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Molecular and structural analysis of mosaic variants of penicillin-binding protein 2 conferring decreased susceptibility to expanded-spectrum cephalosporins in *Neisseria gonorrhoeae*: role of epistatic mutations†

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Abstract

Mutations in penicillin-binding protein 2 (PBP 2) encoded by mosaic *penA* alleles are critical for intermediate resistance to the expanded-spectrum cephalosporins ceftriaxone and cefixime in *Neisseria gonorrhoeae*. Three of the ~60 mutations present in mosaic alleles of *penA*, G545S, I312M, and V316T, have been reported to be responsible for increased resistance, especially to cefixime (Takahata *et al.* 2006. *Antimicrob Agents Chemother* 50:3638-45). However, we observed that the minimum inhibitory concentrations (MICs) of penicillin, ceftriaxone, and cefixime for a wild type strain (FA19) containing a *penA* gene with these three mutations increased only 1.5-, 1.5-, and 3.5-fold, respectively. In contrast, when these three mutations in a mosaic *penA* allele (*penA35*) were reverted back to wild type and the gene transformed into FA19, the MICs of the three antibiotics were reduced to near wild type levels. Thus, these three mutations display epistasis, in that their capacity to increase resistance to β -lactam antibiotics is dependent on the presence of other mutations in the mosaic alleles. We also identified an additional mutation, N512Y, that contributes to decreased susceptibility to expanded-spectrum cephalosporins. Finally, we investigated the effects of a mutation (A501V) currently found only in non-mosaic *penA* alleles on decreased susceptibility to ceftriaxone and cefixime, under the expectation that this mutation may arise in mosaic alleles. Transfer of the mosaic *penA35* allele containing an A501V mutation into FA6140, a chromosomally mediated penicillin-resistant isolate, increased the MICs of ceftriaxone (0.4 μ g/ml) and cefixime (1.2 μ g/ml) to levels above their respective breakpoints. The proposed structural mechanisms of these mutations are discussed in light of the recently published structure of PBP 2.

Neisseria gonorrhoeae is the etiologic agent of the sexually transmitted infection gonorrhea. In 2007, there were over 350,000 infections reported in the United States (1). Gonococcal

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Supporting Information Available

Alignment of wild type PBP 2 with PBP 2^{35/02} showing the location of mutations in each module. This material is available free of charge via the Internet at <http://pubs.acs.org>.

infections are often asymptomatic, especially in females, and if left untreated can cause pelvic inflammatory disease and disseminated infections, as well as contribute to the spread of the disease. Because of the lack of a vaccine, antibiotics are the primary treatment for gonococcal infections.

Traditionally, gonorrhea was treated with penicillin G, but in 1986 this antibiotic was discontinued in the USA due to the onset of resistance. For the same reason, tetracyclines were also withdrawn. More recently, high-level resistance to fluoroquinolones led to the withdrawal of these antibiotics from the recommended list in the USA, leaving only expanded-spectrum cephalosporins such as ceftriaxone and cefixime as reliable treatments for gonococcal infections (2). Importantly, strains with decreased susceptibility to these antibiotics (also called ceph^I strains) are becoming widespread in the community (3,4), making it crucial that we understand the mechanisms that lead to ceph^I resistance so that new antibiotics and new approaches be found for treating gonococcal infections before we are left with no effective antibiotics (4). This need is highlighted by recent reports of treatment failures using oral expanded-spectrum cephalosporins in Japan and Hong Kong (5,6), as well as two confirmed treatment failures of pharyngeal gonococcal infections with ceftriaxone (7), although the latter two cases probably reflect the well-recognized difficulties in eradicating pharyngeal gonorrhea (4).

The most common mechanism of penicillin resistance in *N. gonorrhoeae*, termed chromosomally mediated resistance, involves at least 5 resistance determinants (8,9). These determinants, which are mutated forms of normal genes/loci, can be transferred from a high-level resistant donor to a susceptible strain by homologous recombination and selection, with transfer occurring in a defined order (8,10,11). The first step is transfer of *penA*, which encodes altered forms of penicillin-binding protein 2 (PBP 2), the lethal target of penicillin G in *N. gonorrhoeae* (12). The second determinant transferred is *mtrR*, which leads to increased efflux pump expression and activity (13). The third determinant transferred is *penB*, which encodes altered forms of the major outer membrane porin, PorB_{1B} (14,15). The final steps, which result in high-level penicillin resistance, include acquisition of *ponA*, encoding an altered form of PBP 1, and a resistance determinant whose identity is unknown (8). The sum effect of these determinants is to increase the minimum inhibitory concentration (MIC) of penicillin by 400-fold (from 0.01 µg/ml to 4 µg/ml).

The major difference between penicillin-resistant and ceph^I strains is the presence of mosaic *penA* alleles that encode PBP 2 variants containing up to 60 amino acid changes compared to PBP 2 from wild type strains. We and others have proposed that the rapid emergence of ceph^I strains has occurred by horizontal transfer of mosaic *penA* genes, which were generated by recombination events between *N. gonorrhoeae* and commensal *Neisseria* species, into chromosomally mediated penicillin-resistant strains already harboring the necessary determinants to increase resistance to intermediate levels (16,17).

Takahata *et al.* have reported that three mutations in mosaic variants of PBP 2, G545S, I312M, and V316T, are responsible for decreased susceptibility to cefixime (18). However, when we incorporated these mutations into a wild type *penA* gene and transformed the gene into FA19, resistance to penicillin, ceftriaxone, and cefixime increased only marginally (≤ 3 -fold), putting the original conclusion in doubt (see Results). Therefore, we set out to identify important regions of PBP 2, and amino acids within them, that confer resistance to penicillin and expanded-spectrum cephalosporins. Our data indicate that more complex and subtle mechanisms are at play; that is, mosaic PBP 2 variants display epistasis, in which the three residues identified by Takahata *et al.* (18) are important in decreasing the rate of acylation, but only in the presence of other residues that have little to no apparent effect on their own. We also find that if an A501V mutation, which has been observed recently in non-mosaic

penA alleles (19–21), were to emerge in mosaic *penA* alleles, the MICs of ceftriaxone and ceftriaxone would increase to above the established break points of resistance.

