

Diversity of *penA* Alterations and Subtypes in *Neisseria gonorrhoeae* Strains from Sydney, Australia, That Are Less Susceptible to Ceftriaxone[∇]

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Increasing numbers of *Neisseria gonorrhoeae* strains with decreased susceptibilities to ceftriaxone and other oral cephalosporins widely used for the treatment of gonorrhea have been isolated in Sydney, Australia, over several years. In this study, we examined the complete penicillin-binding protein 2 (PBP 2) amino acid sequences of 109 gonococci, selected on the basis of their diverse temporal and geographic origins and because they exhibited a range of ceftriaxone MICs: ≤ 0.03 $\mu\text{g/ml}$ ($n = 59$), 0.06 $\mu\text{g/ml}$ ($n = 43$), and 0.125 $\mu\text{g/ml}$ ($n = 7$). Auxotyping, serotyping, and genotyping by *N. gonorrhoeae* multiantigen sequence typing sequence-based analysis was also performed. In total, 20 different amino acid sequence patterns were identified, indicating considerable variation in the PBP 2 sequences in this study sample. Only some of the *N. gonorrhoeae* isolates with significantly higher ceftriaxone MICs contained a mosaic PBP 2 pattern, while more isolates exhibited a nonmosaic PBP 2 pattern containing an A501V substitution. Although particular *N. gonorrhoeae* genotypes in our sample were shown to be less susceptible to ceftriaxone, the reduced susceptibility to ceftriaxone was not specific to any particular genotype and was observed in a broad range of auxotypes, serotypes, and genotypes. Overall, the results of our study show that *N. gonorrhoeae* strains exhibiting reduced sensitivity to ceftriaxone are not of a particular subtype and that a number of different mutations in PBP 2 may contribute to this phenomenon.

The options for the effective treatment of *Neisseria gonorrhoeae* infections have been significantly reduced over time by the emergence and spread of gonococci resistant to the penicillins, tetracyclines, and the macrolide and quinolone groups of antibiotics (22). Cephalosporin antibiotics are now the mainstay of treatment in many settings, and the injectable agent ceftriaxone is used extensively in Australia and elsewhere. However, over recent years, reports, most notably from Japan (1, 8, 21), but also elsewhere (7, 27), of *N. gonorrhoeae* strains with decreased susceptibility to cephalosporin agents have appeared with increasing frequency.

The mechanism of gonococcal resistance to ceftriaxone is still unclear. *N. gonorrhoeae* has two penicillin-binding proteins (PBPs 1 and 2) that are the major targets of beta-lactam antibiotics. PBP 2, which is encoded by the *penA* gene, has a 10-fold greater affinity for penicillin than PBP 1 (6). Thus, mutations in PBP 2 are the major contributor to the emergence of chromosomally mediated resistance to beta-lactam antibiotics in *N. gonorrhoeae*, and particular attention has been paid to the *penA* gene. Alterations in *penA* that affect the activities of beta-lactam antibiotics have long been recognized. Insertion

of a single aspartic acid residue preceding Asp-345 (Asp-345A) in PBP 2 resulted in the expression of an altered PBP 2 and decreased susceptibility to penicillins (3). The acquisition by gonococci of other mutations and sequential alterations to other genes was shown to result in cumulative high-level chromosomal resistance to the penicillins (18), but these changes were said to have but little effect on the clinical efficacies of cephalosporin antibiotics (8).

Ameyama et al. (1) described a mosaic *penA* gene that was associated with reduced susceptibility to a number of beta-lactam agents, including cephalosporins, in gonococci from Tokyo, Japan. The mosaic *penA* gene was thought to be acquired by gonococci from commensal *Neisseria* strains by in vivo transformation (19). Reductions in the susceptibilities of in vitro transformants with this mosaic sequence were much greater for oral cephalosporins than for ceftriaxone. Ito et al. (8) examined the *penA* sequences of 70 clinical isolates of *N. gonorrhoeae* from Gifu, Japan, with a range of susceptibilities to ceftriaxone and other beta-lactam agents and described 10 PBP 2 patterns of sequence alteration numbered patterns I to X. The distribution of MICs for penicillin, the oral agents cefixime and cefdinir, and the injectable agent ceftriaxone was correlated with the *penA* patterns determined. Isolates with patterns I to IX contained the Asp-345A insertion, but isolates with pattern X did not; rather, they possessed the mosaic PBP 2 sequence described by Ameyama et al. (1). The MICs of ceftriaxone for gonococci with pattern X that contained the

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mosaic *penA* gene ranged from 0.015 to 0.25 $\mu\text{g/ml}$ but were said to be “significantly higher” than the MICs of isolates with patterns I to IX.

