2x is a 750-amino acid (aa) multidomain protein with a short N-terminal membrane anchor, an N-terminal domain, and a central transpeptidase domain (residues 266 to 626) followed by linker regions and a C-terminal extension (residues 635 to 750) folded into two PASTA domains (30).

Mutations associated with resistance have only been described within the central transpeptidase domain of the protein, which contains three boxes highly conserved in penicillin-sensitive streptococci: S_{337}TMK with the active site serine, S_{394}SN, and K_{547}TG. PBP2x of the sensitive reference strains *S. mitis* M3, NCTC10712, 658, SV01, *S. oralis* ATCC 35037, and *S. infantis* JR that represent donors for mosaic blocks in PBP2x of PRSP differ from the *S. pneumoniae* R6 PBP2x at 215 sites (28.7%), 66 of which are within the transpeptidase domain. This number includes 23 sites listed in a recent study as being associated with resistance (31), and T_{369} mentioned by Jensen et al. (25) was present in the penicillin-susceptible *S. mitis* 658. These mutations were not considered in the following analysis.

The PBP2x alleles were divided into two groups: class 1, which contained at least one of the two mutations at sites 338 and/or 552 known to confer resistance to penicillin and/or cefotaxime, and class 2, which did not. Residues 338 and 552 are near the active site S_{337} and the K_{547}TG box, respectively, and lead to decreased susceptibility to beta-lactams when introduced in penicillin-sensitive *S. pneumoniae* (32, 33). Not counting alterations that occurred in the reference PBP2x, the number of potentially interesting sites in class 1 PBP2x was reduced to 63 sites: 21 were located within the N-terminal region (265 aa), 38 within the transpeptidase domain (351 aa), and 14 in the C-terminus, including the linker region (134 aa). The numbers of sites that were altered in class 2 PBP2x only were eight in the N-terminal domain, 18 in the transpeptidase domain, and five in the C-terminal extension (see Table S2).

**Mutations at sites T_{338} and Q_{552}**. Thirty-two PBP2x alleles contained mutations at T_{338} to A_{338} (21 strains), G_{338} (*S. pneumoniae* SV18 and *S. mitis* 205), P_{338} (*S. pneumoniae* 5919, *S. mitis* U08, and *S. pseudopneumoniae* RRR472), or S_{338} (*S. pneumoniae* 14761, *S. mitis* strains 578 and 661 in cluster X-3). Four PBP2x alleles contained the E_{552} mutation (*S. pneumoniae* strains F2/F3, URA473, EU14 in cluster X-5, and RRR1136 in X-1). *S. pneumoniae* strains 5919 (X-3) and F1 contained mutations at both sites. Strains harboring these mutations were nonsusceptible to beta-lactams in cases where their MIC values were known, but varied in MICs between 0.03 (*S. pseudopneumoniae* RRR472) and \( > 32 \, \mu g/ml \) (*S. mitis* B6) for penicillin G (mutation at 338), and in the case of mutation E_{552}, between 0.12 (*S. pneumoniae* RRR1136) and 0.5 \( \mu g/ml \) (*S. pneumoniae* EU14) for cefotaxime (Fig. 3; see also Table S2). As cutoff values for resistance were \( \pm 0.25 \, \mu g/ml \) for benzylpenicillin and \( < 0.5 \, \mu g/ml \) for cefotaxime as used in a recent study, T_{338} and E_{552} were reported to occur in penicillin-sensitive strains (25).

An indication for the importance of these mutations is the fact that they are occasionally located on a very small mosaic block, for example, in *S. pneumoniae* RRR1136 (Glu_{552}) and SP00058 (Ala_{338}), suggesting a selective advantage. The sequence block in *S. pneumoniae* RRR1136 (codons 550 to 578) was almost identical to the corresponding regions in *S. pneumoniae* EU14 and *S. oralis* JH, all encoding E_{552}, indicating a common source.
The codon usage of the mutations varied depending on the pbp2x allele and the nature of the mosaic block where the mutation occurred (see Table S3). The codon for T338 is ACT in S. pneumoniae R6 PBP2x, whereas it is ACC in S. mitis strains M3 and 658. All mutations at this site were caused by a single nucleotide (nt) change except for the double mutation G338 in M3-related sequence blocks (ACC to GCC). Similarly, the mutation Q552 was caused by a change from CAG or CAA (Q) to GAG/GAA.