MATERIALS AND METHODS

Strains and Plasmids

The strains and plasmids used in this study are shown in Table 1. FA19 is a penicillin- and cephalosporin-susceptible strain that served as the recipient strain for most of the studies described herein (22). In the experiments examining the effect of the A501V mutation, FA6140, a penicillin-resistant, cephalosporin-susceptible strain (23) that contains all of the known resistance determinants (i.e., *penA*, *mtrR*, *penB*, and *ponA*) as well as the unknown determinant, also served as a recipient strain. pUC18us-*penA* and pUC18us-*penA35* contain the wild type *penA* gene from FA19 and the mosaic *penA* gene from the cep^I strain 35/02 (24), respectively, along with 300 bp of downstream sequence and an uptake sequence to facilitate homologous recombination. To construct the different chimeric PBPs, silent restriction sites were introduced into the coding sequences of pUC18us-*penA* and pUC18us-*penA35* by the QuikChange method (Stratagene, Carlsbad, CA). These sites allowed us to swap out individual cassettes from *penA35* with the corresponding cassettes from *penA* to create the “-mod” constructs (Fig. 1, Supplemental Fig. 1). Other point mutations were introduced into the appropriate constructs by overlap extension PCR (25). All constructs were verified by sequencing before proceeding with transformation experiments.

Transformation

Transformation of the chimeric and mutant constructs into FA19 and FA6140 was accomplished as described previously (8). Transformants were selected on GCB plates containing various amounts of antibiotics just above their respective MICs. To verify correct recombination, transformants were passaged on GCB plates, and the next day, colonies were boiled in 30 μ l water, spun briefly to pellet cell debris, and the supernatants were used as templates in PCR with the appropriate *penA* primers. PCR products were sequenced by the UNC sequencing facility or by Eton Bioscience Inc. (Research Triangle Park, NC).

MIC measurements

The MICs for penicillin, ceftriaxone, and cefixime were determined exactly as described previously (16). The antibiotics were tested in ~1.5-fold increments to increase the accuracy of the MIC determination. At least two (and often up to 4) colonies from each transformation were tested, each verified by PCR amplification and sequencing as described above. At least three independent MIC experiments were carried out, and the MICs reported represent the averages of all experiments. Error bars indicate the standard deviation of the determinations.

Purification of PBP 2 variants

PBP 2, PBP 2^{35/02}, and PBP 2^{35/02}-A501V were purified as described previously (26). Briefly, the genes encoding each variant were cloned into derivative of pMAL-C2 (New England Biolabs, Beverly, MA), pMAL-C2KV, which fuses His₆-maltose-binding protein and an intervening tobacco etch virus (TEV) protease site to amino acid 44 of PBP 2, and the proteins were expressed in *E. coli*. The fusion proteins were purified on a Ni²⁺-NTA column (GE Healthcare, Piscataway, NJ), cleaved with His₆-tagged tobacco etch virus (TEV) protease, and the digests were re-chromatographed over a Ni²⁺-NTA column. The purified proteins were eluted in buffer containing 15 mM imidazole, while uncleaved protein, His₆-TEV and His₆-maltose-binding protein were eluted with 250 mM imidazole.

The proteins were dialyzed to remove imidazole, concentrated to ~ 6 mg/ml, and frozen at -80°C .

k_2/K_S measurements of the rate of acylation by β -lactam antibiotics

The reaction of β -lactam antibiotics with PBP 2 is denoted by the following equation:

$E+S \xrightleftharpoons{K_S} E \bullet S \xrightarrow{k_2} E-S' \xrightarrow{k_3} E+P$, where $E \bullet S$ is the non-covalent enzyme-antibiotic complex, $E-S'$ is the acyl-enzyme complex, and P the hydrolyzed antibiotic. k_2/K_S constants, which are a direct measure of the ability of an antibiotic to inhibit a PBP (27), were calculated from first order rates of acylation of purified, soluble PBP 2 variants by [^{14}C]penicillin G (Moravek, Brea, CA) as previously described (26,28). Graphs of PBP 2- [^{14}C]penicillin G complex formation versus time were obtained by incubating 27 μg of protein with 1.0 μM [^{14}C]penicillin G, and aliquots of ~4 μg were removed at 15 sec intervals, precipitated with 5% trichloroacetic acid, filtered over Whatman GC-A filters, and the filters were submitted to scintillation counting. The concentration of [^{14}C]penicillin G was increased to 25 and 50 μM for determination of k_2/K_S values with PBP 2^{35/02} and PBP 2^{35/02}-A501V, respectively. The k_2/K_S values of non-radioactive cephalosporin antibiotics were obtained in competition experiments with [^{14}C]penicillin G using the following

equation: $(k_2/K_S)_{ceph} = (k_2/K_S)_{penG} \left(\frac{[penG]}{[ceph]_{0.5}} \right)$, where $[penG]$ is the concentration of [^{14}C]penicillin G used in the reaction and $[ceph]_{0.5}$ is the concentration of cephalosporin antibiotic that inhibits the binding of [^{14}C]penicillin G by 50% (27).