Tanaka et al. (21) examined the alterations present in a multiresistant gonococcus isolated in Fukuoka City, Japan, that was also less susceptible to ceftriaxone (MIC, 0.5 $\mu\text{g/ml}$). That group found mutations present in *penA*, *mtrR*, *penB*, and *ponA* and postulated that mutations that confer high-level resistance to penicillin also confer resistance to cephalosporins, including ceftriaxone. Although their isolate possessed the mosaic gene described by Ameyama et al. (1) and Ito et al. (8), it also lacked the Asp-345A substitution in PBP 2. Lindberg et al. (12) also noted that polymorphisms were present in *penA*, *mtrR*, *penB*, and *ponA* in 18 gonococci less susceptible to ceftriaxone and concluded that an “unambiguous” association existed between the increased ceftriaxone MICs, these polymorphisms, and *penA* mosaic alleles. Takahata et al. (20) more recently demonstrated that particular additional amino acid substitutions in the mosaic PBP 2 sequence were associated with decreased susceptibility to cefixime but had little effect on susceptibility to ceftriaxone. That study also found only minor reductions in ceftriaxone susceptibility in transformation experiments that introduced the full-length mosaic into a susceptible strain. We (26) described a second mosaic PBP 2 sequence with an Asp-345A change, but without the three key substitutions described by Takahata et al. (20), in a gonococcus fully susceptible to ceftriaxone and suggested that the extent of the contribution of mosaic PBP 2 sequences to reduced ceftriaxone susceptibility was yet to be fully elucidated.

In this study, selected gonococci that were isolated between 1997 and 2005 in Sydney, Australia, and that had altered susceptibilities to ceftriaxone were examined together with isolates susceptible to ceftriaxone. We explored further the relationship between *penA* alteration and ceftriaxone susceptibility by sequencing the *penA* genes of all isolates. Ito et al. (8) and a Danish group (7) have also suggested that the spread of these gonococci with decreased susceptibilities to ceftriaxone may be the result of the wider dissemination of a limited number of gonococcal subtypes. We therefore typed and compared all isolates by phenotypic and genotypic procedures and included in the selection of the strains studied a number of gonococci that were examined in Australia but that were isolated from patients who acquired their infection in a number of different countries.

MATERIALS AND METHODS

***N. gonorrhoeae* strains.** The South Eastern Area Laboratory Services laboratory in Sydney is the reference laboratory for *N. gonorrhoeae* for the state of New South Wales, Australia. It undertakes the primary isolation of *N. gonorrhoeae* from clinical samples from a number of sexual health clinics and hospitals and also receives isolates from public- and private-sector laboratories throughout the state. Approximately 1,200 gonococcal isolates are examined and stored annually, as described previously (11). A total of 109 gonococcal clinical isolates were retrospectively selected on the basis of their ceftriaxone MICs and their geographical and temporal diversities. The ceftriaxone MICs for the isolates were in the range of ≤ 0.008 to 0.03 $\mu\text{g/ml}$ ($n = 59$), 0.06 $\mu\text{g/ml}$ ($n = 43$), and 0.125 $\mu\text{g/ml}$ ($n = 7$). The penicillin G MICs were in the range of < 0.125 $\mu\text{g/ml}$ ($n = 6$), ≥ 0.125 $\mu\text{g/ml}$ to < 0.5 $\mu\text{g/ml}$ ($n = 18$), and ≥ 1.0 $\mu\text{g/ml}$ ($n = 75$); and another 10 isolates were penicillinase-producing gonococci (see Table 3). The isolates examined included local and international strains; 13 isolates were obtained from various parts of Asia (China, $n = 1$; Indonesia, $n = 1$; Japan, $n = 3$; Korea, $n = 5$ [28]; Mongolia, $n = 1$ [13]; not specified, $n = 2$), and the remaining 96 strains