**Other mutations associated with penicillin resistance.** Eight mutations were found in both PBP2x classes: V369, T371, G378, L389, S444, A453, I523, and Q531. Site 597 was changed to S597 (S. mitis RRR720) or D597 (S. mitis 277) in class 2, whereas it was T597 in the class 1 PBP2x of S. mitis B6. D597 also occurs in cefotaxime-resistant laboratory mutants (33, 34).

Strikingly, particular combinations of altered amino acids were observed, and some were specifically associated with one cluster (Table S2). This is especially evident in the X-2b cluster, where the mutations at aa 338 (T338 or G338), F364, T371, K417, S444, T510, and N513 were present in all cases, and T510 and N513 occurred only in this cluster. These mutations were reported to be associated with resistance by many studies (25, 31, 35–39), which is not astounding, given the fact that the 23F family of PBP2x is common and distributed worldwide among S. pneumoniae and SMG (22). L389 and Q531 were associated only with PBP2xM3-related mosaic blocks that included strains in clusters X-2a, X-2b, and X-4. In cluster X-5, only E553 and N568 were noticeable, and one strain (S. pneumoniae EU114) contained Q514 as well, a site not affected in any other PBP2x. T371 plus K531 were associated with S338 (cluster X-5, S. mitis strains 578, 661, and S. pneumoniae 14761) or P338 (S. mitis Uo8 and S. pseudopneumoniae RRR472 in cluster X-3b), but only in one case with A338 (S. mitis 315, cluster X-2a). In cluster X-1, A522 was altered in most of the S. pseudopneumoniae PBP2x.

Noteworthy is that the mutation N568 always occurred in combination with E552, including in PBP2x of S. pneumoniae RRR1136 (cluster X-1). L394 reported in S. pneumoniae isolates with low-level resistance (38) occurred in several class 2 PBP2x, but never in class 1, i.e., always without changes at sites 338 or 552. Sites that are only altered in strains with high-level resistance are 378 (A378 in S. pseudopneumoniae PW2113 and S. pneumoniae SV18 and G378 in S. mitis B6), sites 595 and 599 (L595/W599 in S. mitis B6 and F595/H599 in S. mitis Uo8), and 336 (M336 in S. mitis strains B6 and Uo8). F595 has been reported to be associated with high-level resistance to cefotaxime in S. pneumoniae (37).

Most of the mutations mentioned above are positioned around the enzymatic active site and in the 360 to 394 loop, which affects the stability of this region (30) (see Fig. S3). Surface-orientated mutations include those at positions 322, 444, 493 to 495, 507 to 514, 523, and 531. Interestingly, mutations at sites 417, 424, and 434 are directed toward the noncovalent second beta-lactam located between the transpeptidase and the C-terminal domain (40) (Fig. S3).

**DISCUSSION**

**Phylogeny of S. pseudopneumoniae.** The identification of viridans streptococci species is notoriously difficult due to their genetic competence, resulting in multiple genomic rearrangements and large accessory genomes. Consequently, physiological tests to define species in this group of bacteria remain unsatisfactory, and therefore current approaches to characterize a large number of strains are mainly based on the comparison of DNA sequences derived from housekeeping genes, i.e., by MLST (7) and MLSA (8).

The MLST-based analysis of 41 atypical S. pneumoniae strains and 39 S. pseudopneumoniae genome sequences (strain IS7394 and 38 whole-genome shotgun sequences) defined a total of 35 (atypical collection) plus 10 strains (genomic data) as S. pseudopneumoniae. MLSA of 110 atypical pneumococci obtained from invasive and noninvasive infections in Spain identified 61 strains as S. pseudopneumoniae, and the authors reported on variable phenotypes among these isolates (19). Similar results have been obtained previously using MLST or MLSA with atypical S. pneumoniae. Forty nontype-
able pneumococci from Finland formed a phylogenetic cluster distinct from serotype-
able S. pneumoniae (41). Leegaard et al. described 12 atypical pneumococcal isolates
from HIV-positive patients (42). Only one of these strains clustered with the type strain
S. pseudopneumoniae ATCC BAA-960, which could not clearly be distinguished from S.
mitis by MLSA (8), one represented a truly nontypeable S. pneumoniae, and the
remaining 10 strains formed a subgroup distinct from S. mitis and S. pneumoniae. Some
of the S. pseudopneumoniae strains clustered distinctly from the major group, similar to
the four strains in this study, PW2113, PS886, CAP245, and PS3719 (see Fig. 1).