RESULTS

Analysis of mosaic PBP 2 mutations I312M, V316T, and G545S in $ceph^I$ resistance following incorporation into a wild-type $penA$ background

Takahata *et al.* reported that three amino acid mutations found in mosaic PBP 2 variants, I312M, V316T and G545S, are responsible for decreased susceptibility to cefixime in *N. gonorrhoeae* (18). To confirm these data, we introduced a single mutation (G545S), a double mutation (I312M/V316T), or all three mutations (G545S/I312M/V316T) into the wild type $penA$ gene from FA19, and used these plasmids to transform FA19 to increased penicillin G or cefixime resistance. No transformants could be isolated with either the single or double mutants, presumably because they did not confer an increase in resistance. However, we were able to select for transformants harboring the triple mutant, but the MICs of penicillin, ceftriaxone, and cefixime for these transformants increased by only 1.5-, 1.5-, and 3.5-fold, respectively, compared to 12-, 20-, and 100-fold increases, respectively, with the full $penA_{35}$ gene (16). Thus, it appears that together these three mutations confer only minimal increases in resistance to these antibiotics when incorporated into the $penA$ gene from FA19.

Construction and analysis of PBP 2 chimeras

These data indicated that the mechanisms leading to emergence of $ceph^I$ strains are more complex than originally reported, and thus we constructed chimeras between wild type and mosaic $penA$ genes to identify the regions and amino acids that are important in decreasing the rate of acylation by these antibiotics. As shown in Fig. 1, six regions in $penA_{35}$ bounded by silent restriction sites, termed “modules” and denoted mod0 through mod5, were replaced by their corresponding modules from $penA$, and *vice versa*, and these constructs were used to transform FA19 to increased cefixime or penicillin resistance. Each module contained between 3 and 15 amino acid changes, with mod0 comprising the entire N-terminal domain of PBP 2 and mods 1–5 covering the C-terminal transpeptidase/ β -lactam-binding domain (Figs. 1, S1). No transformants could be selected when individual

penA35 modules were transferred into wild type *penA* in an attempt to show gain-of-function, indicating that no single region contained changes that were capable of conferring increased resistance to the antibiotics.

In contrast, we were able to select for transformants of FA19 with *penA35* constructs in which individual mosaic *penA35* modules were replaced with their wild type counterpart in *penA* (Fig. 2). In all cases, the *penA35/penA* chimeras resulted in lower levels of resistance relative to *penA35*, and except for a few instances, the MICs of the three antibiotics (compared to *penA35*) decreased similarly for each chimera. For clarity, the different chimeric constructs are defined in terms of the fold decrease in MIC compared to that conferred by *penA35*.

Replacing modules 0 or 2 of *penA35* with wild type modules had little to no effect on the MICs of the three antibiotics, while replacing modules 1, 3 or 4 of *penA35* with the corresponding wild type module decreased MICs from 2- to 3-fold compared to *penA35* (Fig. 2). Replacing modules 1 or 4 of *penA35* resulted in a greater decrease in the MICs than replacement of module 3, whereas replacement of modules 3 and 4 together decreased the MICs of the three antibiotics slightly more than replacement of the individual modules. Despite repeated attempts, we could not select transformants in which module 5 of *penA35* was replaced by wild type, most likely because this chimera did not increase the MIC of FA19 above that conferred by wild type *penA*. These data indicate that mutations within module 5 are the most critical for decreased susceptibility, whereas mutations in modules 1 and 4 are important but less so than those in module 5.

Role of G545S and I312M/V316T mutations in cep^I resistance

Importantly, modules 1 and 5 contain the I312M/V316T and G545S mutations, respectively, initially described by Takahata *et al.* (18). To examine the role of the G545S mutation in resistance, we incorporated this mutation back into the $-mod5$ chimera (i.e. $-mod5 + G545S$). Unlike the parent $-mod5$ construct, which did not confer resistance over wild type levels, the resulting transformants were only slightly less resistant to penicillin and ceftriaxone compared to those containing the unmodified *penA35* gene (Fig. 3), demonstrating the importance of this mutation. Resistance to cefixime, however, was nearly 3-fold lower, indicating that the G545S mutation is less important for cefixime resistance and that one or more additional mutations within module 5 appear to be required to reach the level of *penA35*. Overall, these data indicated that G545S was the most important mutation within *mod5* for increasing resistance, but the degree to which the mutation is responsible for resistance depended on the antibiotic being examined.

To examine the role of the I312M and V316T mutations in resistance, we replaced module 1 within the $-mod5 + G545S$ chimera with the wild type sequence of *penA* ($-m1,5 + G545S$). The MICs of all three antibiotics for the resulting transformants decreased by ~2-fold compared to $-mod5 + G545S$ (Fig. 3). Incorporation of the I312M/V316T double mutation back into this construct (i.e. $-m1,5 + I312M/V316T/G545S$) restored the MICs of ceftriaxone and cefixime in these transformants to the same level as $-mod5 + G545S$, whereas the MIC of penicillin did not change. These data suggest that all of the decrease in the MICs of ceftriaxone and cefixime caused by replacement of module 1 of *penA35* with wild-type sequence was due to the I312M/V316T mutations, while the drop in penicillin MIC was due to other residue(s) within the module.

Reversion of G545S and I312M/V316T mutations in mosaic *penA35*

Reversion of the G545S mutation in *penA35* back to wild type decreased the MICs of penicillin, ceftriaxone, and cefixime for the resulting transformants 2-, ~6- and 8-fold,

respectively, compared to *penA35*, whereas reversion of the I312M/V316T mutations to wild type decreased MICs of penicillin, ceftriaxone, and cefixime by 2-, 4- and 4-fold, respectively. When all three mutations were reverted to wild type, the MICs of penicillin, ceftriaxone, and cefixime were decreased by 2.5-, 16-, and 25-fold, respectively, relative to *penA35*. Importantly, the MICs of ceftriaxone and cefixime for the triple reversion mutant were nearly identical to that of FA19. These data are consistent with the idea that the three mutations are important in resistance, but only in the context of some or all of the other 55 mutations found in mosaic *penA* alleles.