were collected from various regions of the Australian state of New South Wales. The years of isolation included 1988 ($n = 3$), 1997 ($n = 1$), 1999 ($n = 1$), 2000 ($n = 3$), 2001 ($n = 12$), 2002 ($n = 11$), 2003 ($n = 7$), 2004 ($n = 12$), and 2005 ($n = 59$). Included among the isolates examined were gonococci used as reference cultures in World Health Organization and Australian quality assurance programs (2, 24, 27) and gonococci less susceptible to ceftriaxone, which were kindly supplied by T. Muratani (16).

Nucleotide sequencing of *penA* genes. The full-length *penA* gene sequences of the isolates were amplified and sequenced as described previously (8). Briefly, three separate PCRs were used to amplify three fragments of the *penA* gene. The PCR products were purified and then sequenced by using an ABI PRISM BigDye sequencing kit (Applied Biosystems).

Phenotypic and genotypic characterization and MIC determinations. Phenotypic and genotypic characterization and MIC determinations were performed for all 109 isolates as described previously (11, 24). Briefly, the extended phenotype of each isolate was derived by determination of the antibiotic susceptibility by the agar plate dilution methods of the Australian Gonococcal Surveillance Programme (24), which uses Sensitest agar (Oxoid, Basingstoke, United Kingdom) with 8% saponin-lysed horse blood as the test medium and an inoculum of 10^4 CFU per spot, and by auxotype/serovar classification. The ceftriaxone MIC was determined in triplicate examinations on separate occasions for those gonococci for which the MIC was raised. Auxotyping was done by the method of La Scolea and Young (10), and serovar determinations were done with a panel of 14 monoclonal reagents (Boule, Huddinge, Sweden) and by use of the nomenclature of Bygdeman (4). Genotyping was performed by *N. gonorrhoeae* multiantigen sequence typing (NG-MAST), as described previously (15).

Statistical analysis. Data analysis was performed by using the Mann-Whitney U test. Significance was set at a P value of < 0.05 .

RESULTS

Mutations of PBP 2 in clinical isolates. The complete nucleotide sequences of the *penA* genes of 109 clinical isolates of *N. gonorrhoeae* were determined. In total, 20 different amino acid sequence patterns were identified (patterns I to XXIII; Fig. 1), indicating the presence of considerable variation among the PBP 2 proteins in this study sample. Briefly, the first 340 amino acids of all 20 patterns except pattern X were identical to those in the GenBank reference sequence (accession number M32091). Various substitutions in the remaining sequence, particularly amino acids 500 to 580, accounted for most of the variation in the sequence patterns observed. Sequence patterns I to X were previously described by Ito et al. (8), whereas patterns XI to XXIII were identified in this study. Overall, sequence pattern XII was the most common (28 isolates), followed by patterns X and XIII (11 isolates each). Patterns III, VI, and VIII, previously described by Ito et al. (8), were not identified in this sample.

Sequence pattern X ($n = 11$) represented the previously described mosaic PBP 2 sequence, confirming the presence of this sequence in Australian gonococcal strains. Sequence pattern XXIII was a variation of the previously described pattern X mosaic sequence (26) (Fig. 1). Briefly, the first 430 amino acids of this pattern XXIII mosaic sequence were nearly identical to those in the GenBank reference sequence (accession number M32091) except for an additional Asp-345A codon, whereas the remaining amino acids were similar to those of the pattern X mosaic sequence but with the three substitutions described by Takahata et al. (20).

Association of PBP 2 with antimicrobial susceptibility. Overall, considerable variation in the ceftriaxone and penicillin G MICs was observed for each of the PBP 2 sequence patterns (Table 1). Significant associations with ceftriaxone susceptibility were observed for five PBP 2 patterns. Significantly higher ceftriaxone MICs were observed for the *N. gonorrhoeae* strains