Some shortcomings of the MLST analysis became apparent in the single locus
analysis (see Fig. S2 in the supplemental material). Only aroE and recP sequences placed
the majority of the S. pseudopneumoniae strains into one cluster, whereas several
smaller clusters were observed with gki, gdh, and spi. A detailed genomic comparison
of a significant number of strains will certainly help to identify more genetic determin-
ants specific for the S. pseudopneumoniae core genome for defining the genetic
relationship between the atypical strains that clustered outside and within the main S.
pseudopneumoniae cluster (Fig. 1). Similarly, features that distinguish members of the
different S. mitis and S. oralis branches remain to be clarified. The gdh and spi MLST
alleles singled out a group of S. mitis strains (see Fig. S2D and S2E). MLST placed the
recently described strains S. tigurinus AZ-3a (43) and S. dentisani 7747 (44) in the IgA
protease-negative S. oralis and the previous S. mitis biovar 2 subcluster, respectively
(10), and genomic comparison suggested three subclusters, namely, S. oralis subsp.
oralis subsp. nov., S. oralis subsp. tigurinus comb. nov., and S. oralis subsp. dentisani
comb. nov., within the coherent phylogenetic clade S. oralis (45). Most important is the
strain collection used for phylogenetic analysis, which ideally should consist of ran-
donically collected isolates from different parts of the world. Isolates were frequently only
collected from diseased patients, from one hospital, from one particular geographic
area, or from a study concentrated on antibiotic-resistant isolates. Given the rapidly
growing number of genomes, this shortcoming will be overcome in the near future.

**PBP2x in S. pseudopneumoniae and close relatives.** The aims of this study were
to see whether PBP genes of penicillin-sensitive S. pseudopneumoniae are highly
conserved, as is the case with S. pneumoniae, and whether they are distinct to alleles
identified in truly penicillin-sensitive viridans streptococci. The PBP2x genes in 12 S.
pseudopneumoniae strains from Germany, Russia, Denmark, Canada, and Spain were
very similar to pbp2x in S. pneumoniae R6 but interspersed with small diverse sequence
blocks of unknown origin, frequently between codons 246 to 268 and 308 to 329. MIC
values of PW2108, CAP156, CAP160, and SK674 were identical to that of S. pneumoniae
R6 (MIC values for the other strains within cluster X-1 were not available). Moreover,
none of the pbp2x alleles in penicillin-sensitive S. pseudopneumoniae were found
among S. pneumoniae or S. mitis genome sequences. Therefore, the mosaic makeup of
pbp2x could serve as a marker for penicillin-sensitive S. pseudopneumoniae. Complex
mosaic pbp2x sequences were found only in S. pseudopneumoniae isolates with MIC
values above those of S. pneumoniae R6, structures which are also typical for penicillin-
resistant S. pneumoniae. Three penicillin-resistant strains, IS7394, PW2113, and RRR468,
contained the typical sequence block of the widespread 23F family of PBP2x genes
(cluster X-2b).

The PBP2x genes in penicillin-sensitive S. pseudopneumoniae strains differed from
that in the S. pneumoniae R6 by 35 to 65 nt (1.8 to 2.5%) or 9 to 16 aa, and from each
other by up to 23 aa (3.1%) or 120 nt (5.3%). This is clearly above the diversity of pbp2x
found among penicillin-susceptible S. pneumoniae isolates, which is almost completely
conserved. We found a maximum of 12 nt encoding 2 aa differences in pbp2x between
the R6 strain and S. pneumoniae INV104 scattered throughout the gene. One exception
appears to be in pbp2x in S. pneumoniae 645.35L98 (29) in that it is identical to pbp2x
in S. mitis M3, but no genomic information was available and the species identification
has not been verified.
On the other hand, the variability of pbp2x sequences in penicillin-sensitive S. mitis is astounding, where variation between the individual alleles ranged between 16 and 22%, and a similar phenomenon was observed in S. infantis and S. oralis, where three and five unique alleles and variants, respectively, were detected (Fig. 4). What is the reason for this apparent reduction of pbp2x variation from S. mitis to S. pseudopneumoniae and further to S. pneumoniae in the gene encoding a major peptidoglycan synthesizing enzyme? There is no indication that the various pbp2x alleles in S. mitis specify subgroups of this species according to the MLST data, and the presence of diverse mosaic genes within sensitive strains suggests that these variations do not result solely from selection pressure imposed by beta-lactam treatment. Since PBP2x is part of a complex protein machinery required for septum synthesis, and thus for cell division (for a review, see reference 46), it would be interesting to study the diversity of genes encoding other cell wall components. Such considerations add to the problem of defining mutations in mosaic PBP2x genes that are associated with resistance as discussed below. We did not consider mutations in the N- or C-terminal domains, as it is generally assumed that only mutations within the transpeptidase domain are important for the expression of resistance. However, they might affect structural features associated with other functions of the proteins or interactions with other partners.