Role of module 4 amino acids in antibiotic resistance

Replacement of module 4 from *penA35* with wild type sequence resulted in a ~3-fold decrease in the MIC of the three antibiotics when the chimera was transformed into FA19 (Fig. 2). This result was intriguing, since module 4 has only three amino acid changes, F504L, A510V, and N512Y, the first two of which are also found in the non-mosaic *penA4* allele from FA6140 (26), a high-level, penicillin-resistant but cephalosporin-susceptible strain (8,26). These three mutations are clustered on the β 3- β 4 loop of PBP 2, which is just C-terminal to the KTG active site motif. We therefore reverted the three changes individually in *penA35* back to the wild type sequence, and determined the MICs of penicillin, ceftriaxone, and cefixime of the resulting transformants.

Surprisingly, the individual mutations had different effects on the MIC depending on the antibiotic being examined. For the expanded-spectrum cephalosporins, the most important reversion was Y512N, which was responsible for most if not all of the decrease in the MICs observed in strains containing *penA35* –mod4. For penicillin, the Y512N mutation had a minor effect on resistance; instead, both the L504F and V510A reversions decreased resistance to the same level as the –mod4 construct (Fig. 4). These data highlight the importance of the β 3- β 4 loop for the reactivity of β -lactam antibiotics toward PBP 2 and demonstrate the differential effects of these mutations on the different antibiotics.

Effects of the A501V mutation in PBP 2^{35/02}

We also examined the effects of the A501V mutation in both mosaic and non-mosaic *penA* genes by incorporating this mutation into the *penA4* and *penA35* genes and transforming these genes into FA19 and FA6140 (Fig. 5). When transformed into FA19, the *penA4*-A501V and *penA35*-A501V mutant alleles decreased the MIC of penicillin by 20% and 50%, respectively, compared to levels conferred by the parental alleles (Fig. 5a). In contrast, *penA4*-A501V and *penA35*-A501V both increased ceftriaxone and cefixime MICs between 2-fold and 4-fold above those conferred by *penA4* and *penA35*, respectively. In general, the effects of the A501V mutation were greater in the *penA35* background than in the *penA4* background.

Consistent with our results in FA19, when the *penA35*-A501V gene was transformed into FA6140, the MIC of penicillin for FA6140 *penA35*-A501V was nearly half of that for FA6140 *penA35* (Fig. 5b), whereas the MICs of both ceftriaxone and cefixime increased by over 2-fold. Importantly, the MICs of ceftriaxone and cefixime for the resulting strains, 0.4 and 1.2 μ g/ml, respectively, are well above the breakpoints for “resistance” (>0.25 μ g/ml for both antibiotics). These data suggest that emergence of this mutation in mosaic *penA* genes, which to date has not occurred, could render both ceftriaxone and cefixime ineffective for treating gonococcal infections.

k_2/K_s constants of wild type PBP 2, PBP 2^{35/02}, and PBP 2^{35/02}-A501V

To compliment our MIC data, we determined the acylation rate constants of penicillin, ceftriaxone, and cefixime for purified wild type PBP 2, PBP 2^{35/02}, and PBP 2^{35/02}-A501V.

For wild type PBP 2, [¹⁴C]penicillin G had a k_2/K_S of 76,000 M⁻¹s⁻¹, whereas ceftriaxone and cefixime displayed k_2/K_S constants that were markedly higher (1.7×10^6 and 1.5×10^6 M⁻¹s⁻¹, respectively; Table 2), consistent with their low MICs for wild type gonococcal strains. The acylation rates of penicillin G, ceftriaxone, and cefixime for PBP 2^{35/02} decreased 150- to 200-fold compared to wild type, underscoring the marked effect of the multiple substitutions in mosaic PBP 2 on the kinetics of β -lactam binding. The acylation rate of penicillin for PBP 2^{35/02}-A501V increased ~2.7-fold compared to PBP 2^{35/02}, while the k_2/K_S of cefixime decreased by 2.3-fold (Table 2). Surprisingly, the rate for ceftriaxone increased 1.8-fold, opposite of what we expected based on MIC experiments. Except for this latter result, these data are consistent with the changes observed in strains transformed with the mosaic *penA* allele.

Discussion

Takahata *et al.* reported that three amino acid mutations found in mosaic PBP 2 variants, I312M, V316T and G545S, are responsible for decreased susceptibility to cefixime in *N. gonorrhoeae* (18). In that study, the authors determined the MICs of a range of expanded-spectrum cephalosporins for transformants of FA1090 harboring *penA*^{FA1090} alleles containing either the G545S mutation alone or G545S together with either the I312M or V316T mutation (a strain containing all three mutations could not be isolated). The MICs of ceftriaxone and cefixime increased 2-fold for the G545S transformant and 4-fold and 8-fold, respectively, for the G545S/I312M or G545S/V316T double transformants. From these data, the authors concluded that these three mutations were responsible for the decreased MICs conferred by mosaic *penA* alleles.

However, when we incorporated the three mutations into the wild type *penA* allele from FA19 and transformed it into FA19, the MICs of ceftriaxone and cefixime increased by only 1.5- and 3.5-fold, respectively, well below the levels obtained with the full mosaic *penA*³⁵ gene (Fig. 2). An important difference between our work and the study by Takahata *et al.* is the choice of the parental *penA* allele. Whereas we used *penA* from FA19, which we consider to be a true “wild type” allele (22), Takahata *et al.* (18) used *penA* from FA1090, which contains an Asp345a insertion that decreases the k_2/K_S acylation rate constant of penicillin G by 6-fold (26). Presumably, the increased levels of resistance observed by Takahata *et al.* were due to the Asp-345a insertion adding to or amplifying the effects of the mutations. Since the Asp-345a insertion is not present in mosaic *penA* alleles, the *penA* gene from FA1090 is not an optimal background to examine the effects of these mutations.