	330	340	350	360	370	380	390	400	410
M32091	LDAGKTDLNE	RLNTQPYKIG	PSPVR-DTHV	YPSLDVRGIM	QKSSNVGTSK	LSARFGAEEM	YDFYHELIGIG	VRMHSGFPGE	TAGLLRNWRR
I (6)D....
II (5)D....
IV (2)D....
V (9)D....
VII (3)D....
IX (9)D....
X (11)	.S..V.ATD	TF..L....	SAT.Q-	..T.....M.TPK..	.D..V....S....
XI (2)D....
XII (28)D....
XIII (11)D....
XIV (1)D....
XV (2)	-.....
XVI (1)D....
XVII (3)D....
XVIII (3)D....
XIX (3)D....
XX (3)D....
XXI (3)D....
XXII (3)D....
XXIII (1)D....
	420	430	440	450	460	470	480	490	500
M32091	WRPIEQATMS	FGYGLQLSLL	QLARAYTALT	HGVLPLLSF	EKQAVAPQGK	RIFKESTARE	VRNLMVSVTE	PGGTGTAGAV	DGFDVGAKTG
I (6)
II (5)
IV (2)
V (9)
VII (3)
IX (9)
X (11)	.QK.....V....	..E..V..	..K....	.VI.A..KK	.E.....	.A.....
XI (2)
XII (28)
XIII (11)
XIV (1)
XV (2)
XVI (1)
XVII (3)
XVIII (3)
XIX (3)
XX (3)L....
XXI (3)
XXII (3)
XXIII (1)V....	..E..V..	..K....	.VI.A..KK	.E.....
	510	520	530	540	550	560	570	580	
M32091	TARKFVNGRY	ADNKHVATFI	GFAPAKNPRV	IVAVTIDEPT	AHGYYGGVVA	GPPFKKIMGG	SLNILGISPT	KPLT-AAAVK	TPS
I (6)
II (5)	..L....	V....G..
IV (2)	..L....	V....G..S.....
V (9)	..L....	V....G..S.....V....	.NV....
VII (3)	.V..L....	V....G..S.....
IX (9)	..L....	V....G..L.....
X (11)	..L....	V.Y....N..S..T	.V..QV..	..V....	.NV....
XI (2)	.V..L....	V....G..L.....
XII (28)	..L....	V....G..S.....
XIII (11)	.V..L....	V....G..S.....
XIV (1)	..L....	V....G..N.....
XV (2)N.....
XVI (1)	..L....	V....G..NV....
XVII (3)	.V..L....	V....G..S.....V....	.NV....
XVIII (3)	.T..L....	V....G..S.....V....	.NV....
XIX (3)	..L....	V....G..N.....V....	.NV....
XX (3)	..L....	V....G..N.....V....	.NV....
XXI (3)	.V..L....	V....G..V..QV..	..V....	.NV....
XXII (3)	..L....	V....G..N.....	.V..QV..	..V....	.NV....
XXIII (1)	..L....	V.Y....N..T..	.V..LV..	..V....	.NV....

FIG. 1. Partial PBP 2 amino acid sequences (positions 321 to 581) from the 109 *N. gonorrhoeae* isolates used in this study. The sequences are classified into different amino acid patterns (patterns I to XXIII) and are aligned with a GenBank *N. gonorrhoeae* sequence (accession number M32091). Sequence patterns I to X were previously described by Ito et al. (8), whereas patterns XI to XXIII were identified in this study. The numbers of isolates with each pattern are indicated in parentheses. Patterns III, VI, and VIII, previously described by Ito et al. (8), were not identified in this sample.

TABLE 1. Ceftriaxone MICs for the various PBP 2 amino acid patterns and associated *N. gonorrhoeae* NG MAST sequence types

PBP 2 pattern	No. of isolates with the following ceftriaxone MIC ($\mu\text{g/ml}$):						No. of NG MAST STs ^a
	≤ 0.008	0.016	0.03	0.06	0.12	Total	
I	6					6	6
II	5					5	3
IV				1	1	2	2
V	3		3	3		9	5
VII				2	1	3	3
IX	1	1		7		9	3
X			1	10		11	7
XI				2		2	2
XII	9	5	10	4		28	12
XIII				8	3	11	7
XIV	1					1	1
XV	2					2	2
XVI	1					1	1
XVII				2	1	3	2
XVIII				2	1	3	2
XIX	2	1				3	3
XX		1		2		3	1
XXI	3					3	2
XXII	2	1				3	3
XXIII	1					1	1
Total	36	9	14	43	7	109	64