As shown in Fig. 3, the borders of the divergent sequence blocks frequently coincided with the domain structure of PBP2x, i.e., they covered the regions encoding the N- and C-terminal domains and the spacer regions flanking the central transpeptidase domain. Similar observations were obtained with S. pneumoniae R6 transformants obtained with heterologous pbp2x (23). Recombination of small blocks has also been observed by Jensen et al. (25) in S. pneumoniae clones (termed microrecombination) (47) and in S. pneumoniae transformants obtained with S. mitis and S. oralis DNA (48, 49). These data support the conclusion that it is not possible to deduce the number of gene transfer events that contribute to a mosaic gene structure.

The mosaic makeup of pbp2x, especially in penicillin-resistant strains, depends on the strain collection. For example, pbp2x in S. pneumoniae strains from Japan, SP00058, SP00059, and SP00087, display a highly complex mosaic structure. It would be interesting to analyze pbp2x sequences in commensal streptococci from Japan to see whether they are related to the three pneumococcal sequences and represent novel alleles. S. pneumoniae appears to be the most promiscuous species, with genes containing sequence blocks of S. mitis, S. oralis, and S. infantis. The pbp2x 23F family predominates among resistant strains. In fact, this cluster (X-2b) is the only one that includes all five species investigated in the study, confirming the successful global spread of this gene family. No S. mitis alleles apart from S. mitis M3-related sequences were detected in S. infantis or S. oralis in the small number of alleles investigated here.

**PBP2x mutations associated with penicillin resistance.** There have been several approaches for identifying mutations associated with beta-lactam resistance in S. pneumoniae, but, although they resulted in important conclusions, every single study has drawbacks. The mutations identified in beta-lactam-resistant mutants in the laboratory differed from those in clinical isolates (33, 34). On the other hand, a mutation at 338, which can be selected with oxacillin (32, 33), is one of the few that are clearly associated with resistance in many clinical isolates, but is not changed in laboratory mutants. The most important conclusion was that different pathways for modulating PBP2x exist, and that the mutational pattern is related to the selective beta-lactam. A comparison of the structures of PBP2x derived from the clinical isolates from the 23F family (S. pneumoniae 328 and 5204) (30) and a PBP2x identical with S. pneumoniae F2 (cluster X-5) (50) confirmed this notion.

It is important to note that the two mutations near conserved motifs and linked to resistance by experimental and structural evidence, T/G/P338 and E552, mediate only a 1.5-fold increase in the resistance level for penicillin, which is 0.03 µg/ml in the penicillin-sensitive strain S. pneumoniae R6 and up to 2.25-fold for cefotaxime (MIC, 0.02 µg/ml) depending on the mutation (32, 33, 36, 51). Not surprisingly, the MICs of strains...
carrying these mutations varied largely, clearly implicating other sites relevant for the resistance phenotype. Thus, it is impossible to directly correlate a mutation in PBP2x and the MIC of a resistant clinical isolate.