While the three mutations (I312M, V316T, and G545S) do not increase resistance markedly when incorporated into a wild type *penA* gene, they have a striking effect on resistance when reverted back to the wild type in the mosaic *penA*³⁵ background, indicating that these mutations require at least some of the other amino acid changes found in 35/02 to decrease susceptibility. This phenomenon, referred to as epistasis, has been elegantly described by Thornton and colleagues (29,30), who identified several “permissive” mutations in steroid hormone receptors that, while having no functional importance on their own, stabilized function-switching mutations and facilitated the evolution of new steroid hormone-binding activity. Weinreich *et al.*, studying the evolution of hydrolytic activity against cefotaxime in TEM β -lactamase, have described a similar epistatic phenomenon (31). These authors showed that only a few pathways to full resistance are permissible, since some mutations do not increase cefotaxime resistance in certain allelic backgrounds. In an example involving PBPs, Hedge and Spratt (32) defined the steps for acquisition of resistance of *E. coli* to a range of cephalosporins through serial mutagenesis of PBP 3 (the functional equivalent of PBP 2 in *N. gonorrhoeae*). The authors were able to obtain high-level resistance to cephalosporins in four steps. Importantly, two of the steps had little effect on resistance, but

increased protein stability and thermostability of PBP 3 and allowed for subsequent isolation of resistance-conferring mutations (32).

To suggest how the three mutations described in this study might impact the rate of acylation, we examined our recent crystal structure of PBP 2 from *N. gonorrhoeae*, solved in the apo form. G545S, which is the most important mutation for resistance in mosaic *penA* alleles, is present at the start of the $\alpha 11$ helix, one of the two helices that pack on top of the 5 anti-parallel β strands typical of penicillin-interacting proteins. The main chain amides of G545 and G546 are within hydrogen-bonding distance to the side chain hydroxyls of Thr498 and Thr500, respectively, located within the KTG(T) active-site motif (Fig. 6A). By analogy with other PBPs such as *E. coli* PBP 5, the main chain amide of Thr500 is predicted to stabilize the oxyanion hole for the transition state, and so one effect of the G545S mutation might be to lower acylation by compromising the geometry of the transition state/tetrahedral intermediate. Alternatively, because the hydroxyl side chains of the equivalent residues to Thr498 and Thr500 in *S. pneumoniae* PBP 2x (Ser548 and Thr550) interact with the β -lactam carboxylate in the covalent complex, alteration of these contacts may be another mechanism that lowers acylation. Ile312 and Val316 are located on the opposite side of the helix as Ser310 and Lys313 of the SxxK motif and pack into a hydrophobic pocket (Fig. 6B). Mutation to larger (I312M) or more hydrophilic (V316T) side chains might disrupt these interactions such that the position of the SxxK motif is altered, leading to decreases in acylation rates with β -lactam antibiotics. A similar argument has been put forth for the M339F mutation in *S. pneumoniae* PBP 2X (33).

In addition to the three mutations discussed above, we also showed that reversion of N512Y in *penA35* decreases resistance to ceftriaxone and cefixime by 2-fold without affecting penicillin resistance. This result is consistent with its emergence in mosaic *penA* alleles and its relative absence in non-mosaic *penA* alleles found in penicillin-resistant strains (18,19,21). Asn512 is relatively distant from the active site (Fig. 6B) and the exact impact of the N512Y mutation on acylation is difficult to predict, but it is located on the $\beta 3$ - $\beta 4$ loop that contains mutations that are known to be important for resistance to penicillin in non-mosaic forms of PBP 2 (26). One possibility is that such a mutation perturbs the architecture of the KTG motif of $\beta 3$.

Our work has also highlighted the impending danger of the A501V mutation arising in mosaic *penA* alleles, because if it happens this would increase the MICs of ceftriaxone and cefixime above their breakpoints for resistance. The alteration of A501 in PBP 2 appears to be a gonococcal-specific alteration, as it has not yet been observed in commensal *Neisseria* species, which suggests that it arose through mutation in response to selective pressure with expanded-spectrum cephalosporins instead of by transformation. Indeed, Takahata *et al.* (18) reported the isolation of a spontaneous A501V mutation during transformation experiments. Thus, it may be a matter of when and not if mutations in Ala501 arise in mosaic *penA* genes. Ala501 resides on the $\beta 3$ - $\beta 4$ loop where mutation of the methyl side chain to the bulkier side chain of valine could clash with the R1 substituent of the cephalosporin (Fig. 6B). Consistent with this hypothesis, an A501T mutation, which introduces a similar branched side chain, affects resistance in a similar manner as A501V (MU, unpublished observations).

The k_2/K_S acylation rate constants of the three antibiotics for wild type PBP 2, PBP 2^{35/02} and PBP 2^{35/02}-A501V were determined to assess directly the effects of the PBP mutations on reactivity with β -lactam antibiotics. The k_2/K_S rate constants for acylation are largely, but not entirely, consistent with MIC values. For example, acylation rates are dramatically impaired in PBP 2^{35/02} compared to wild type, with 150- to 200-fold decreases in acylation rate for all three antibiotics. However, since the MICs increase 10-, 20- and 100-fold for

penicillin, ceftriaxone, and cefixime, respectively, when *penA35* is transformed into FA19, it is clear that other intrinsic factors, e.g. diffusion through wild type porin channels, likely play an important role in defining the MIC. One anomaly in the acylation data is the ~2-fold increase in k_2/K_S of ceftriaxone for the PBP 2^{35/02}-A501V variant, which is inconsistent with the ~2-fold increase in the MIC for transformants harboring this mutant.