^a STs, sequence types.

with the mosaic PBP 2 pattern X ($P < 0.01$), as well as those with pattern XIII ($P < 0.001$). Significantly lower ceftriaxone MICs were observed for strains with patterns I ($P < 0.01$), II ($P < 0.01$), and XII ($P < 0.05$). Ten of the 11 strains with mosaic pattern X were less susceptible to ceftriaxone (MICs = 0.06 $\mu\text{g/ml}$), whereas one strain with pattern X and the one strain with variant mosaic pattern XXIII were both sensitive (MICs = 0.03 $\mu\text{g/ml}$ and 0.008 $\mu\text{g/ml}$, respectively; Table 1).

Significant associations with penicillin G susceptibility were observed for five PBP 2 patterns (Table 2). Significantly higher penicillin G MICs were observed for *N. gonorrhoeae* strains with mosaic PBP 2 pattern X ($P < 0.001$) as well as those with pattern XX ($P < 0.01$). Strains with patterns I, XV, and XXII demonstrated significantly lower penicillin G MICs ($P < 0.01$). Only 4 of 95 *N. gonorrhoeae* strains with the additional Asp-345A codon were susceptible to penicillin G, including strains with PBP 2 patterns I (1 isolate), XII (1 isolate), and XXII (2 isolates). The two gonococcal strains with PBP 2 pattern XV, which lacks the Asp-345A codon, were both susceptible to penicillin G, with low MICs (MIC = 0.0016 $\mu\text{g/ml}$).

Association of *N. gonorrhoeae* typing with ceftriaxone susceptibility. There was considerable diversity among the isolates tested, with seven auxotypes, 23 serotypes, and 64 NG MAST sequence types observed. The PBP 2 amino acid patterns, ceftriaxone MICs, auxotypes, serotypes, and sequence types for the 50 *N. gonorrhoeae* isolates with reduced susceptibilities to ceftriaxone are provided in Table 3. Briefly, the strains with reduced susceptibilities to ceftriaxone did not have any particular type and were found to have a broad range of auxotypes, serotypes, and genotypes.

Also of note was the presence of 14 isolates of sequence type 225, all with PBP 2 pattern XII. The ceftriaxone MICs for this sequence type were ≤ 0.008 $\mu\text{g/ml}$ ($n = 6$), 0.016 $\mu\text{g/ml}$ ($n = 4$),

TABLE 2. Penicillin G MICs for the various PBP 2 amino acid patterns

PBP 2 pattern	No. of isolates with the following penicillin G MIC ($\mu\text{g/ml}$) or characteristic:							Total
	≤ 0.06	0.125	0.25	0.5	1	2	PPNG ^a	
I	1	3	1			1		6
II		1	3				1	5
IV			1			1		2
V					5	4		9
VII					1	2		3
IX			1		7	1		9
X					2	9		11
XI				1	1			2
XII	1			2	19	5		27
XIII					9	1	1	11
XIV							1	1
XV	2							2
XVI							1	1
XVII					2	1		3
XVIII						3		3
XIX					1		2	3
XX							3	3
XXI			1	1			1	3
XXII	2			1				3
XXIII			1					1
Total	6	4	8	5	47	28	10	

^a PPNG, penicillinase-producing *N. gonorrhoeae*.

and 0.03 $\mu\text{g/ml}$ ($n = 4$). There were another 14 isolates also of PBP 2 pattern XII but of seven other sequence types (data not shown) and with ceftriaxone MICs ≤ 0.008 $\mu\text{g/ml}$ ($n = 3$), 0.016 $\mu\text{g/ml}$ ($n = 1$), 0.03 $\mu\text{g/ml}$ ($n = 6$), and 0.06 $\mu\text{g/ml}$ ($n = 4$). Sequence type 225 was the "clone" identified in the Danish cluster (7), but no PBP 2 sequencing was performed in that study.