Depending on the strain collection, the mutations identified as being associated with penicillin resistance vary considerably, since different clones, and thus, the frequency of the various mosaic structures of PBP2x, prevail in different geographic areas. PBP2x of the 23F type is common among penicillin-resistant *S. pneumoniae*, *S. mitis*, *S. infantis*, and *S. oralis* (22), and as shown here, also in *S. pseudopneumoniae*. Thus it is not surprising that amino acid changes associated with this PBP2x family (F364, T371, L389, K417, S444, T510, N513, and T605; Fig. 5) are mentioned in many studies (23, 25, 35, 37, 38, 52, 53). However, other mutations listed in these studies frequently included alterations that occur in the penicillin-sensitive reference strains of this study. Two crystal structures of PBP2x of the 23F family are available, namely, for *S. pneumoniae* strain 328 (30) and strain 5204 (35). It was necessary to mutate F364 back to leucine to prevent proteolysis of the protein in the PBP2x328, an indication that this site plays a critical role in the structure of PBP2x. In vitro mutagenesis of *pbp2x*5204 combined with biochemical and microbiological tests singled out six mutations critical for resistance: T371, A338F, F339G, T400, and T605 (35). However, only T371, A338F, and T605 are common mutations, and G384 is frequent in PBP2x variants of penicillin-susceptible *Streptococcus* sp. (Fig. 3; see also Table S2). When these six mutations were introduced into *S. pneumoniae* R6, the PBP2x protein differed by an order of magnitude in acylation efficiency compared with that of the 5204-PBP2x, suggesting a cooperative role of some other substitutions (35). Cooperativity of alterations might explain why the reversal of G384, common in penicillin-susceptible strains, decreased benzylpenicillin and cefotaxime MICs in a *S. pneumoniae* with high-level resistance (37).

Most importantly, the mutational makeup of mosaic PBP2x variants differed between the PBP2x groups. This supports the notion of distinct mutational pathways for PBP2x, but puts this conclusion in the context of the parental PBP2x variants of sensitive strains. Given the fact that the amino acid sequences of PBP2x in penicillin-
susceptible *S. mitis* vary by up to >22%, it is conceivable that certain mutations are allowed only in the particular structural context of individual PBP2x. For example, A$_{338}$ was mutated to S$_{338}$ or P$_{338}$ in cluster X-3, whereas mutations A$_{338}$ or G$_{338}$ were associated with X-2a/2b, and E$_{552}$ was frequent in cluster X-5. A comparison of structural data available from a PBP2x identical to that of strain *S. pneumoniae* F2 (cluster X-5) (50) with PBP2x of the 23F family (30, 35) fully supports alternative mechanisms of penicillin resistance.

N$_{386}$, which always occurred in combination with E$_{552}$, is also an active-site mutation with its side chain lying across from that of E$_{552}$ as deduced from the PBP2x structure identical to PBP2xF$_2$ (X-5) (50). The mutations K$_{117}$, S$_{424}$, and S$_{434}$ are located between the between the transpeptidase and the C-terminal domains near the second noncovalent cefuroxime molecule (40), similar to mutations in laboratory mutants (54), supporting the importance of the C-terminal domain for antibiotic and, most likely, substrate recognition. Some mutations proposed to play a role in resistance, and which also have structural effects on the active site, are present in sensitive *S. mitis*, including M$_{447}$ and A$_{449}$, signature residues for PBP2x of cluster X-5 (Fig. 4), and G$_{384}$. Such changes must affect the interaction with the actual substrate as well, suggesting different enzymatic properties of PBP2x variants already in penicillin-susceptible streptococci.

In conclusion, penicillin-sensitive *S. pseudopneumoniae* strains contain particular *pbp2x* alleles distinct from those of the close relatives, *S. pneumoniae* and *S. mitis*. PBP2x genes in resistant *S. pseudopneumoniae* display complex mosaic structures that are typical for resistant strains of other viridans streptococci. PBP2x sequences of four penicillin-sensitive *S. mitis* isolates account for most of the diverse sequence blocks in resistant *S. pneumoniae* and *S. mitis*, and are widespread among sensitive *S. mitis* as well. Altered sites associated with resistant strains revealed specific patterns depending on the parental sequence block, in agreement with distinct mutational pathways during the development of beta-lactam resistance. Several sites have been shown to affect acylation kinetics to beta-lactam antibiotics, and it is conceivable that the proteins display slightly different enzymatic properties with their actual substrate, which still needs to be characterized. Characterization of the mosaic structure of PBPs in penicillin-resistant isolates by comparison to parental genes of sensitive strains is not only helpful but probably essential for identifying mutations related to resistance, which needs to be evaluated by further genetic, biochemical, and structural data. Given this scenario, it is unlikely that any novel antibiotic will target all PBP2x variants equally well, unless by a mechanism completely distinct from that of beta-lactams.