In conclusion, the emergence of decreased susceptibility to expanded-spectrum cephalosporins through remodeling of the active site of PBP 2 is more complicated than originally envisioned, with important resistance-conferring mutations showing epistasis. The acquisition of further mutations, such as A501V, in mosaic *penA* alleles is likely to increase resistance to levels that render expanded-spectrum cephalosporins ineffective in treating gonococcal infections. The need to identify new antibiotics against this organism is therefore of prime importance in the treatment of STIs.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Abbreviations and Textual Footnotes

PBP	penicillin-binding protein
MIC	minimum inhibitory concentration
ceph^I	cephalosporin-intermediate resistance
NTA	nitriloacetic acid
TEV	tobacco etch virus
STI	sexually transmitted infection

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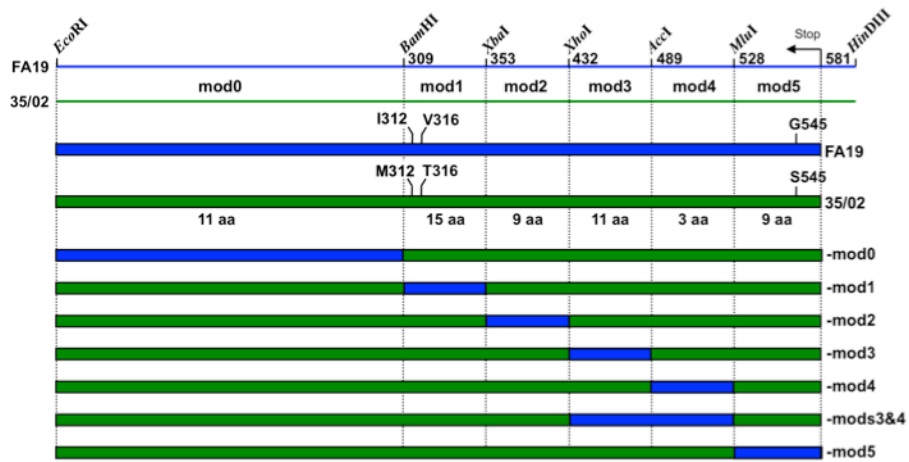


Fig. 1. Chimeric *penA* genes used in this study

Silent restriction sites were incorporated into the coding sequences of the *penA* genes from FA19 (blue) and 35/02 (green). The modules, designated mod0 through mod5, were used to create chimeric *penA* genes in which modules from *penA35* were replaced with the corresponding modules from wild type *penA*. These chimeric constructs were then used to create the strains listed in Table 1. The number of amino acid alterations in *penA35* relative to *penA* for each module is shown below 35/02. The lines represent DNA, and the rectangles represent the proteins encoded.

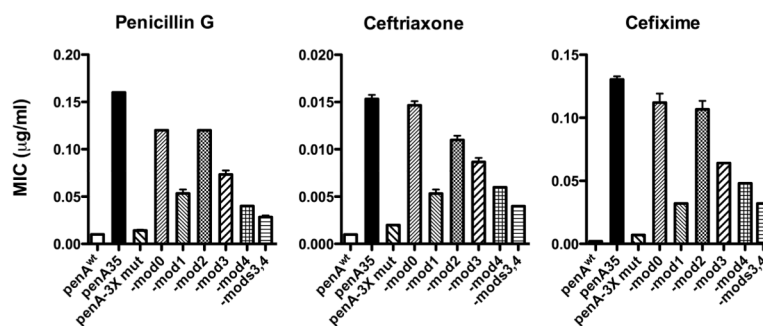


Fig. 2. MICs of penicillin, ceftriaxone, and cefixime for FA19 harboring 3X mutant and chimeric *-mod penA* genes

The MICs of penicillin, ceftriaxone, and cefixime for FA19 transformants containing the indicated 3X mutant (*penA*-I312M/V316T/G545S) and chimeric (*-mod*) *penA* genes (see Fig. 1 and Table 1) were determined as described in Materials and Methods. The MICs represent the averages for at least two transformants in a minimum of three independent experiments, and error bars represent the standard deviation of the values.

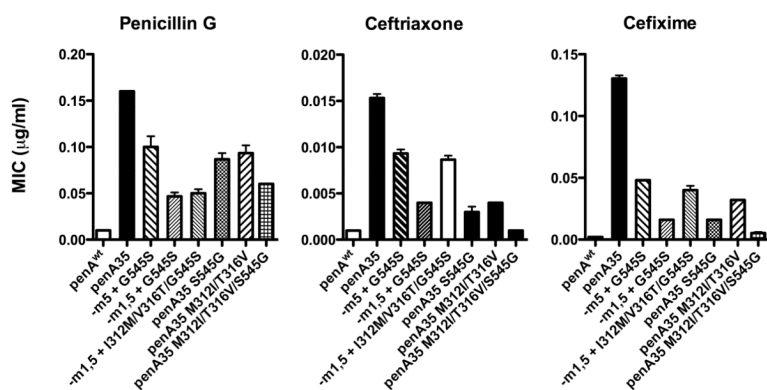


Fig. 3. The G545S and I312M/V316T mutations are critical for conferring resistance to penicillin and expanded-spectrum cephalosporins only when present in the *penA35* background
 The contributions of the G545S and I312M/V316T mutations within their respective modules were probed by either incorporating the mutations in the $-mod1$, $-mod5$, and $-mod1,5$ chimeric constructs or by reverting the mutations back to wild type in the *penA35* gene. The indicated constructs were transformed into FA19 and the MICs of penicillin, ceftriaxone, and cefixime were determined.