DISCUSSION

This study shows the considerable diversity that exists in PBP 2 proteins in *N. gonorrhoeae* strains less susceptible to ceftriaxone. The 20 different amino acid sequences detected included the 7 previously described by Ito et al. (8) and 13 additional variations. Many of the individual PBP 2 sequence patterns displayed a range of ceftriaxone MICs and were found in multiple gonococcal subtypes, although some were associated with significantly higher (patterns X and XIII) or lower (patterns I, II, and XII) ceftriaxone MICs. As demonstrated in a previous study (8), we also found a significant association between the presence of the pattern X mosaic sequence and reduced ceftriaxone susceptibility. However, in contrast to the previous report (8), the mosaic sequences in our sample did not account for a large proportion of strains with reduced sensitivities to ceftriaxone, and only 6 of the 50 strains with reduced sensitivities (MICs = 0.06 $\mu\text{g/ml}$ or more) in this study possessed the mosaic PBP 2 sequence (Table 3). Thus, the presence of a mosaic PBP 2 sequence, previously assigned a pivotal role in this phenomenon, could not account for the majority of instances of this reduced sensitivity to ceftriaxone.

We raised further questions regarding the role of mosaic sequences in conferring reduced ceftriaxone susceptibility when we described a strain with variant mosaic pattern XXIII that was fully susceptible to ceftriaxone (26). In this context, the highly significant association between PBP 2 pattern XIII

TABLE 3. Profiles for the 50 *N. gonorrhoeae* isolates with reduced susceptibilities to ceftriaxone (MIC \geq 0.06 μ g/ml), indicating the PBP 2 amino acid pattern, ceftriaxone MIC, auxotype, serotype, and genotype data^a

PBP 2 pattern	Ceftriaxone MIC (μ g/ml)	NG MAST ST	Auxotype	Serotype ^b	No. of isolates
IV	0.06	69	NR	Brpyut	1
IV	0.12	327	Pro	Brpust	1
V	0.06	1,410	Pro	Broypst	2
V	0.06	1,688	Pro	Broypst	1
VII	0.06	1,619	Pro	Brpyust	1
VII	0.06	1,622	Pro	Byust	1
VII	0.12	1,422	Pro	Brpyust	1
IX	0.06	5	NR	Broypst	6
IX	0.06	21	NR	Bopyst	1
X	0.06	1,414	Pro	Bpyust	3
X	0.06	1,424	Pro	Bpyust	3
X	0.06	1,677	Pro	Byust	1
X	0.06	326	Pro	Broypst	1
X	0.06	835	Pro	Bpyust	1
X	0.06	925	Pro	Boys	1
XI	0.06	1,420	Pro	Bro	1
XI	0.06	5	NR	Broypst	1
XII	0.06	1,426	NR	Brpust	1
XII	0.06	1,703	NR	Bopt	1
XII	0.06	604	NR	Broypst	1
XII	0.06	604	NR	Broypst	1
XIII	0.06	1,409	NR	Broypst	2
XIII	0.06	1,623	NR	Broypst	4
XIII	0.06	1,624	NR	Byvut	1
XIII	0.06	1,676	NR	Byvut	1
XIII	0.12	1,148	Pro	Broypst	1
XIII	0.12	1,413	NR	Broypst	1
XIII	0.12	1,425	NR	Broypst	1
XVII	0.06	1,342	Pro	Broypst	2
XVII	0.12	1,620	Pro	Broypst	1
XVIII	0.06	864	Pro	Bopt	2
XVIII	0.12	1,621	Pro	Broypst	1
XX	0.06	211	NR	Arst	2

^a Nine profiles were identified for more than one isolate. ST, sequence type; NR, nonrequiring; Pro, proline requiring.

^b Serotype by use of the nomenclature of Bygdeman (4).

and reduced susceptibility to ceftriaxone was also of particular interest. This PBP 2 pattern contained a substitution at position 501 (A501V), and all 11 strains with pattern XIII were less susceptible to ceftriaxone, with MICs of 0.06 μ g/ml or greater. In fact, 22 of the 25 strains in our sample containing a substitution at position 501 (A501V or A501T; patterns VII, XI, XIII, XVII, XVIII, and XXI) were less susceptible to ceftriaxone and comprised 6 of the 7 strains with MICs of 0.125 μ g/ml. During transformation experiments with mosaic *penA* sequences, Takahata et al. (20) found that a spontaneous mutant with a single A501V mutation was selected with cephalixin. Subsequent analysis revealed that this A501V mutation led to two- to fourfold increases in the MICs of cefixime and other cepheims (20). Ito et al. (8) also observed that all five strains with PBP 2 pattern VII were less susceptible to ceftriaxone (8). These changes at position 501 were also present in some gonococci less susceptible to ceftriaxone in the studies of Ito et al. (8) and Lindberg et al. (12) but were not specifically remarked upon by either group. Overall, this suggests that the mutations at position 501 are also of interest and, based on our data, are more significant as a contribution to ceftriaxone susceptibility than the mosaic sequences. However, questions are