### MATERIALS AND METHODS

**Characterization of bacterial strains and antibiotic susceptibility testing.** Forty-one strains isolated between 1997 and 2011 from the German National Reference Center for Streptococci (GNRCS) were included in the present study (see Table S1A in the supplemental material). They were characterized as atypical *S. pneumoniae* by serotyping and optochin and bile (deoxycholate) susceptibility, and were tested for the presence of the capsular biosynthesis gene *cpsA*.

Pneumococcal isolates were serotyped by the Neufeld Quellung reaction using type and factor sera provided by the Statens Serum Institute, Copenhagen, Denmark. LytA PCR/restriction was performed as described previously (15). PCR amplification of 16S rRNA genes and subsequent cleavage with the restriction endonuclease BsiHKAI was performed according to published procedures (55).

Identification by MLST was performed as described previously (7) (Table S1B). In addition, the species from 38 genomes available in GenBank and listed under “whole-genome contigs/*S. pseudopneumoniae*” were characterized by *map* and *pyk*, which differentiate *S. pseudopneumoniae* from *S. mitis* in MLSA (8), and by *pbp3*, which differentiates streptococcal species. Those genomes identified as *S. mitis* and *S. pseudopneumoniae* were further characterized by MLST, and details are listed in Table S1B.

All strains were tested for antibiotic MICs using the broth microdilution method as recommended by the CLSI (56). The microtiter plates (Sensititre NLMCS10; TREK Diagnostic Systems, Ltd., East Grinstead, UK) contained penicillin G (PEN), cefotaxime (CEF), clarithromycin/erythromycin (CLA/ERY), clindamycin (CLI), tetracycline (TET), levofloxacin (LEV), chloramphenicol (CHL), and trimethoprim-sulfamethoxazole (SXT) with cation-adjusted Mueller-Hinton broth (Oxoid, Wesel, Germany) and 5% lysed horse blood. Oxacinil susceptibility was tested using 1-μg discs (Oxoid, Wesel, Germany). In some isolates, MICs for beta-lactams were additionally specified using narrow dilutions of the antibiotic as described previously (23).
DNA isolation and PCR amplification. Chromosomal DNAs from streptococci were isolated as described previously (23). PCR products were purified using a JetQuick DNA purification kit (Genomic). PCRs were performed using either Goldstar Red Taq polymerase (Eurigenotec) or DreamTaq polymerase (Fermentas) according to the manufacturer’s instructions. The oligonucleotides used in this study were obtained from Eurofins. PBP2x gene fragments were amplified with the primers pn2xup and pn2xdown (23), and direct sequencing of PCR products was performed with consecutive primers.

Bioinformatic tools and analysis. Neighbor-joining trees were generated with MEGA6.06 (57) and Clustal alignments using standard parameters. Bootstrap analysis was based on 1,000 replicates. PBP2x sequences were aligned by ClustalX2 (58) and further processed by Gedeneroc (http://www.psc.edu/biomed/genedoc). PBP2x gene sequences were aligned with the each of the reference sequences. Codon sites were included manually and trimmed by the program Clustal Formatter (http://nbc11.biologie.uni-kl.de/) to reveal only sites that differ from the reference sequence as shown in Table S2. Sequence blocks that differ by <5% are defined as distinct sequences as shown by different colors in Fig. 3 and 4.

Accession number(s). Accession numbers of pbp2x sequences are KY292535, KY292538, KY292539, KY292542, KY292549 to KY292551, KY292559, KY292560, KY292564, AJ238585, KY292528, KY292530, KY292531, KY292533, KY292534, KY292536, KY292540, KY292541, KY292543, KY292544, KY292548, KY292552 to KY292557, and KY292561 to KY292563 and are listed in Table S2.

SUPPLEMENTAL MATERIAL
Supplemental material for this article may be found at https://doi.org/10.1128/AAC.02646-16.

SUPPLEMENTAL FILE 1, XLSX file, 0.1 MB.
SUPPLEMENTAL FILE 2, XLSX file, 0.1 MB.
SUPPLEMENTAL FILE 3, XLSX file, 0.1 MB.
SUPPLEMENTAL FILE 4, PDF file, 1.2 MB.

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The authors have no conflict of interest to declare.

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