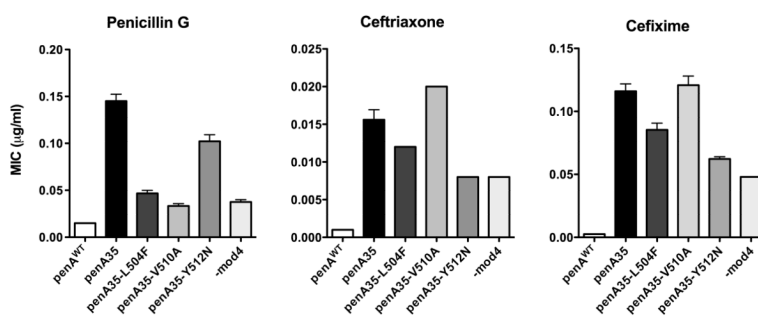


Fig. 4. MICs of penicillin, ceftriaxone, and cefixime for FA19 harboring *penA35* containing reversion mutations from module 4

FA19 was transformed with the indicated *penA35* alleles in which the three mutations in module 4 (F504L, A510V, and N512Y) were reverted individually back to wild type. The MICs of penicillin G, ceftriaxone, and cefixime of the resulting transformants were determined as described in Materials and Methods.

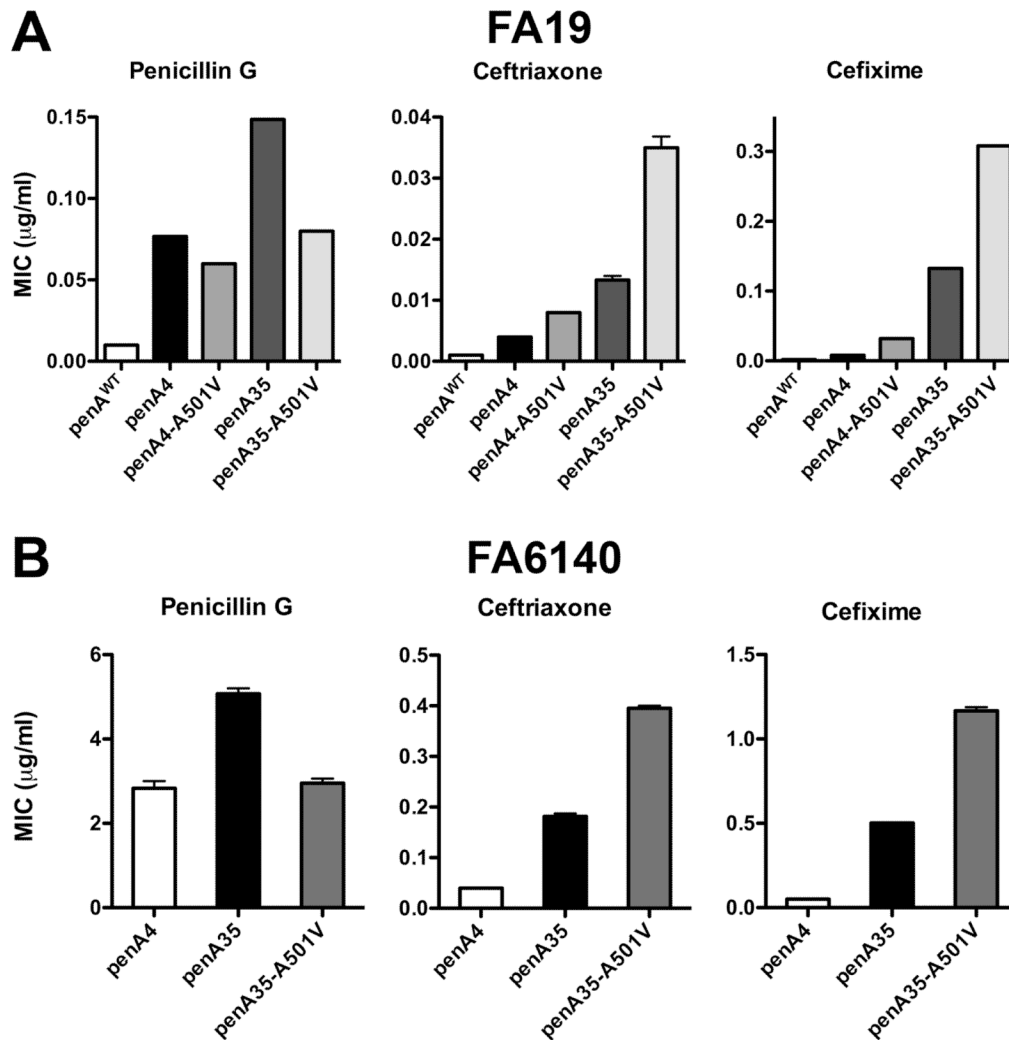


Fig. 5. MICs of penicillin, ceftriaxone, and cefixime for FA19 and FA6140 harboring the *penA4* (from FA6140) or *penA35* (from 35/02) allele with or without an A501V mutation. FA19 (A) or FA6140 (B) was transformed with *penA4*, *penA4*-A501V, *penA35*, or *penA35*-A501V, and the MICs of penicillin, ceftriaxone, and cefixime for the resulting transformants were assessed as described in Materials and Methods.