again raised over the relevance of this mutation in isolation, given that all three strains in our study with pattern XXI (which contains the A501V mutation) were fully susceptible to ceftriaxone (MICs = 0.008 μ g/ml).

It was also shown in this study that while particular *N. gonorrhoeae* genotypes, e.g., sequence type 5, were associated with decreased susceptibility to ceftriaxone, in general, this reduced susceptibility to ceftriaxone was not restricted to any particular genotype and was observed in strains of a broad range of auxotypes, serotypes, and genotypes. Thus, the mechanisms responsible for the reduced susceptibility to ceftriaxone appear to be present in a broad range of gonococcal types and strains and were not associated with a particular subtype, as has been observed previously (7, 8). Both of those earlier studies were with gonococci isolated in a limited geographic area over a defined period, whereas this study was of gonococci selected for their temporal and geographic diversities. Other broadly based studies of the diagnostic features (11, 14) and the pathogenic features (17) of gonococci have also revealed a wider distribution of gonococcal subtypes with a particular characteristic than that revealed by earlier and more narrowly based studies. The findings of those earlier studies and those of this study are thus consistent with the known population dynamics of gonococci (9, 23), and the earlier data probably reflect the expansion of a particular subtype within a wider gonococcal population at a particular time and place. The presence in both Denmark and Australia of *N. gonorrhoeae* NG MAST sequence type 225 strains with some increase in the ceftriaxone MIC (MIC = 0.03 μ g/ml) also suggests the possibility of the international spread of these strains. The international spread of gonococci with other specific characteristics has also been documented recently (25).

The clinical significance of these findings is difficult to determine at present. The history of antimicrobial resistance in *N. gonorrhoeae* indicates that problems with any antibiotic that is used extensively for the treatment of *N. gonorrhoeae* disease will eventually be encountered (5, 22). The emergence of *N. gonorrhoeae* strains with reduced susceptibilities to ceftriaxone and other expanded-spectrum cephalosporins suggests that the gonococci are now developing mechanisms of resistance to yet another important group of antibiotics. These isolates with reduced susceptibilities to ceftriaxone are often also quinolone resistant (7), and this was also the case with the isolates studied here (data not shown). Although treatment failure with oral expanded-spectrum agents, including cefixime, which is not available in Australia, have been recorded in Japan (16), it should be noted that no documented treatment failures have been associated with the injectable agent ceftriaxone when the higher 250-mg dose recommended for use in Australia is used, even for strains containing the mosaic *penA* sequence (22). Thus, it would appear that additional resistance mechanisms are required before we will see gonococci that are fully resistant to ceftriaxone.

Overall, the results of our study show that mosaic PBP 2 sequences alone cannot account for the reduced ceftriaxone sensitivity observed in our *N. gonorrhoeae* strains and that strains exhibiting this phenotype are not necessarily of a particular subtype. Furthermore, the results suggest that gradual mutation in the *N. gonorrhoeae penA* gene sequence, in addition to genetic recombination, which occurs frequently in

gonococci (23), contributes not only to reduced susceptibility to ceftriaxone but also to the diversity of changes observed here. The separate phenomenon of strain diversity seen also, but separately, reflects the high rate of transformation that occurs in *N. gonorrhoeae*. Other mechanisms that may confer reduced susceptibility to newer cephalosporins, including mutations in the *penA* (PBP 1) gene, have been proposed (21). Thus, further investigation is needed to fully understand the impacts of these genetic mechanisms both alone and in combination. We emphasize that *N. gonorrhoeae* strains with reduced susceptibilities to ceftriaxone represent a potential threat to current treatment regimens. Hence, the continued spread of these strains needs to be closely monitored.

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