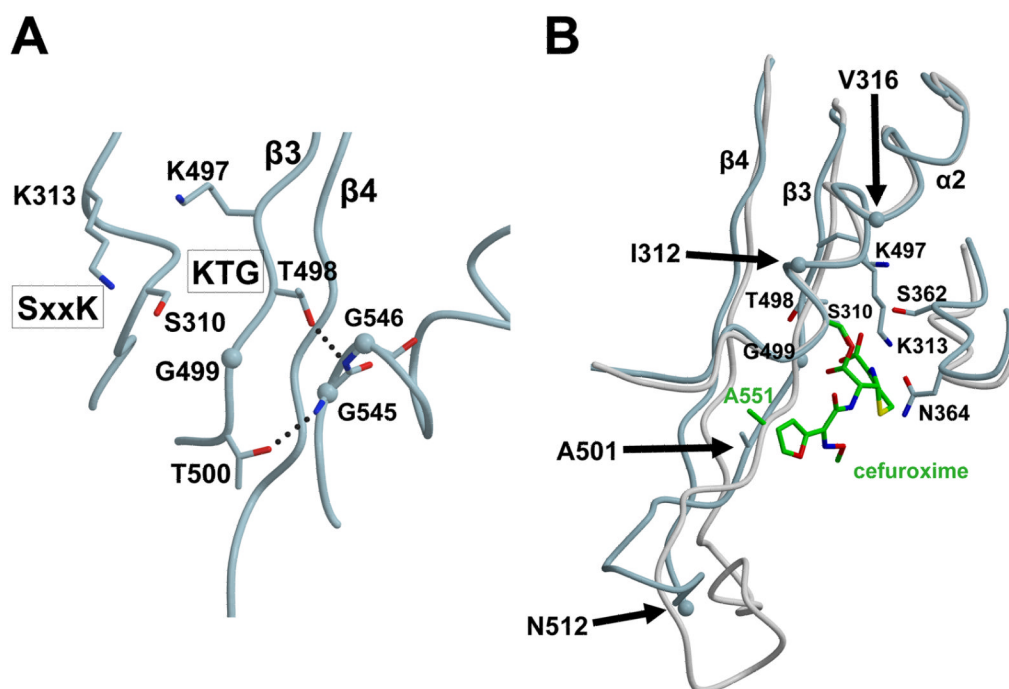


Fig. 6. Structure of PBP 2 showing the location of important mutations and the active site sequence motifs

A, Interaction of Thr498 and Thr500 within the KTG(T) active site motif with the main chain amides of Gly545 and Gly546. Mutation of Gly545 to Ser incorporates a hydroxylated side chain that potentially could perturb the interactions with the two Thr residues on $\beta 3$ with the main chain of the $\alpha 11$ helix. B, The crystal structure of apo-PBP 2 from strain CDC84 of *N. gonorrhoeae* (2.1Å, unpublished data) was superimposed onto that of *S. pneumoniae* PBP 2X in complex with cefuroxime (34) using the SUPERPOSE program of CCP4 (35). The structure of PBP 2 is from a construct containing the six C-terminal mutations (but missing the Asp345a insertion) of strain CDC-84 primarily because its $\beta 3$ - $\beta 4$ loop is more ordered than other PBP 2 structures. In this calculation, the main chain atoms of 18 residues comprising the three conserved active site motifs superimposed with a root mean square difference of 0.97Å. Note the proximity of the R1 furyl group of cefuroxime to the $\beta 3$ strand nearest to the A501V mutation. The view in this image is from underneath the $\alpha 2$ helix containing the active site SxxK motif. Also shown are the locations of the I312M, V316T, and N512Y mutations discussed in the text.

Table 1

Strains and plasmids used in this study.

Strain or Plasmid	Description	Reference
Plasmids		
pUC18us- <i>penA35</i> *	Plasmid containing the <i>penA35</i> gene from strain 35/02 and an uptake sequence	(16) and this study
pUC18us- <i>penA</i> *	Plasmid containing the <i>penA</i> gene from strain FA19 and an uptake sequence	(16) and this study
pUC18us- <i>penA35</i> *-A501V	pUC18us- <i>penA35</i> * harboring an A501V mutation	This study
pUC18us- <i>penA4</i>	Plasmid containing the <i>penA4</i> gene from strain FA6140 and an uptake sequence	(16)
pUC18us- <i>penA4</i> -A501V	pUC18us- <i>penA4</i> harboring an A501V mutation	This study
Strains		
FA19	Penicillin- and cephalosporin-susceptible recipient strain	(22)
FA6140	Penicillin-resistant but cephalosporin-susceptible recipient strain	(23)
WT-3X	FA19 transformed with pUC18us- <i>penA</i> containing I312M, S316V, and G545S mutations	This study
-mod0, -mod1, etc.	FA19 transformed with pUC18us- <i>penA35</i> * in which the indicated module (see Fig. 1) was replaced with the corresponding module from <i>penA</i> *	This study
-mod5 + G545S	FA19 transformed with pUC18us- <i>penA35</i> * in which mod5 was replaced with corresponding mod5 from wild type; mod5 also contains G545S mutation	This study
-m1,5 + G545S	FA19 transformed with pUC18us- <i>penA35</i> * in which mods 1 and 5 were replaced with corresponding mods from wild type; mod5 also had a G545S mutation	This study
-m1,5 + I312M/V316T/G545S	FA19 transformed with pUC18us- <i>penA35</i> * in which mods 1 and 5 were replaced with corresponding mods from wild type; mod1 additionally had I312M and V316T mutations and mod5 had a G545S mutation	This study
FA19 <i>penA35</i> -S545G	FA19 containing the <i>penA35</i> gene with reversion of the G545S mutation	This study
FA19 <i>penA35</i> -M312I/T316V	FA19 containing the <i>penA35</i> gene with reversion of I312M/V316T mutations	This study
FA19 <i>penA35</i> -S545G/M312I/T316V	FA19 containing the <i>penA35</i> gene with reversion of G545S/I312M/V316T mutations	This study

* Genes contains silent restriction sites incorporated into the coding sequence as depicted in Fig. 1.

Table 2
 k_2/K_S acylation rates for wild type PBP 2, PBP 2^{35/02}, and PBP 2^{35/02}-A501V

Proteins were purified and the k_2/K_S values were determined as described in Materials and Methods. The values shown are average \pm standard deviation (number of determinations).

PBP 2 Protein	k_2/K_S values ($M^{-1}s^{-1}$)		
	Penicillin	Ceftriaxone	Cefixime
Wild Type	75,700 \pm 2,300 (7)	1,710,000 \pm 90,000 (3)	1,480,000 \pm 22,000 (3)
35/02	510 \pm 90 (12)	11,300 \pm 400 (3)	7,200 \pm 300 (4)
35/02-A501V	1,400 \pm 140 (6)	20,000 \pm 400 (3)	3,100 \pm 100 (4)